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# Synthases

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#### (54) SYNTHASES

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- (60) Provisional application No. 60/150,262, filed on Aug. 23, 1999, provisional application No. 60/130,628, filed on Apr. 22, 1999, and provisional application No. 60/100,993, filed on Sep. 18, 1998.
- (51) **Int. Cl.**<sup>7</sup> ...... **C12N 9/00**; C12N 9/88; C07N 21/04

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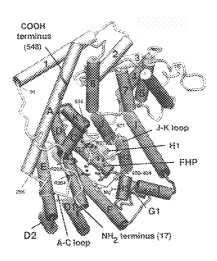
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#### (57) ABSTRACT

Novel synthases and the corresponding nucleic acids encoding such synthases are disclosed herein. Such synthases possess an active site pocket that includes key amino acid residues that are modified to generate desired terpenoid reaction intermediates and products. Synthase modifications are designed based on, e.g., the three-dimensional coordinates of tobacco 5-epi-aristolochene synthase, with or without a substrate bound in the active site.

## 23 Claims, 4 Drawing Sheets



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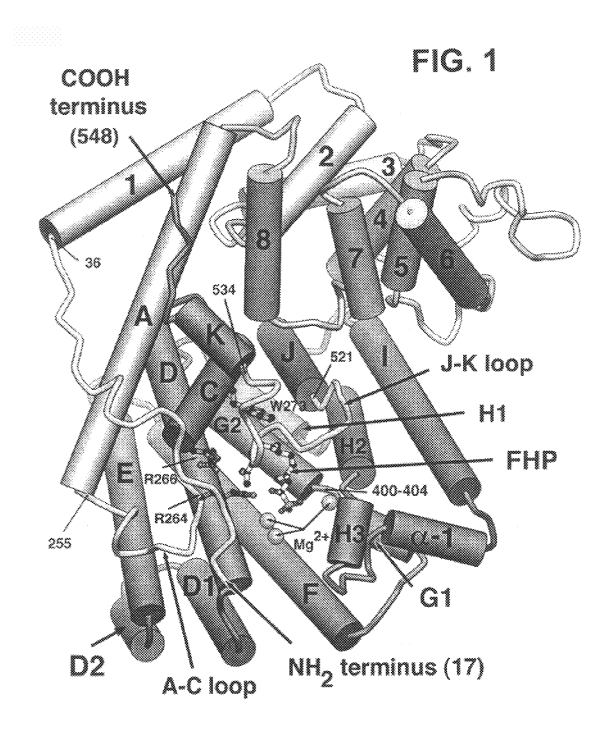
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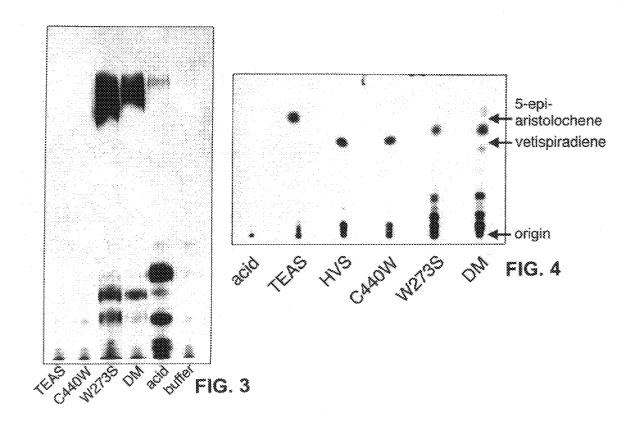
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Genbank Accession No: U20190.



May 6, 2003



## **SYNTHASES**

#### CROSS REFERENCE TO RELATED APPLICATIONS

This application is a divisional of and claims priority under 35 U.S.C. §120 to U.S. application Ser. No. 09/398, 395, filed Sep. 17, 1999, which claims benefit under 35 USC §119(e) of U.S. Provisional Application No. 60/150,262, filed Aug. 23, 1999, U.S. Provisional Application No. 60/130,628, filed Apr. 22, 1999, and U.S. Provisional Application No. 60/100,993, filed Sep. 18, 1998.

This work was supported, in part, with funding from NIH (GM54029 and GM07240) and NSF (IBW-9408152). Therefore, the Unites States Federal Government may have certain rights in the invention.

#### BACKGROUND OF THE INVENTION

Isoprenoid compounds are organic molecules produced by a wide range of organisms (e.g., plants, bacteria, fungi, etc). To date, over 23,000 individual isoprenoid molecules have been characterized with tens to hundreds of new structures identified each year. These molecules can fulfill a variety of roles. For example, monoterpenes can be used as fragrances and flavors. Sesquiterpenes and diterpenes can serve as pheromones, defensive agents, visual pigments, antitumor drugs, and components of signal transduction pathways. Triterpenes can serve important functions as membrane constituents and precursors of steroid hormones and bile acids. Polyprenols function as photoreceptive agents and cofactor side chains, and can also exist as natural polymers.

The diverse molecular compounds produced by the isoprenoid pathway are created from diphosphate esters of monounsaturated isoprene units. Isoprenes are added together in multiples of 2, 3, or 4 by prenyl transferases to make  $C_{10}$ ,  $C_{15}$ , and  $C_{20}$  units, respectively. The  $C_{10}$ ,  $C_{15}$ , and C<sub>20</sub> molecules, named geranyl diphosphate (GPP), farnesyl diphosphate (FPP), and geranylgeranyl diphosphate (GGPP), respectively, serve as substrates for terpene syn- 40 isoprenoid products. thases.

Terpene synthases catalyze the production of isoprenoid compounds via one of the most complex reactions known in chemistry or biology. In general, terpene synthases are about 40 to 100 kD. As an enzyme, terpene synthases can be classified as having low to moderate turnover rates coupled with exquisite reaction specificity and preservation of chirality. Turnover comprises binding of substrate to the enzyme, establishment of substrate conformation, conversion of substrate to product and product release. Reactions can be performed in vitro in aqueous solvents, typically require magnesium ions as cofactors, and the resulting products, which are often highly hydrophobic, can be recovered by partitioning into an organic solvent.

Terpene synthase genes are found in a variety of organisms including bacteria, fungi and plants. Swapping regions approximating exons between different terpene synthases has identified functional domains responsible for terminal enzymatic steps. For example, work performed on 5-epiaristolochene synthase (TEAS) from Nicotiana tabacum (tobacco) and Hyoscyamus muticus vetispiradiene synthase (HVS) from henbane revealed that exon 4 and exon 6, respectively, were responsible for reaction product specificity. Combining functional domains resulted in novel enzymes capable of synthesizing new reaction products (U.S. Pat. No. 5,824,774).

Studies have led to proposed reaction mechanisms for isoprenoid production; see, e.g., Cane et al., 1985, Bioorg. Chem., 13:246–265; Wheeler and Croteau, 1987, Proc. Natl. Acad. Sci. USA, 84:4856–4859; and Pyun et al., 1994, Arch. Biochem. Biophys., 308:488-496. The studies used substrate analogs and suicide inhibitors (Croteau, 1994, Arch. Biochem. Biophys., 251:777-782; Cane et al., 1995, Biochemistry, 34:2471-2479; and Croteau et al., 1993, Arch. Biochem. Biophys., 307:397-404), as well as 10 chemical-modifying reagents and site-directed mutagenesis in efforts to identify amino acids essential for catalysis (Cane et al., 1995, Biochemistry, 34:2480-2488; Rajaonarivony et al., 1992, Arch. Biochem. Biophys., 296:49-57; and Rajaonarivony et al., 1992, Arch. Biochem. Biophys., 15 299:77–82). However, these studies have resulted in limited success in defining the active site due to inherent limitations with these techniques.

#### SUMMARY OF THE INVENTION

The invention describes a method of identifying alphacarbon atoms found in the active site of a terpene synthase and describes these atoms in three-dimensional space as well as the spatial relationships among them. The present invention also describes R-groups associated with such alphacarbons and methods of altering these R-groups in order to create novel terpene synthases capable of generating novel reaction products.

Until the invention taught in this present application, the active site of synthase proteins, the amino acid residues located therein, the amino acid residues involved in catalysis, and the configuration of α-carbons and R-groups within the active site have not been known. The current invention now teaches the structure of synthases, as well as provides the means of making and using the information obtained therefrom to develop and produce new and novel synthases having new and novel synthetic capabilities. The data generated using the methods described herein are useful for creation and production of synthase mutants that can use a variety of isoprenoid substrates and produce a variety of

In one embodiment, the invention features an isolated terpene synthase having about 20% or greater sequence identity to residues 265 to 535 of SEQ ID NO: 2. Such a synthase comprises nine α-carbons having interatomic dismoderately sized enzymes having molecular weights of 45 tances in Angstroms between the α-carbons that are ±2.3 Angstroms of the interatomic distances shown in Table 6. The center point of each  $\alpha$ -carbon is positioned within a sphere having a radius of 2.3 Angstroms. The center point of each such sphere has the structural coordinates given in Table 5. Each α-carbon has an associated R-group, and the synthase has an ordered arrangement of R-groups associated with each alpha-carbon other than the ordered arrangements of R-groups shown in Table 9. The synthase can have about 25% or greater sequence identity to residues 265 to 535 of SEQ ID 2, or about 35% or greater sequence identity to residues 265 to 535 of SEQ ID 2. Such a synthase can catalyse the formation of a terpenoid product from a monoterpene substrate, a sesquiterpene substrate, or a diterpene substrate. The product can be a cyclic terpenoid hydrocarbon or an acyclic terpenoid hydrocarbon. Either type of product can be hydroxylated or non-hydroxylated. The R-group associated with  $\alpha$ -carbon 1 can be selected from one of the following groups: the group consisting of Cys, Ser, and Thr, the group consisting of Phe, Tyr and Trp, the group consist-65 ing of Pro, Gly, and Ala, the group consisting of Glu and Asp, the group consisting of Met, Ile, Val and Leu, the group consisting of Arg and Lys, and the group consisting of Gln,

As and His. R-groups associated with  $\alpha$ -carbons 2 to 9 can be any amino acid except those having the ordered arrangements of Table 9. Similarly, the R-group associated with each of  $\alpha$ -carbons 2–9 can be selected independently from the group consisting of Cys, Ser and Thr, the group consisting of Phe, Tyr and Trp, the group consisting of Pro, Gly, and Ala, the group consisting of Glu and Asp, the group consisting of Met, Ile, Val and Leu, the group consisting of Arg and Lys, and the group consisting of Gin, Asn and His. In these embodiments, R-groups associated with the remaining eight α-carbons except those having the ordered arrangements of Table 9.

In some embodiments, the ordered arrangement of R-groups associated with  $\alpha$ -carbons 1 to 9 is Trp, Ile, Thr, Thr, Tyr, Leu, Cys, Thr and Phe, respectively, Ser, Ile, Thr, Thr, Tyr, Leu, Cys, Thr and Tyr, respectively, Trp, Ile, Thr, Thr, Tyr, Leu, Trp, Thr and Tyr, respectively, Ser, Ile, Thr, Thr, Tyr, Leu, Trp, Thr and Tyr, respectively, or Glu, Ile, Thr, Thr, Tyr, Leu, Cys, Thr and Tyr, respectively.

The invention also features a terpene synthase made by 20 aligning the primary amino acid sequence of a preselected terpene synthase polypeptide to the amino acid sequence of residues 265 to 535 of SEQ ID NO: 2, mutating a nucleic acid encoding the preselected polypeptide at one or more codons for nine amino acid residues in a region of the polypeptide primary amino acid sequence having about 20% or greater sequence identity to residues 265 to 535 of SEQ ID NO: 2, the nine residues in the polypeptide aligning with residues 273, 294, 402, 403, 404, 407, 440, 519 and 520 of SEQ ID NO: 2; and expressing the mutated nucleic acid so 30 that a mutated terpene synthase is made.

The invention also features an isolated terpene synthase having about 20% or greater sequence identity to residues 265 to 535 of SEQ ID NO: 2, the synthase comprising sixteen a-carbons having interatomic distances in Angstroms between the  $\alpha$ -carbons that are  $\pm 2.3$  Angstroms of the interatomic distances given in Table 4. The center point of each α-carbon is positioned within a sphere having a radius of 2.3 Angstroms. The center point of each of the spheres has the structural coordinates given in Table 3. Each  $\alpha$ -carbon 40 has an associated R-group, and the synthase has an ordered arrangement of R-groups other than the ordered arrangements of R-groups given in Table 8. The synthase can have about 25% or greater sequence identity to residues 265 to identity to residues 265 to 535 of SEQ ID NO: 2. The synthase can catalyse the formation of a terpenoid product from a monoterpene substrate, a sesquiterpene substrate, or a diterpene substrate. The product can be, for example, a cyclic terpenoid hydrocarbon. The ordered arrangement of 50 R-groups in the synthase associated with  $\alpha$ -carbons 1 to 16 can be Cys, Trp, Ile, Ile, Ser, Thr, Thr, Tyr, Leu, Cys, Val, Thr, Tyr, Asp, Phe and Thr, respectively.

The invention also features an isolated terpene synthase having about 20% or greater sequence identity to residues 55 265 to 535 of SEQ ID NO: 2, the synthase comprising nineteen α-carbons having interatomic distances in Angstroms between the  $\alpha$ -carbons that are  $\pm 2.3$  Angstroms of the interatomic distances given in Table 2. The center point of each α-carbon is positioned within a sphere having a radius of 2.3 Angstroms. The center points of each sphere have the structural coordinates given in Table 1. Each α-carbon has an associated R-group, and the synthase has an ordered arrangement of the R-groups other than the ordered arrangements of R-groups given in Table 7. The synthase can have about 25% or greater sequence identity to residues 265 to 535 of SEQ ID NO: 2, or about 35% or greater sequence

identity to residues 265 to 535 of SEO ID NO: 2. The synthase can catalyse the formation of a terpenoid product from a monoterpene substrate, a sesquiterpene substrate, or a diterpene substrate. The product can be, for example, a cyclic terpenoid hydrocarbon.

The invention also features an isolated protein comprising a first domain having an amino terminal end and a carboxyl terminal end. The first domain comprises amino acids that align structurally in three-dimensional space with a glycosyl 10 hydrolase catalytic core, the glycosyl hydrolase catalytic core selected from the group consisting of amino acids 36 to 230 of glucoamylase protein databank (PDB) code 3GLY of Aspergillus awamori and amino acids 36 to 230 of endoglucanase CelD PDB code 1CLC. The isolated protein also comprises a second domain having an amino terminal end and carboxyl terminal end. The second domain comprises amino acids that align structurally in three-dimensional space with avian FPP synthase. The carboxyl terminal end of the first domain is linked to the amino terminal end of the second domain. The second domain has about 20% or greater sequence identity to residues 265 to 535 of SEQ ID NO: 2, and comprises nine  $\alpha$ -carbons having interatomic distances in Angstroms between the  $\alpha$ -carbons that are  $\pm 2.3$ Angstroms of the interatomic distances given in Table 6. The center point of each  $\alpha$ -carbon is positioned within a sphere having a radius of 2.3 Angstroms, the center point of each sphere having the structural coordinates given in Table 5. Each  $\alpha$ -carbon has an associated R-group, and the synthase has an ordered arrangement of R-groups other than the ordered arrangements of R-groups given in Table 9. The protein can have about 25% or greater sequence identity to SEQ ID NO: 2, or about 35% or greater sequence identity to SEQ ID NO: 2. The synthase can catalyse the formation of a terpenoid product from a monoterpene substrate, a sesquiterpene substrate, or a diterpene substrate. The product can be, for example, a cyclic terpenoid hydrocarbon.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 343 to 606 of SEQ ID NO: 20, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 348, 351, 372, 375, 376, 454, 479, 480, 481, 482, 485, 519, 523, 597, 600, 601, 605, 607 and 608 of SEQ ID NO: 20 are residues other than amino acids Y, L, C, I, T, Y, S, C, G, H, S, L, G, F, G, Y, D, Y and S, 535 of SEQ ID NO: 2, or about 35% or greater sequence 45 respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

> The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 316 to 586 of SEQ ID NO: 22, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 321, 324, 345, 348, 349, 427, 452, 453, 454, 455, 458, 492, 496, 569, 572, 573, 577, 579 and 580 of SEO ID NO: 22 are residues other than amino acids C, W, N, I, T, Y, S, I, S, G, M, L, D, A, M, Y, D, H and G, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more ordered arrangements of residues as given in Table 7 are not found in such a synthase.

> The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 352 to 622 of SEQ ID NO: 58, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 357, 360, 381, 384, 385, 463, 487,

488, 489, 490, 493, 528, 532, 606, 609, 610, 614, 616 and 617 of SEQ ID NO: 58 are residues other than amino acids Y, M, C, V, T, F, V, S, S, G, I, L, G, F, V, Y, D, Y and T, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to amino acid residues 272 to 540 encoded by SEQ ID NO: 33, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 277, 280, 301, 304, 305, 383, 408, 409, 410, 411, 414, 448, 452, 524, 527, 528, 532, 534 and 535 encoded by SEQ ID NOS: 33 are residues other than amino acids G, W, I, A, S, Y, T, S, G, Y, L, C, D, M, L, Y, D, Y and T, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 319 to 571 of SEQ ID NO: 42, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 324, 327, 348, 351, 352, 430, 455, 456, 457, 458, 461, 495, 499, 571, 574, 575, 579, 581 and 582 of SEQ ID NO: 42 are residues other than amino acids I, W, V, I, S, Y, T, T, G, L, V, I, N, T, S, Y, D, Y, and T, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 579 to 847 of SEQ ID NO: 44, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 584, 587, 606, 609, 610, 688, 713, 714, 715, 716, 719, 753, 757, 831, 834, 835, 839, 841 and 842 of SEQ ID NO: 44 are residues other than amino acids V, S, G, Q, V, Y, S, V, G, L, C, W, N, V, F, Y, D, Y and G, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 495 to 767 of SEQ ID NO: 46, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 500, 503, 524, 527, 528, 606, 631, 632, 633, 634, 637, 674, 678, 751, 754, 755, 759, 761 and 762 of SEQ ID NO: 46 are residues other than amino acids 557, F. L. A. Q. T. Y. S. I. G. Q. L. S. D. T. I. F. D. F. and G. respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 295 to 564 of SEQ ID NO: 48, wherein one or more amino acid residues of the synthase that align with amino 65 acid residues at positions 300, 303, 324, 327, 328, 406, 431, 432, 433, 434, 437, 471, 475, 548, 551, 552, 556, 558 and

559 of SEQ ID NO: 48 are residues other than amino acids Y, W, A, C, T, Y, S, S, G, M, L, G, D, L, I, Y, D, L and Y, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 307 to 578 of SEQ ID NO: 50, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 312, 315, 336, 339, 340, 419, 444, 445, 446, 447, 450, 484, 488, 562, 565, 566, 570, 572 and 573 of SEQ ID NO: 50 are residues other than amino acids F, W, A, M, T, Y, N, T, G, M, L, S, D, I, M, Y, D, F and S, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 264 to 533 of SEQ ID NO: 52, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 269, 272, 293, 296, 297, 375, 401, 402, 403, 404, 407, 441, 445, 517, 520, 521, 525, 527 and 528 of SEQ ID NO: 52 are residues other than amino acids C, W, L, T, S, Y, S, A, G, Y, I, A, N, A, L, Y, D, Y and S, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 585 to 853 of SEQ ID NO: 56, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 590, 593, 614, 617, 618, 696, 721, 722, 723, 724, 727, 761, 765, 837, 840, 841, 845, 847 and 848 of SEQ ID NO: 56 are residues other than amino acids I, S, S, T, V, Y, S, I, A, L, V, G, N, M, F, Y, D, L and T, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 307 to 574 of SEQ ID NO: 54, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 312, 315, 336, 339, 340, 418, 443, 444, 445, 446, 449, 483, 487, 560, 563, 564, 566, 568 and 569 of SEQ ID NO: 54 are residues other than amino acids C, W, I, I, T, Y, S, I, S, A, I, L, D, A, I, Y, D, D and G, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 309 to 577 of SEQ ID NO: 24, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 314, 317, 338, 341, 342, 420, 446, 447, 448, 449, 452, 485, 489, 560, 563, 564, 569, 571 and 572 of SEO ID NO: 24 are residues other than amino acids

C, W, N, V, T, Y, I, G, G, I, L, L, D, A, I, Y, D, F and G, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 315 to 584 of SEQ ID NO: 26, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 320, 323, 344, 347, 348, 426, 451, 452, 453, 454, 457, 492, 496, 568, 571, 572, 576, 578 and 579 of SEQ ID NO: 26 are residues other than amino acids S, W, I, A, T, Y, S, V, A, S, I, L, D, A, I, Y, D, F, and G, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 265 to 536 of SEQ ID NO: 28, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 270, 273, 294, 297, 298, 376, 401, 402, 403, 404, 407, 440, 444, 518, 521, 522, 528, 530 and 531 of SEQ ID NO: 28 are residues other than amino acids A, W, V, C, G, F, T, S, C, I, M, G, N, C, S, Y, D, Y and S, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 342 to 612 of SEQ ID NO: 30, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 347, 350, 371, 374, 375, 453, 478, 479, 480, 481, 483, 518, 522, 596, 599, 600, 604, 606 and 607 of SEQ ID NO: 30 are residues other than amino acids F, L, C, V, T, Y, S, S, A, Y, V, L, G, L, L, Y, D, F and S, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more of the ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features an isolated synthase having a region with about 40% or greater sequence identity to residues 273 to 541 of SEQ ID NO: 32, wherein one or more amino acid residues of the synthase that align with amino acid residues at positions 278, 281, 302, 305, 306, 384, 409, 410, 411, 412, 415, 448, 452, 524, 527, 528, 533, 535 and 536 of SEQ ID NO: 32 are residues other than amino acids C, W, I, I, S, Y, T, S, T, Y, L, C, D, I, T, Y, D, Y and T, respectively. In some embodiments, the sequence identity can be about 20% or greater, 25% or greater, or 35% or greater. In some embodiments, one or more ordered arrangements of residues as given in Table 7 are not found in such a synthase.

The invention also features a method for making a terpene synthase, comprising identifying, in a preselected polypeptide having a region with 20% or greater sequence identity to residues 265 to 535 of SEQ ID NO: 2, nine amino acid residues whose  $\alpha$ -carbons have interatomic distances in Angstroms between the  $\alpha$ -carbons that are  $\pm 2.3$  Angstroms of the interatomic distances given in Table 6. The center point of each  $\alpha$ -carbon is positioned within a sphere having

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a radius of 2.3 Angstroms. The center point of each sphere has the structural coordinates given in Table 5. The method then comprises synthesizing a polypeptide that is modified from the preselected polypeptide. The modified polypeptide has one or more R-groups associated with the nine α-carbons other than the R-groups associated with the  $\alpha$ -carbons in the preselected polypeptide. The synthesizing step can comprise the formation of a nucleic acid encoding the preselected polypeptide in which the coding sequence for one or more amino acids corresponding to the nine α-carbons is replaced by a coding sequence that codes for an amino acid different from the amino acid present in the preselected polypeptide. The preselected polypeptide can be, for example, any one of the polypeptides given in SEQ ID NOS: 2, 4, 6, 8, 10, 12, 20, 22, 24, 26, 28, 30, 32, 34–40, 42, 44, 46, 48, 50, 52, 54, 56, or 58.

The invention also features a method of using a terpene synthase, comprising identifying, in a preselected polypeptide having a region with 20% or greater sequence identity to residues 265 to 535 of SEQ ID NO: 2, amino acid residues at nine positions that align with amino acid residues 273, 294, 402, 403, 404, 407, 440, 519 and 520 of SEQ ID NO: 2; and synthesizing a polypeptide that is modified from the preselected polypeptide. The novel polypeptide is modified by having amino acid residues at one or more of the nine positions other than the amino acid residues present in the preselected polypeptide. In some embodiments, the identifying step can comprise identifying sixteen amino acid residues in the preselected polypeptide that align with amino acid residues 270, 273, 294, 297, 298, 402, 403, 404, 407, 440, 516, 519, 520, 525, 527 and 528 of SEQ ID NO: 2, and the synthesizing step can comprise synthesizing a polypeptide that is modified from the preselected polypeptide, the modified polypeptide having amino acid residues at one or 35 more of the sixteen positions other than the amino acid residues present in the preselected polypeptide. In some embodiments, the identifying step can comprise identifying nineteen amino acid residues in the preselected polypeptide that align with amino acid residues 270, 273, 294, 297, 298, 376, 401, 402, 403, 404, 407, 440, 444, 516, 519, 520, 525, 527 and 528 of SEQ ID NO: 2, and the synthesizing step can comprise synthesizing a polypeptide that is modified from the preselected polypeptide, the modified polypeptide having amino acid residues at one or more of the nineteen 45 positions other than the amino acid residues present in the preselected polypeptide. The synthesizing step can comprise the formation of a nucleic acid encoding the preselected polypeptide in which the coding sequence in the nucleic acid coding for one or more of the identified amino acid residues is replaced by a coding sequence that encodes an amino acid different from the amino acid present in the preselected polypeptide. The preselected polypeptide can be, for example, any one of the polypeptides given in SEQ ID NOS: 2, 4, 6, 8, 10, 12, 20, 22, 24, 26, 28, 30, 32, 34–40, 42, 44, 46, 48, 50, 52, 54, 56, or 58. The method can further comprise: contacting the modified polypeptide with an isoprenoid substrate under conditions effective for the compound to bind the polypeptide; and measuring the ability of the modified polypeptide to catalyze the formation of a reaction product from the isoprenoid substrate. The isoprenoid substrate can be a monoterpene, a sesquiterpene, or a diterpene.

The invention also features a method of making a terpene synthase, comprising creating a population of nucleic acid molecules that encode polypeptides, the population having members that differ from one another at one or more of nine codons specifying amino acids of a preselected terpene

synthase having a region with about 20% or greater sequence identity to residues 265 to 535 of SEQ ID NO: 2, α-carbons of the nine amino acids having interatomic distances in Angstroms between the  $\alpha$ -carbons that are  $\pm 2.3$ Angstroms of the interatomic distances given in Table 6. The center point of each  $\alpha$ -carbon is positioned within a sphere having a radius of 2.3 Angstroms, and the center point of each sphere has the structural coordinates given in Table 5. In some embodiments, the codons specify amino acids as described in Tables 1-2 or 3-4 of a preselected terpene synthase. A portion, or all, of the nucleic acid population is expressed so that a population of polypeptides is made. At least one member of the population of polypeptides is a mutant terpene synthase. The expressing step can comprise in vitro transcription and in vitro translation of the nucleic acid population. In some embodiments, the expressing step comprises cloning members of the nucleic acid population into an expression vector, introducing the expression vector into host cells and expressing the cloned nucleic acid population members in the host cells so that the population 20 of polypeptides is made. The preselected terpene synthase polypeptide can be a monoterpene synthase, a sesquiterpene synthase, or a diterpene synthase. The host cells can be prokaryotic cells or eukaryotic cells, including, without limitation, bacterial cells, fungal cells, and animal cells, e.g., mammalian cells or insect cells. The host cells can also be plant cells, e.g., a cell from a Graminaceae plant, a cell from a Legumineae plant, a cell from a Solanaceae plant, a cell from a Brassicaeae plant or a cell from a Conifereae plant.

The invention also features a nucleic acid encoding a 30 synthase as described herein, and a host cell containing such a nucleic acid. The invention also features a transgenic plant containing such a nucleic acid, or a transgenic animal cell culture containing such a nucleic acid.

In some embodiments, a synthase polypeptide of the 35 invention comprises a domain that contains an active site comprised of nine α-carbon atoms having the coordinates of Table 5, and interatomic distances between the  $\alpha$ -carbons ±2.3 angstroms of the distances given in Table 5. The α-carbon atoms align structurally in three dimensional space 40 in the presence or absence of bound substrate or substrate analogue, with avian FPP synthase. In another embodiment, a synthase of this invention comprises the following: (i) a first domain containing amino acid residues that align in either having a bound or unbound substrate) with a glycosyl hydrolase catalytic core selected from the group consisting of (a) amino acids 36-230 of glycosyl hydrolase (PDB code 3GLY) of Aspergillus awarmori, and (b) amino acids 36-230 of endogluconase CellB (PDB code 1CLC), and (ii) 50 a second domain that aligns structurally in three dimensional space with or without substrate or substrate analogues bound in the active site with avian FPP synthase. The second domain contains an active site comprised of nine, sixteen or nineteen α-carbon atoms having the structural coordinates 55 and interatomic distances of Tables 1-2, 3-4 or 5-6. These α-carbon atoms have R-groups attached thereto that can interact, either directly or indirectly, with an isoprenoid substrate.

The invention also features a method for generating 60 mutant terpene synthases possessing catalytic activity. The method comprises the steps of (a) providing a crystallographic model of a preselected catalytically active terpene synthase having an active site, and (b) using the model to design a terpene synthase having at least one altered 65 R-group in the active site relative to the preselected synthase. The invention also features terpene synthases having

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altered substrate specificity, methods of making the same, and procedures for generating three-dimensional structures thereof.

Although methods and materials similar or equivalent to those described herein can be used to practice the invention, suitable methods and materials are described below. All publications, patent applications, patents and other references mentioned herein are incorporated by reference in their entirety.

Other aspects, embodiments, advantages, and features of the present invention will become apparent from the specification.

#### BRIEF DESCRIPTION OF DRAWINGS

FIG. 1. Schematic representation of tobacco 5-epiaristolochene synthase (TEAS) with bound farnesyl hydroxyphosphonate (FTP), prepared using the RIBBONS software program of Carson, M. and Bugg, C., J. Mol. Graphics 4:121 (1986). Cylinders 1-8 and A represent α-helices in the NH<sub>2</sub>-terminal domain; cylinders C, D, D1, D2, E, F, G1, G2, H1, H2, H3, I and  $\alpha$ -1 represent  $\alpha$ -helices in the COOH-terniinal domain.

FIG. 2. Structure of twenty natural amino acids showing α-carbons and associated R-groups.

FIG. 3. Autoradiogram of an argentation thin-layer chromatogram of terpenoid hydrocarbon products made by TEAS and mutant TEAS enzymes using GGPP as a substrate. DM: W273S/C440W mutant TEAS enzyme.

FIG. 4. Autoradiogram of an argentation thin-layer chromatogram of terpenoid hydrocarbon products made by TEAS and mutant TEAS enzymes using FPP as a substrate.

#### BRIEF DESCRIPTION OF TABLES

Table 1. X-ray crystallographic structural coordinates for 19  $\alpha$ -carbons found in the active site of a terpene synthase.

Table 2. Interatomic distances in Angstroms between each  $\alpha$ -carbon of Table 1. Each  $\alpha$ -carbon occupies a sphere having a radius of 2.3 Angstroms. Interatomic distances are calculated from the center point of each sphere.

Table 3. X-ray crystallographic structural coordinates for 16  $\alpha$ -carbons found in the active site of a terpene synthase.

Table 4. Interatomic distances in Angstroms between each three-dimensional space (in solution or crystal form, and 45 α-carbon of Table 3. Each α-carbon occupies a sphere having a radius of 2.3 Angstroms. Interatomic distances are calculated from the center point of each sphere.

> Table 5. X-ray crystallographic structural coordinates for nine  $\alpha$ -carbons found in the active site of a terpene synthase.

> Table 6. Interatomic distances in Angstroms between each α-carbon of Table 5. Each α-carbon occupies a sphere having a radius of 2.3 Angstroms. Interatomic distances are calculated from the center point of each sphere.

Table 7. Ordered arrangement of Regroups not found associated with the  $\alpha$ -carbons of Table 1.

Table 8. Ordered arrangement of R-groups not found associated with the  $\alpha$ -carbons of Table 3.

Table 9. Ordered arrangement of R-groups not found associated with the  $\alpha$ -carbons of Table 5.

Table 10. X-ray structural coordinates for TEAS having the substrate analog FHP bound in the active site.

Table 11. X-ray structural coordinates for TEAS in the absence of substrate.

Table 12. Alignment of residues 265–535 of TEAS with a limonene synthase, SEQ ID NO: 22, using the BLASTp alignment program.

Table 13. Alignment of residues 579 to 847 of SEQ ID NO:44 with SEQ ID NO:26, using the BLASTp program.

Table 14. Alignment of residues 265 to 535 of TEAS with SEQ ID NO:48, using the BLASTp program.

Table 15. Alignment of residues 307 to 593 of SEQ ID NO:50 with SEQ ID NO:56 using the BLASTp program.

## BRIEF DESCRIPTION OF THE SEQUENCE LISTING

SEQ ID NO:1 is the DNA coding sequence for a tobacco 5-epi-aristolochene synthase (TEAS) protein. Genbank No: O40577.

SEQ ID NO:2 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:1.

SEQ ID NO:3 is the DNA coding sequence for a TEAS protein in which the codon for Trp273 has been changed to a codon for Glu.

SEQ ID NO:4 is the amino acid sequence for the W273E protein encoded by the TEAS DNA of SEQ ID NO:3.

SEQ ID NO:5 is the DNA coding sequence for a TEAS protein in which the codon for Tyr520 has been changed to a codon for Phe.

SEQ ID NO:6 is the amino acid sequence for the Y520F 25 exons encoded by the DNA of SEQ ID NO:33. protein encoded by the TEAS DNA of SEQ ID NO:5. SEO ID NO:41 is the DNA coding sequence at the DNA coding sequen

SEQ ID NO:7 is the DNA coding sequence for a TEAS protein in which the codon for Tyr527 has been changed to a codon for Phe.

SEQ ID NO:8 is the amino acid sequence for the Y527F <sup>30</sup> protein encoded by the TEAS DNA of SEQ ID NO:7.

SEQ ID NO:9 is the DNA coding sequence for a TEAS protein in which the codon for Trp273 has been changed to a codon for Ser and the codon for Cys440 has been changed to a codon for Trp.

SEQ ID NO:10 is the amino acid sequence for the W273S/C440W protein encoded by the TEAS DNA of SEQ ID NO:9.

SEQ ID NO:11 is the DNA coding sequence for TEAS  $_{\rm 40}$  proteins in which the codons for Tyr406 and Leu407 have each been changed to the nucleotides NNS.

SEQ ID NO:12 is the amino acid sequence for the population of Y406X/L407X proteins encoded by the TEAS DNA of SEQ ID NO:11, where X is any naturally occurring 45 amino acid.

SEQ ID NO:13 is a DNA primer sequence.

SEQ ID NO:14 is a DNA primer sequence.

SEQ ID NO:15 is a DNA primer sequence.

SEQ ID NO:16 is a DNA primer sequence.

SEQ ID NO:17 is a DNA primer sequence. SEQ ID NO:18 is a DNA primer sequence.

SEQ ID NO:19 is the DNA coding sequence for a grand fir pinene synthase. Genbank Accession No: U87909.

SEQ ID NO:20 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:19.

SEQ ID NO:21 is the DNA coding sequence for a spearmint limonene synthase. Genbank Accession No: L13459.

SEQ ID NO:22 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:21.

SEQ ID NO:23 is the DNA coding sequence for a sage 1, 8 cincole synthase. Genbank Accession No: AF051899.

SEQ ID NO:24 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:23.

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SEQ ID NO:25 is the DNA coding sequence for a sage bornyl diphosphate synthase. Genbank Accession No: AF051900.

SEQ ID NO:26 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:25.

SEQ ID NO:27 is the DNA coding sequence for a mint E-b-farnesene synthase. Genbank Accession No: AF024615.

SEQ ID NO:28 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:27.

SEQ ID NO:29 is the DNA coding sequence for a grand fir myrcene synthase. Genbank Accession No: U87908.

SEQ ID NO:30 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:29.

SEQ ID NO:31 is the DNA coding sequence for a potato vetaspiradiene synthase. Genbank Accession No: AB022598.

SEQ ID NO:32 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:31.

SEQ ID NO:33 is the genomic DNA coding sequence for a cotton delta-cadinene synthase. Genbank Accession No: Y18484.

SEQ ID NOS:34-40 are the amino acid sequences for the exons encoded by the DNA of SEO ID NO:33.

SEQ ID NO:41 is the DNA coding sequence for a castor bean casbene synthase. Genbank Accession No: L32134.

SEQ ID NO:42 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:41.

SEQ ID NO:43 is the DNA coding sequence for a yew taxadiene synthase. Genbank Accession No: U48796.

SEQ ID NO:44 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:43.

SEQ ID NO:45 is the DNA coding sequence for a grand fir E-alpha-bisabolene synthase. Genbank Accession No: AF006194.

SEQ ID NO:46 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:45.

SEQ ID NO:47 is the DNA coding sequence for a grand fir delta-selinene synthase. Genbank Accession No: U92266.

SEQ ID NO:48 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:47.

SEQ ID NO:49 is the DNA coding sequence for a grand fir gamma-humulene synthase. Genbank Accession No: U92267.

SEQ ID NO:50 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:49.

SEQ ID NO:51 is the DNA coding sequence for a tomato germacrene C synthase. Genbank Accession No: AF035631.

SEQ ID NO:52 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:51.

SEQ ID NO:53 is the DNA coding sequence for a sage+sabinene synthase. Genbank Accession No: AF051901.

SEQ ID NO:54 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:53.

SEQ ID NO:55 is the DNA coding sequence for a grand fir abietadiene synthase. Genbank Accession No: U50768.

SEQ ID NO:56 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:55.

SEQ ID NO:57 is the DNA coding sequence for a grand  $_{65}\,$  fir limonene synthase. Genbank Accession No. AF006193.

SEQ ID NO:58 is the amino acid sequence for the protein encoded by the DNA of SEQ ID NO:57.

#### DETAILED DESCRIPTION

The following terms are used herein:

" $\alpha$ -carbon" refers to the chiral carbon atom found in an amino acid residue. Four substituents are covalently bound to the  $\alpha$ -carbon, including an amino group, a carboxyl group, a hydrogen atom, and an R-group.

"R-group" refers to a substituent attached to the  $\alpha$ -carbon of an amino acid residue that is not involved in peptide bond formation in a protein. An R-group is an important determinant of the overall chemical character of an amino acid. The twenty naturally occurring amino acids found in proteins and the R-groups associated with the  $\alpha$ -carbon of each amino acid are listed in FIG. 2. The three-letter and one-letter abbreviations for naturally occurring amino acids are 15 sometimes used herein to refer to the R-group associated with a particular amino acid.

"Naturally occurring amino acid" includes L-isomers of the twenty amino acids naturally occurring in proteins. Naturally occurring amino acids are glycine, alanine, valine, 20 leucine, isoleucine, serine, methionine, threonine, phenylalanine, tyrosine, tryptophan, cysteine, proline, histidine, aspartic acid, asparagine, glutamic acid, glutamine, arginine, and lysine. Unless specially indicated, all amino acids referred to in this application are in the 25 L-form. Three-letter and one-letter abbreviations are sometimes used herein to refer to naturally occurring amino acids. These abbreviations are known in the art.

"Unnatural amino acid" includes amino acids that are not naturally found in proteins. Examples of unnatural amino 30 acids included herein are racemic mixtures of selenocysteine and selenomethionine. In addition, unnatural amino acids include the D or L forms of norleucine, paranitrophenylalanine, homophenylalanine, parafluorophenylalanine, 3-amino-2-benzylpropionic acid, 35 homoarginine, D-phenylalanine, and the like.

"Positively charged amino acid" includes any naturally occurring or unnatural amino acid having an R-group that carries a positive charge under normal physiological conditions. Examples of positively charged, naturally occurring amino acids include arginine and lysine.

"Negatively charged amino acid" includes any naturally occurring or unnatural amino acid having an R-group that carries a negative charge under normal physiological conditions. Examples of negatively charged, naturally occurring amino acids include aspartic acid and glutamic acid.

"Hydrophobic amino acid" includes any naturally occurring or unnatural amino acid having an uncharged, nonpolar side chain under normal physiological conditions. Examples of naturally occurring hydrophobic amino acids are leucine, isoleucine, valine and methionine.

"Hydrophilic amino acid" includes any naturally occurring or unnatural amino acid having a charged polar side chain. Examples of naturally occurring hydrophilic amino 55 acids include serine, threonine and cysteine.

"Mutant terpene synthase" or "mutated terpene synthase" refers to a synthase polypeptide having a primary amino acid sequence. The center point of the  $\alpha$ -carbon of nine residues of the polypeptide is positioned within a sphere having a radius of 2.3 Angstroms; the center points of the nine spheres have the structural coordinates of Table 5 or coordinates which can be rotated and/or translated to coincide with the coordinates of Table 5. The relative interatomic distances between the nine  $\alpha$ -carbons is  $\pm 2.3$  angstroms of the nine amino code. The interatomic distances given in Table 6. Each  $\alpha$ -carbon has an associated R-group. A mutant synthase differs from a acid reference with the coordinates of Table 5.

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non-mutant synthase in the ordered arrangement of R-groups associated with the nine  $\alpha$ -carbons. A mutant synthase has an ordered arrangement of R-groups on the nine  $\alpha$ -carbons other than the ordered arrangements of R-groups listed in Table 9. R-groups associated with other  $\alpha$ -carbons of the synthase primary amino acid sequence may or may not be the same as in a non-mutated synthase.

In some embodiments, a mutant synthase refers to a synthase in which the center point of the  $\alpha$ -carbon of sixteen residues of the polypeptide is positioned within a sphere having a radius of 2.3 Angstroms; the center points of the sixteen spheres have the structural coordinates of Table 3 or coordinates which can be rotated and/or translated to coincide with the coordinates of Table 3. The relative interatomic distances between the nine  $\alpha$ -carbons is  $\pm 2.3$  angstroms of the interatomic distances given in Table 4. Each  $\alpha$ -carbon has an associated R-group. A mutant synthase differs from a non-mutant synthase in the ordered arrangement of R-groups associated with the sixteen  $\alpha$ -carbons. A mutant synthase has an ordered arrangement of R-groups on the sixteen \alpha-carbons other than the ordered arrangements of R-groups listed in Table 8. R-groups associated with other α-carbons of the synthase primary amino acid sequence may or may not be the same as in a non-mutated synthase.

In some embodiments, a mutant synthase refers to a synthase in which the center point of the  $\alpha$ -carbon of nineteen residues of the polypeptide is positioned within a sphere having a radius of 2.3 Angstroms; the center points of the nineteen spheres have the three dimensional coordinates of Table 1 or coordinates which can be rotated and/or translated to coincide with the coordinates of Table 1. The relative interatomic distances between the nineteen  $\alpha$ -carbons is  $\pm 2.3$  angstroms of the interatomic distances given in Table 2. Each α-carbon has an associated R-group. A mutant synthase differs from a non-mutant synthase in the ordered arrangement of R-groups associated with the nineteen α-carbons. A mutant synthase has an ordered arrangement of R-groups on the nineteen α-carbons other than the ordered arrangements of R-groups listed in Table 7. R-groups associated with other  $\alpha$ -carbons of the synthase primary amino acid sequence may or may not be the same as in a non-mutated synthase.

"Nonmutated synthase" or "non-mutant synthase" includes a synthase having a primary amino acid sequence comprising nine, sixteen, or nineteen amino acid residues.

The center point of each α-carbon of these residues is positioned within a sphere having a radius of 2.3 Angstroms; the center points of the spheres have the three dimensional coordinates of Tables 5, 3, or 1, respectively, or coordinates which can be rotated and/or translated to coincide with the coordinates of Tables 5, 3, or 1. The relative interatomic distances between the nine, sixteen, or nineteen α-carbons is ±2.3 angstroms of the interatomic distances given in Tables 6, 4, or 2, respectively. Each α-carbon has an associated R-group. A non-mutant synthase has an ordered arrangement of R-groups on the nine, sixteen, or nineteen α-carbons as listed in Tables 9, 8, or 7, respectively.

"Degenerate variations thereof" refers to variants of a gene coding sequence by which the same polypeptide is encoded by different nucleotide sequences, due to the degeneracy of the genetic code. For example, synthases of the present invention have a primary amino acid sequence. Degenerate synthase variations are different nucleic acid coding sequences that nevertheless encode the same primary amino acid sequence due to the degeneracy of the genetic code.

"Expression" refers to transcription of a gene or nucleic acid molecule and the translation of that nucleic acid into a

polypeptide. Expression of genes also involves processing of RNA into mRNA in eukaryotic systems. It is not necessary for the genes to integrate into the genome of a cell in order to achieve expression. This definition is not limited to expression in a particular system or a particular cell type and includes, without limitation, stable, transient, in vitro, and in vivo expression.

"Promoter" and "promoter regulatory element", refers to a nucleic acid that is involved in controlling expression of a gene. Promoter regulatory elements, and the like, from a <sup>10</sup> variety of sources can be used efficiently to promote gene expression. Promoter regulatory elements include constitutive, tissue-specific, developmental-specific, inducible, subgenomic promoters, and the like. Promoter regulatory elements may also include certain enhancer ele- <sup>15</sup> ments or silencing elements that improve or regulate transcriptional efficiency.

"Active Site" refers to a site in a terpene synthase that binds the hydrophobic portion of a terpene substrate, GPP, FPP, and/or GGPP. The active site can, under certain conditions, catalyze a biosynthetic reaction that allows one or more reaction products to be produced.

"Altered enzymatic specificity" includes an alteration in the ability of a mutant synthase to use a particular terpene substrate or a change in the profile of reaction product(s) from a mutant synthase, compared to the substrate specificity of and the reaction products made by a corresponding non-mutated synthase. Altered specificity may include the ability of a synthase to exhibit different enzymatic parameters relative to a non-mutated synthase (Km, Vmax, etc), and/or to produce products that are different from those that are produced by a corresponding non-mutant synthase.

"Structure coordinates" or "structural coordinates" refers to Cartesian coordinates (x, y, and z positions) derived from mathematical equations involving Fourier synthesis as determined from patterns obtained via diffraction of a monochromatic beam of X-rays by the atoms (scattering centers) of a synthase molecule in crystal form. Diffraction data are used to calculate electron density maps of repeating protein 40 lated gene. units in the crystal (unit cell). Electron density maps are used to establish the positions of individual atoms within a crystal's unit cell. The absolute values for structural coordinates listed herein convey relative spatial relationships between atoms because the absolute values ascribed to 45 structural coordinates can be changed by rotational and/or translational movement along the x, y and/or z axes, together or separately, while maintaining the same relative spatial relationships among atoms. Thus, a terpene synthase whose absolute values for a set of structural coordinates can be 50 rotationally or translationally adjusted to coincide with the particular values listed in Tables 1, 3, or 5 is considered to have the same structural coordinates as those of Tables 1, 3 or 5. An example of structural coordinates that coincide with the absolute values listed herein after rotation and/or trans- 55 lation are the coordinates of Table 11.

"Heavy atom derivatization" refers to a method of producing a chemically modified form of a synthase crystal. In practice, a crystal is soaked in a solution containing heavy atom salts or organometallic compounds, e. g., lead chloride, 60 gold thiomalate, thimerosal, uranyl acetate and the like, which can diffuse through the crystal and bind to the protein's surface. Locations of the bound heavy atoms can be determined by X-ray diffraction analysis of the soaked crystal. The information is then used to construct phase 65 information which can then be used to construct three-dimensional structures of the enzyme as described in

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Blundel, T. L., and Johnson, N. L., Protein Crystallography, Academic Press (1976).

"Unit cell" refers to a basic parallelepiped shaped block. Regular assembly of such blocks may construct the entire volume of a crystal. Each unit cell comprises a complete representation of the unit pattern, the repetition of which builds up the crystal.

"Mutagenesis" refers to the substitution of a different amino acid residue at a particular position in the primary amino acid sequence of a protein, thereby changing the R-group present at that position. Mutagenesis can be most easily performed by changing the coding sequence of a nucleic acid encoding the protein so that the coding sequence in the nucleic acid specifies an amino acid residue different from the residue initially present at that position.

"Space Group" refers to the arrangement of symmetry elements within a crystal.

"Molecular replacement" refers to the generation of a 20 preliminary model of a synthase whose structural coordinates are unknown, by orienting and positioning a molecule whose structural coordinates are known within the unit cell of the unknown crystal so as best to account for the observed diffraction pattern of the unknown crystal. Phases can then be calculated from this model and combined with the observed amplitudes to give an approximate Fourier synthesis of the structure whose coordinates are unknown. This in turn can be subject to any of the several forms of refinement to provide a final, accurate structure of the unknown crystal (Lattman, E., 1985, in Methods in Enzymology, 115:55-77; Rossmann, M G., ed., "The Molecular Replacement Method" 1972, Int. Sci. Rev. Ser., No. 13, Gordon & Breach, New York). Using structure coordinates and interatomic distance matrices, molecular 35 replacement may be used to determine the structural coordinates of a crystalline mutant, homologue, or a different crystal form of terpene synthase.

"Recombinant protein" includes a protein that is chemically synthesized or derived biosynthetically from an isolated gene.

"Gene" includes naturally derived or genetically manipulated nucleic acids that contain the information needed to produce a polypeptide.

"Nucleic acid" includes any genetic material comprised of the nucleotides guanine, adenine, thymine, cytosine, uracil, inosine and the like. Nucleic acids may be single-, double-, or triple stranded. Nucleic acids may be deoxyribonucleic acid or ribonucleic acid.

"Genetically manipulated" includes genes that have been modified to contain a different nucleotide sequence from that present in a preselected nulceic acid. Genes can be manipulated by synthetically or via traditional cloning, PCR, chemical gene synthesis, direct or random mutagenesis, and gene shuffling. Genetically manipulated also includes the process of making genes that are degenerate variations of nucleic acids encoding preselected proteins.

"First domain" includes polypeptides having a first and second end wherein the first end can have an amino terminal amino acid with a free amino group and can be linked by a peptide bond to a second amino acid. The first end may also be modified through acetylation and the like. The second end of the first domain may or may not have a free carboxyl terminal group.

"Second domain" includes polypeptides having a first and second end wherein the first end can have an amino terminal amino acid and can be linked by a peptide bond to a second

amino acid. The second end of the second domain may or may not have a carboxyl terminal group. Typically, the first end of the second domain is linked to the second end of the first domain via a peptide bond.

"Isoprenoid substrate" refers to the  $C_{10}$ ,  $C_{15}$ , and  $C_{20}$  molecules, named geranyl diphosphate (GPP), farnesyl diphosphate (FPP), and geranylgeranyl diphosphate (GGPP), respectively.

"Sequence identity" or "percent sequence identity" refers to the percentage of amino acids or nucleotides that occupy the same relative position when two protein sequences or nucleic acid sequences, a query sequence and a subject sequence, are aligned. The number of amino acid or nucleotide residues that are identical between both the subject and query sequences are counted, divided by the number of residues in the query sequence, and multiplied by 100. The process is repeated until the alignment resulting in the highest percent sequence identity is found. Percent sequence identity can be determined by visual inspection and/or by using various computer programs, e.g., MegAlign (DNASTAR, Inc., Madison, Wis.) or BLAST programs available on the world wide web from the National Center for Biotechnology Information (NCBI). Gaps of one or more residues may sometimes be inserted to maximize sequence alignments to structurally conserved domains of the query sequence, i.e.,  $\alpha$ -helices,  $\beta$ -sheets and loops.

"Monoterpene product" refers to linear, cyclized, and/or hydroxylated reaction products made from the substrate GPP. "Sesquiterpene product" refers to linear, cyclized, and/or hydroxylated reaction products made from the substrate FPP. "Diterpene product" refers to linear, cyclized, and/or hydroxylated reaction products made from the substrate GGPP.

The present invention relates to terpene synthases and mutants thereof from which the position of specific α-carbon atoms and R-groups associated therewith comprising the active site can be determined in three-dimensional space. The invention also relates to structural coordinates of the synthases, use of the structural coordinates to develop structural information related to synthase homologues, mutants, and the like, and to crystal forms of such synthases. Furthermore, the invention provides a method whereby α-carbon structural coordinates for atoms comprising the active site of a preselected terpene synthase can be used to develop synthases in which R-groups associated with active site α-carbon atoms are different from the R-groups found in the preselected terpene synthase. In addition, the present invention provides for the production of novel terpene synthases based on the structural information provided herein and for the use of such synthases to make a variety of isoprenoid compounds.

The present invention further provides, for the first time, crystals of a synthase, as exemplified by tobacco 5epi-aristolochene synthase (TEAS), which are grown in the 55 presence or absence of substrate and substrate analogues, thus allowing definition of the structural coordinates associated therewith. The structural coordinates allow determination of the  $\alpha$ -carbon atoms comprising the active site and R-groups associated therewith. The crystals of the present invention belong to the tetragonal space group P4<sub>1</sub>2<sub>1</sub>2; the unit cell dimensions vary by a few angstroms between crystals but on average a=126 angstroms, c=122 angstroms, a=b,  $\alpha$ =90°,  $\beta$ =90°, and  $\gamma$ =90°.

Structural coordinates are preferably obtained at a reso- 65 lution of about 2.2 to about 2.8 angstroms for a synthase in the presence and in the absence of bound substrate or

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substrate analog. Coordinates for a synthase with a substrate analog bound in the active site are given in Table 10. Coordinates for a synthase in the absence of a substrate analog bound in the active site are given in Table 11. Those skilled in the art understand that a set of structure coordinates determined by X-ray crystallography is not without standard error. Therefore, for the purpose of this invention, any set of structure coordinates wherein the active site  $\alpha$ -carbons of a synthase, synthase homologue, or mutants thereof, have a root mean square deviation less than  $\pm 2.3$  angstroms when superimposed using the structural coordinates listed in Table 1, 3, or 5, are considered identical.

A schematic representation of the three-dimensional shape of a synthase is shown in FIG. 1 which was prepared by RIBBONS (Carson and Bugg, 1986, J. Mol. Graphics, 4:121). The synthase shown in FIG. 1 consists entirely of  $\alpha$ -helices and short connecting loops and turns, organized into first and second structural domains.

In one embodiment, an isolated synthase of the invention  $_{20}\,$  comprises sixteen active site  $\alpha$ -carbons having the structural coordinates of Table 3 and the relative distances ±2.3 angstroms of the distances given in Table 4. The active site α-carbons of Table 3 generally are not all contiguous, i.e., are not adjacent to one another in the primary amino acid sequence of a synthase, due to intervening amino acid residues between various active site  $\alpha$ -carbons. On the other hand, it should be appreciated that some of the active site  $\alpha$ -carbons can be adjacent to one another in some instances. In the embodiment depicted in the TEAS Y527F protein (SEQ ID NO:8), for example, active site  $\alpha$ -carbons are adjacent to one another in the primary amino acid sequence at positions 402, 403 and 404, respectively, whereas active site  $\alpha$ -carbons at residues 273 and 294 are separated and thus are not adjacent. Thus, the numbering of active site  $\alpha$ -carbons given in Tables 1, 2, 3, 4, 5, or 6 is merely for convenience and such  $\alpha$ -carbons may reside at any position in the primary amino acid sequence that achieves the structural coordinates given in Tables 1, 3, or 5 and the relative interatomic distances ±2.3 angstroms given in Tables 2, 4, or

An appropriate combination of R-groups, linked to active site \alpha-carbons, can facilitate the formation of one or more desired reaction products. The combination of R-groups selected for use in a terpene synthase of the invention can be 45 any combination other than the ordered arrangements of R-groups and corresponding active site  $\alpha$ -carbons shown in Tables 7, 8, or 9. An illustrative example of a suitable arrangement of R-groups and  $\alpha$ -carbons is Cys, Trp, Ile, Ile, Ser, Thr, Thr, Tyr, Leu, Cys, Val, Thr, Phe, Asp, Tyr and Thr, associated with active site  $\alpha$ -carbons 1 to 16, respectively, of Table 3. Another example of a suitable arrangement of R-groups and  $\alpha$ -carbons is Cys, Trp, Ile, Ile, Ser, Thr, Thr, Tyr, Leu, Cys, Val, Thr, Tyr, Asp, Phe, and Thr at active site alpha-carbons 1 to 16, respectively, of Table 3. In some embodiments, a synthase of the invention may have primary amino acid sequences as listed in SEO ID NO:4, SEO ID NO:6, SEQ ID NO:8, and SEQ ID NO:10, DNA molecules encoding the same, which are listed in SEQ ID NO:3, SEQ ID NO:5, SEQ ID NO:7, and SEQ ID NO:9, respectively, and degenerate variations thereof. Typically, R-groups found on active site  $\alpha$ -carbons are those found in naturally occurring amino acids. See, e.g., FIG. 2. In some embodiments, however, R-groups other than naturally occurring amino acids can be used.

Some arrangements of R-groups and active site  $\alpha$ -carbons result in mutant terpene synthases that form reaction products. Such enzymatically active synthases and their corre-

sponding genes are useful to make known terpenoid hydrocarbons, e.g., monoterpenes such as pinene, sesquiterpenes such as delta-cadinene and diterpenes such as abietadiene. Other enzymatically active synthases can be used to make novel terpenoid products.

Some arrangements of R-groups and active site α-carbons may result in mutant terpene synthases that do not form reaction product(s) at a desired rate. Such synthases and their genes are useful as controls in analyses of product formation by enzymatically active mutant synthases. Such synthases and their genes can also be useful in analyses of translation of enzymatically active mutant synthase genes, or as nutritional supplements. Such synthases can be attached to Sepharose beads and used for affinity purification of isoprenoid compounds from crude preparations. In addition, such synthases and their genes can also be useful to develop reagents for various purposes, e.g., immunological reagents to monitor expression of a terpene synthase protein or nucleic acid probes or primers to monitor inheritance of a terpene synthase gene in a plant breeding

In some embodiments, the  $\alpha$ -carbon backbone of a synthase first domain aligns structurally with the catalytic core of glycosyl hydrolases, as exemplified by glucoamylase (Brookhaven Protein Database (PDB) code 3GLY) from Aspergillus awamori (Aleshin et al., 1994, J. Mol. Biol., 25 238:575) and endoglucanase CelD (PDB code ICLC) from Clostridium thermocelum (Juy et al., 1992, Nature, 357:89), and the  $\alpha$ -carbon backbone of a synthase second domain, which contains the active site, aligns structurally with avian farnesyl diphosphate synthase (FPS), wherein the active site is comprised of 9, 16, or 19 amino acid residues with α-carbon structural coordinates as listed in Tables 1, 3, or 5 and interatomic distances as described in Tables 2, 4, or 6. Such α-carbons have an ordered arrangement of R-groups different from that observed in a non-mutated synthase.

In the present invention, the first domain forms a twisted α-barrel made up of eight short (10 to 15 amino acid residues) helices surrounding a surface cavity filled by ordered water molecules when hydrated. The second domain comprises a two-layered barrel of  $\alpha$ -helices surrounding a  $_{40}$ hydrophobic and aromatic-rich active site pocket. Typically, the second domain contains a substrate binding site. As exemplified in FIG. 1, helix H is disrupted between segments H1 and H2 by an amino acid such as proline, but its corresponding kink in helix G between G1 and G2. Within this kink, hydrogen bonds between a hydroxyl group, such as that found on a threonine, and the carbonyl oxygen of other amino acids disrupt the main chain intrahelical hydrostructure as determined.

As exemplified by TEAS, terpene synthases of the present invention can have a first domain segment comprising helices A and C (an A-C loop), and a second domain comprising helices I and K (a I-K loop) (FIG. 1). The 55 ordering of these loops upon substrate binding results in a closed, solvent-inaccessible active site pocket As the I-K loop becomes ordered, a lid-type structure is formed that clamps down over the active site entrance in the presence of substrate and an extended aromatic patch deep within the active site pocket is formed. As the A-C loop becomes ordered, it translates inward toward the active site, positioning certain R groups in this loop at or near the active site. Thus, substrate binding to the active site results in a change in protein conformation.

To identify or create mutant terpene synthases, sequence alignments can be performed to locate specific residues and 20

α-carbons in a preselected polypeptide that have the structural coordinates and interatomic distances of Tables 1-2, 3-4 or 5-6. The preselected polypeptide is used as the subject sequence in the alignment, e.g., the full-length primary amino acid sequence, a region 190 residues in length, a region 220 residues in length, or a region 300 residues in length. The alignment can use residues 265 to 535 of TEAS (SEQ ID NO: 2), which includes the  $\alpha$ -carbons of Tables 1, 3 or 5, as the query sequence to align with the 10 preselected polypeptide. The preselected polypeptide and the query sequence can be aligned using the BLASTp 2.0.9 computer program with a BLOSUM 62 scoring matrix, an expect value of 10, a gap open value of 11, an x\_dropoff value of 50, a gap extension value of 1, a wordsize of 3 and no filtering of low complexity sequences. As an alternative, the BLASTp 2.0.9 program can be used with a BLOSUM 50 scoring matrix, an expect value of 10, a gap open value 13, an x\_dropoff value of 50, a gap extension value of 2, a wordsize of 3 and no filtering of low complexity sequences. Other parameter values can also be used, e.g., a gap extension value from 0 to 4. See Altschul, et al., Nucl. Acids Res. 25:3389-3402.

Regions of the preselected polypeptide with significant sequence identity to residues 265–535 of TEAS, e.g., 20% or greater sequence identity, 25% or greater sequence identity, 35% or greater sequence identity, 40% or greater sequence identity, 50% or greater sequence identity, 60% or greater sequence identity, 70% or greater sequence identity, or 80% or greater sequence identity are examined for specific residues that align with the TEAS residues corresponding to those listed in Tables 1, 3, or 5. In some cases, the output of the computer program alignment identifies a specific residue in the preselected polypeptide for each of the nine, sixteen, or nineteen residues in the query sequence 35 having the structural coordinates and interatomic distances of Tables 1–2, 3–4 or 5–6, with or without gaps introduced by the alignment program. In other cases, a gap is introduced by the alignment program in either the query sequence or the subject sequence such that no direct alignment or a misalignment occurs between one or more of the nine, sixteen, or nineteen residues in the query sequence that are of interest. In either case, the output can be visually inspected, and specific residues can be chosen in the subject sequence after adjusting the alignment so that alpha-helices and interhelical packing with helix G is accommodated by a 45 beta-sheet regions in the query sequence are maintained and that gaps or insertions in the subject sequence align with loop regions of the query sentence.

Sequence alignments suggest that other terpene synthases have regions with 20% or greater sequence identity to gen bonding of helix G thus assisting in producing the 50 residues 265-535 of TEAS. Therefore, a region of a terpene synthase other than TEAS can be used as the query sequence, e.g., regions of terpene synthases given in SEQ ID NOS: 4, 6, 8, 10, 12, 20, 22, 24, 26, 28, 30, 32, 34–40, 42, 44, 46, 48, 50, 52, 54, 56, or 58, that have significant sequence identity to residues 265–535 of SEQ ID NO: 2. For example, large sequence insertions are present at the amino terminus in taxadiene synthase (SEQ ID NO: 44) with respect to TEAS, or are within solvent-exposed loops in the amino-terminal domain. Thus, regions of taxadiene synthase with greater than 20% sequence identity to SEQ ID NO: 2 are closer to the carboxy-terminal end, e.g., from residue 579 to residue 847 of SEO ID NO: 44.

> Useful regions of other terpene synthases that can be used as the query sequence include, without limitation, residues 343 to 606 of SEQ ID NO: 20, 316 to 586 of SEQ ID NO: 22, residues 352 to 622 of SEQ ID NO: 58, residues 272 to 540 encoded by SEQ ID NO: 33, residues 319 to 571 of SEQ

ID NO: 42, residues 579 to 847 of SEQ ID NO: 44, residues 495 to 767 of SEQ ID NO: 46, residues 295 to 564 of SEQ ID NO: 48, residues 307 to 578 of SEQ ID NO: 50, residues 264 to 533 of SEQ ID NO: 52, residues 585 to 853 of SEQ ID NO: 56, residues 307 to 574 of SEQ ID NO: 54, residues 309 to 577 of SEQ ID NO: 24, residues 315 to 584 of SEQ ID NO: 26, residues 265 to 536 of SEQ ID NO: 28, residues 342 to 612 of SEQ ID NO: 30 and residues 273 to 541 of SEQ ID NO: 32.

One or more of the specific residues in the subject sequence that align with residues in the query sequence are mutated in the preselected polypeptide, e.g, by making mutations in a nucleic acid encoding the polypeptide. The mutant terpene synthase thus created can then be expressed in a host cell and the protein evaluated for enzymatic activity, if desired.

Mutant proteins of the present invention may be prepared in a number of ways including but not limited to oligonucleotide-directed mutagenesis, deletion, chemical mutagenesis, and the like. One or more R-groups associated 20 with the active site  $\alpha$ -carbon atoms in a terpene synthase are changed by altering the nucleotide sequence of the corresponding gene. For example, a mutation can be introduced into SEQ ID NO:1, the nucleotide sequence for TEAS, at codons encoding one or more of the following sixteen 25 α-carbons: α-carbon 1=Cys 270; α-carbon 2=Trp 273; α-carbon 3=Ile 294; α-carbon 4=Ile 297; α-carbon 5=Ser298;  $\alpha$ -carbon 6=Thr 402;  $\alpha$ -carbon 7=Thr 403; α-carbon 8=Tyr 404; α-carbon 9=Leu 407; α-carbon 10=Cys 440; α-carbon 11=Val 516; α-carbon 12=Thr 519; α-carbon 13=Tyr 520; α-carbon 14=Asp 525; α-carbon 15=Tyr 527; or α-carbon 16=Thr 528. The protein encoded by the mutant gene is then produced by expressing the gene in, for example, a bacterial or plant expression system. Alternatively, synthase mutants may be generated by site specific replacement of a particular amino acid with an unnaturally occurring amino acid. As such, synthase mutants may be generated through replacement of an amino acid residue or a particular cysteine or methionine residue with selenocysteine or selenomethionine. This may be achieved by growing a host organism capable of expressing either the wild-type or mutant polypeptide on a growth medium depleted of natural cysteine or methionine or both and growing on medium enriched with either selenocysteine, described in Sambrook et al., (Molecular Cloning, A Laboratory Manual, 2<sup>nd</sup> Ed. (1989) Cold Spring Harbor Laboratory Press).

Another suitable method of creating mutant synthases of the present invention is based on a procedure described in 50 Noel and Tsal (1989) J. Cell. Biochem., 40:309–320. In so doing, the nucleic acid encoding the synthase can be synthetically produced using oligonucleotides having overlapping regions, the oligonucleotides being degenerate at specific bases so that mutations are induced.

According to the present invention, nucleic acid sequences encoding a mutated synthase can be produced by the methods described herein, or any alternative methods available to the skilled artisan. In designing the nucleic acid sequence (gene) of interest, it may be desirable to reengineer the gene for improved expression in a particular expression system. For example, it has been shown that many bacterially derived genes do not express well in plant systems. In some cases, plant-derived genes do not express well in bacteria. This phenomenon may be due to the non-optimal G+C content or A+T content of the gene relative to the expression system being used. For example, the very low

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G+C content of many bacterial genes results in the generation of sequences mimicking or duplicating plant gene control sequences that are highly A+T rich. The presence of A+T rich sequences within the genes introduced into plants (e.g., TATA box regions normally found in gene promoters) may result in aberrant transcription of the gene(s). In addition, the presence of other regulatory sequences residing in the transcribed mRNA (e.g. polyadenylation signal sequences (AAUAAA) or sequences complementary to small nuclear RNAs involved in pre-mRNA splicing) may lead to RNA instability. Therefore, one goal in the design of genes is to generate nucleic acid sequences that have a G+C content that affords mRNA stability and translation accuracy for a particular expression system.

Due to the plasticity afforded by the redundancy of the genetic code (i.e., some amino acids are specified by more than one codon), evolution of the genomes of different organisms or classes of organisms has resulted in differential usage of redundant codons. This "codon bias" is reflected in the mean base composition of protein coding regions. For example, organisms with relatively low G+C contents utilize codons having A or T in the third position of redundant codons, whereas those having higher G+C contents utilize codons having G or C in the third position. Therefore, in reengineering genes for expression, one may wish to determine the codon bias of the organism in which the gene is to be expressed. Looking at the usage of the codons as determined for genes of a particular organism deposited in GenBank can provide this information. After determining the bias thereof, the new gene sequence can be analyzed for restriction enzyme sites as well as other sites that could affect transcription such as exon:intron junctions, polyA addition signals, or RNA polymerase termination signals.

Genes encoding synthases can be placed in an appropriate vector, depending on the artisan's interest, and can be expressed using a suitable expression system. An expression vector, as is well known in the art, typically includes elements that permit replication of said vector within the host cell and may contain one or more phenotypic markers for selection of cells containing said gene. The expression vector will typically contain sequences that control expression such as promoter sequences, ribosome binding sites, and translational initiation and termination sequences. Expression vectors may also contain elements such as selenomethionine, or both. These and similar techniques are 45 subgenomic promoters, a repressor gene or various activator genes. The artisan may also choose to include nucleic acid sequences that result in secretion of the gene product, movement of said product to a particular organelle such as a plant plastid (see U.S. Pat. Nos. 4,762,785; 5,451,513 and 5,545,817), or other sequences that increase the ease of peptide purification, such as an affinity tag.

A wide variety of expression control sequences are useful in expressing mutated synthases when operably linked thereto. Such expression control sequences include, for example, the early and late promoters of SV40 for animal cells, the lac system, the trp system, major operator and promoter systems of phage λ, and the control regions of coat proteins, particularly those from RNA viruses in plants. In *E. coli*, a useful transcriptional control sequence is the T7 RNA polymerase binding promoter, which can be incorporated into a pET vector as described by Studier et al., (1990) Methods Enzymology, 185:60–89.

For expression, a desired gene should be operably linked to the expression control sequence and maintain the appropriate reading frame to permit production of the desired synthase. Any of a wide variety of well-known expression vectors are of use in the present invention. These include, for

example, vectors consisting of segments of chromosomal, non-chromosomal and synthetic DNA sequences such as those derived from SV40, bacterial plasmids (including those from  $E.\ coli$  such as col E1, pCR1, pBR322 and derivatives thereof, pMB9), wider host range plasmids such as RP4, phage DNA such as phage  $\lambda$ , NM989, M13, and other such systems as described by Sambrook et al., (Molecular Cloning, A Laboratory Manual,  $2^{nd}$  Ed. (1989) Cold Spring Harbor Laboratory Press).

A wide variety of host cells are available for expressing synthase mutants of the present invention. Such host cells include, without limitation, bacteria such as *E. coli*, Bacillus and Streptomyces, fungi, yeast, animal cells, plant cells, insect cells, and the like. Preferred embodiments of the present invention include terpene synthase mutants that are expressed in *E. coli* or in plant cells. Said plant cells can either be in suspension culture or a culture on a solid support such as an agar-based medium.

Genes encoding synthases of the present invention can also be expressed in transgenic plant cells. In order to 20 produce transgenic plants, vectors containing a nucleic acid construct encoding a mutant terpene synthase are inserted into the plant genome. Preferably, these recombinant vectors are capable of stable integration into the plant genome. One variable in making a transgenic plant is the choice of a selectable marker gene. A selectable marker gene is used to identify transformed cells against a high background of untransformed cells. Such selectable marker genes include but are not limited to aminoglycoside phosphotransferase gene of transposon Tn5 (Aph II) Which encodes resistance to the antibiotics kanamycin, neomycin, and G418, as well as those genes which encode for resistance or tolerance to glyphosate, hygromycin, methotrexate, phosphinothricin, imidazolinones, sulfonylureas, and triazolophyrimidine herbicides, such as chlorosulfuron, bromoxynil, dalapon and the like. In addition to a selectable marker gene, it may be desirable to use a reporter gene. In some instances a reporter gene may be used with a selectable marker. Reporter genes allow the detection of transformed cell and may be used at the discretion of the artisan. A list of these reporter genes is 40 provided in K. Weising et al., 1988, Ann. Rev. Genetics,

The genes are expressed either by promoters expressing in all tissues at all times (constitutive promoters), by promoters expressing in specific tissues (tissue-specific promoters), 45 promoters expressing at specific stages of development (developmental promoters), and/or promoter expression in response to a stimulus or stimuli (inducible promoters). The choice of these is at the discretion of the artisan.

Several techniques exist for introducing foreign genes 50 into plant cells, and for obtaining plants that stably maintain and express the introduced gene. Such techniques include acceleration of genetic material coated directly into cells (U.S. Pat. No. 4,945,050). Plant may also be transformed using Agrobacterium technology (U.S. Pat. Nos. 5,177,010, 55 5,104,310, 5,149,645, 5,469,976, 5,464,763, 4,940,838, 4,693,976, 5,591,616, 5,231,019, 5,463,174, 4,762,785, 5,004,863, and 5,159,135; European Patent Applications 116718, 290799, 320500, 604662, 627752, 0267159, and 0292435. Other transformation technologies include whiskers technology, see U.S. Pat. Nos. 5,302,523 and 5,464, 765. Electroporation technology has also been used to transform plants, see WO 87/06614, WO 92/09696 and WO 93/21335 and U.S. Pat. Nos. 5,472,869 and 5,384,253. Viral vector expression systems can also be used such as those 65 described in U.S. Pat. Nos. 5,316,931, 5,589,367, 5,811,653, and 5,866,785.

In addition to numerous technologies for transforming plants, the type of tissue that is contacted with the genes of interest may vary as well. Suitable tissue includes, but is not limited to, embryogenic tissue, callus tissue, hypocotyl, meristern and the like. Almost all plant tissues may be transformed during dedifferentiation using the appropriate techniques described herein.

Regardless of the transformation system used, a gene encoding a mutant synthase is preferably incorporated into a gene transfer vector adapted to express said gene in a plant cell by including in the vector an expression control sequence (plant promoter regulatory element). In addition to plant promoter regulatory elements, promoter regulatory elements from a variety of sources can be used efficiently in plant cells to express foreign genes. For example, promoter regulatory elements of bacterial origin, such as the octopine synthase promoter, the nopaline synthase promoter, the mannopine synthase promoter may be used. Promoters of viral origin, such as the cauliflower mosaic virus (35S and 19S) are also desirable. Plant promoter regulatory elements also include, but are not limited to, ribulose-1,6bisphosphate carboxylase small subunit promoter, betaconglycinin promoter, phaseolin promoter, ADH promoter, heat-shock promoters, and tissue specific promoters and the like. Numerous promoters are available to skilled artisans for use at their discretion.

It should be understood that not all expression vectors and expression systems function in the same way to express the mutated gene sequences of the present invention. Neither do all host cells function equally well with the same expression system. However, one skilled in the art may make a selection among these vectors, expression control sequences, and host without undue experimentation and without departing from the scope of this invention.

Once a synthase of the present invention is expressed, the protein obtained therefrom can be purified so that structural analysis, modeling, and/or biochemical analysis can be performed, as exemplified herein. The nature of the protein obtained can be dependent on the expression system used. For example, genes, when expressed in mammalian or other eukaryotic cells, may contain latent signal sequences that may result in glycosylation, phosphorylation, or other posttranslational modifications, which may or may not alter function. Once the proteins are expressed, they can be easily isolated and purified using techniques common to the person having ordinary skill in the art of protein biochemistry and as described in Colligan et al., (1997) Current Protocols in Protein Science, Chanda, V. B., Ed., John Wiley & Sons, Inc. Such techniques often include the use of cation-exchange or anion-exchange chromatography, gel filtration-size exclusion chromatography, and the like. Another technique that may be commonly used is affinity chromatography. Affinity chromatography can include the use of antibodies, substrate analogs, or histidine residues (His-tag technology).

Once purified, mutants of the present invention may be characterized by any of several different properties. For example, such mutants may have altered active site surface charges of one or more charge units. In addition, the mutants may have an altered substrate specificity or spectrum of reaction product relative to a non-mutated synthase.

The present invention allows for the characterization of mutant terpene synthase by crystallization followed by X-ray diffraction. Polypeptide crystallization occurs in solutions where the polypeptide concentration exceeds it solubility maximum (i.e., the polypeptide solution is supersaturated). Such solutions may be restored to equilib-

rium by reducing the polypeptide concentration, preferably through precipitation of the polypeptide crystals. Often polypeptides may be induced to crystallize from supersaturated solutions by adding agents that alter the polypeptide surface charges or perturb the interaction between the polypeptide and bulk water to promote associations that lead to crystallization.

Compounds known as "precipitants" are often used to decrease the solubility of the polypeptide in a concentrated solution by forming an energetically unfavorable precipitating depleted layer around the polypeptide molecules (Weber, 1991, Advances in Protein Chemistry, 41:1–36). In addition to precipitants, other materials are sometimes added to the polypeptide crystallization solution. These include buffers to adjust the pH of the solution and salts to reduce the solubility of the polypeptide. Various precipitants are known in the art and include the following: ethanol, 3-ethyl-2-4 pentanediol, and many of the polyglycols, such as polyethylene glycol.

Commonly used polypeptide crystallization methods include the following techniques: batch, hanging drop, seed initiation, and dialysis. In each of these methods, it is important to promote continued crystallization after nucleation by maintaining a supersaturated solution. In the batch method, polypeptide is mixed with precipitants to achieve supersaturation, the vessel is sealed and set aside until crystals appear. In the dialysis method, polypeptide is retained in a sealed dialysis membrane that is placed into a solution containing precipitant. Equilibration across the membrane increases the polypeptide and precipitant concentrations thereby causing the polypeptide to reach supersaturation levels.

In the preferred hanging drop technique (McPherson, 1976, J. Biol. Chem., 6300-6306), an initial polypeptide mixture is created by adding a precipitant to a concentrated polypeptide solution. The concentrations of the polypeptide and precipitants are such that in this initial form, the polypeptide does not crystallize. A small drop of this mixture is placed on a glass slide that is inverted and suspended over a reservoir of a second solution. The system is then sealed. Typically, the second solution contains a higher concentration of precipitant or other dehydrating agent. The difference in the precipitant concentrations causes the protein solution to have a higher vapor pressure than the solution. Since the system containing the two solutions is sealed, an equilibrium is established, and water from the polypeptide mixture transfers to the second solution. This equilibrium increases the polypeptide and precipitant concentration in the polypeptide solution. At the critical concentration of polypeptide and precipitant, a crystal of the polypeptide may

Another method of crystallization introduces a nucleation site into a concentrated polypeptide solution. Generally, a concentrated polypeptide solution is prepared and a seed crystal of the polypeptide is introduced into this solution. If the concentration of the polypeptide and any precipitants are correct, the seed crystal will provide a nucleation site around which larger crystal forms. In preferred embodiments, the crystals of the present invention are formed in hanging drops with 15% PEG 8000; 200 mM magnesium acetate or magnesium chloride, 100 mM 3-(N-morpholino)-2-hydroxypropanesulfonic acid (pH 7.0), 1 mM dithiothreitol as precipitant.

Some proteins may be recalcitrant to crystallization. However, several techniques are available to the skilled 65 artisan to induce crystallization. The removal of polypeptide segments at the amino or carboxyl terminal end of the 26

protein may facilitate production of crystalline protein samples. Removal of such segments can be done using molecular biology techniques or treatment of the protein with proteases such as trypsin, chymotrypsin, subtilisin. Such procedures can result in the removal of flexible polypeptide segments that may negatively affect crystallization.

The crystals so produced have a wide range of uses. For example, high quality crystals are suitable for X-ray or neutron diffraction analysis to determine the three-dimensional structure of a mutant synthase and to design additional mutants thereof. In addition, crystallization can serve as a further purification method. In some instances, a polypeptide or protein will crystallize from a heterogeneous mixture into crystals. Isolation of such crystals by filtration, centrifugation, etc., followed by redissolving the polypeptide affords a purified solution suitable for use in growing the high-quality crystals needed for diffraction studies. The high-quality crystals may also be dissolved in water and then formulated to provide an aqueous solution having other uses as desired.

Because synthases may crystallize in more than one crystal form, the structural coordinates of  $\alpha$ -carbons of an active site determined from a synthase or portions thereof, as provided by this invention, are particularly useful to solve the structure of other crystal forms of synthases. The structural coordinates, as provided herein, may also be used to solve the structure of synthases having  $\alpha$ -carbons position within the active sites in a manner similar to the wild-type yet having R-groups that may or may not be identical. Furthermore, the structural coordinates disclosed herein may be used to determine the structure of the crystalline form of other proteins with significant amino acid or structural homology to any functional domain of a synthase. One method that may be employed for such purpose is molecular replacement. In this method, the unknown crystal structure, whether it is another crystal form of a synthase, a synthase having a mutated active site, or the crystal of some other protein with significant sequence identity and/or structural homology of a synthase may be determined using the coordinates given in Tables 10 and/or 11. This method provides sufficient structural form for the unknown crystal more efficiently than attempting to determine such information ab initio. In addition, this method can be used to 45 determine whether or not a given synthase in question falls within the scope of this invention.

As further disclosed herein, synthases and mutants thereof may be crystallized in the presence or absence of substrates and substrate analogs. The crystal structures of a series of 50 complexes may then be solved by molecular replacement and compared to that of the wild-type to assist in determination of suitable replacements for R-groups within the active site, thus making synthase mutants according to the present invention.

All mutants of the present inventions may be modeled using the information disclosed herein without necessarily having to crystallize and solve the structure for each and every mutant. For example, one skilled in the art may use one of several specialized computer programs to assist in the process of designing synthases having mutated active sites. Examples of such programs can be as follows: GRID (Goodford, 1985, J. Med. Chem., 28:849–857); MCSS (Miranker and Karplus, 1991, Proteins: Structure, Function and Genetics, 11:29–34); AUTODOCK (Goodsell and Olsen, 1990, Proteins: Structure, Function, and Genetics, 8:195–202); and DOCK (Kuntz et al., 1982, J. Mol. Biol., 161:269–288). In addition, specific computer programs are

also available to evaluate specific substrate-active site interactions and the deformation energies and electrostatic interactions resulting therefrom. MODELLER is a computer program often used for homology or comparative modeling of the three-dimensional structure of a protein. A. Sali & T. L. Blundell. J.Mol.Biol. 234, 779–815, 1993. A preselected polypeptide sequence to be modeled is aligned with one or more terpene synthases whose crystal structures are known and the MODELLER program is used to calculate a fullatom model, based on optimum satisfaction of spatial 10 restraints. Such restraints can include, inter alia, homologous structures, fluorescence spectroscopy, NMR experiments, or atom—atom potentials of mean force.

The present invention enables synthase mutants to be made and crystal structures thereof to be solved. Moreover, by virtue of the present invention, the location of the active site and the interface of substrate therewith permit the identification of desirable R-groups for mutagenesis. The particular embodiments of this invention are further exemplified in the Examples. However, those skilled in the art will readily appreciate that the specific experiments detailed are only illustrative of the invention as described more fully in the claims, which follow thereafter.

#### **EXAMPLE 1**

Generation of Mutant TEAS Genes

Construct Generation and Expression

All mutant enzymes were constructed by the Quick-Change method (Stratagene). Manufacturer's instructions 30 were followed, except as noted. Mutations were confirmed by DNA sequencing, and plasmids containing the desired mutation were used to transform BL-21 (DE3) expression cells. Protein was expressed, purified, and stored at -80° C.

from a TEAS-pET28b(+) template using the following primers: GTTGAATGCTACTTTTCGGCATTAGGAGTTTAT (sense) (SEQ ID NO:13) and ATAAACTCCTAATGC-CGAAAAGTAGCATTCAAC (antisense) (SEQ ID NO:14). Mutagenesis was carried out according to the manufacturers 40 ID No:2 was changed to a phenylalanine residue by siteinstructions, except that sense and antisense strands were generated in separate reactions. For each, 30 plasmidcopying cycles of one minute, annealing at 55° C. and 16 minutes extension at 68° C. were carried out. The two reaction mixtures were then combined, heated to 95° C. for 45 shown in SEQ ID No: 7. 2.5 minutes, and cooled to room temperature before Dpnl treatment.

TEAS C440W. The TEAS C440W mutant was generated from the TEAS-pET28b(+) template using the following primers: GCTAGTGTAATTATATGGCGAGTTATC-GATGAC (sense) (SEQ ID NO:15) and GTCATC-GATAACTCGCCATATAATTACACTAGC (antisense) (SEO ID NO:16)

TEAS W273S/C440W. The TEAS C440W/W273S mutant was constructed from a TEAS W273S-pET28b(+) 55 template using the primers described for generation of TEAS C440W.

TEAS 406/407 random library. For generation of a library of TEAS mutants with random amino acids at positions 406 and 407, two 50 microliter QuickChange reactions were carried out with the TEAS-pET28b(+) template and the GCACTAGCAACTACCACATATprimers TACNNSNNSGCGACAACATCGTATTTGGGCATG (sense) (SEQ ID NO:17) and CATGCCCAAATACGATGT-TGTCGCSNNSNNGTAATATGTGGTAGTTGCTAGTGC (antisense) (SEQ ID NO:18), in which N denotes A, C, G, or T and S denotes C or G. By this choice of nucleotides, the

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reaction included primers which coded for all possible amino acid combinations at positions 406 and 407. No adjustment was made for differing numbers of codons among amino acids. In order to ensure efficient reactions, and to minimize the preference for hybridization of wildtype primers to the template, the primers were designed to be longer than those used to generate the mutations described above. In addition, they were HPLC purified prior to use. After 18 cycles of plasmid copying, the reaction was incubated for two hours with Dpnl, ethanol precipitated, and redissolved in 5 microliters water. Each of four 40 microliter aliquots of E. coli NovaBlue (Novagen) cells were electroporated with 1.5 microliters of the redissolved DNA. After a recovery period, the cells were plated on kanamycin-LB-agar plates. In order to transfer the newly constructed plasmids to expression cells, the colonies were scraped from all four plates, and used to start an 8 mL culture grown in liquid LB medium at 37° C. for 8 hours. Plasmid purified from this culture was used to transform 20 microliters of competent BL-21 (DE3) cells.

For storage of the constructs, each individual colony was used to inoculate 100 microliters of LB medium containing kanamycin (50 micrograms/mL) in 96-well culture plates. The cells were grown at  $37^{\circ}$  C. until the  $A_{600}$  reached 25 approximately one; 100 microliters of 30% glycerol in LB were then added, and the plates were frozen at -80° C. A set of randomly selected colonies were grown from individual glycerol stocks of some colonies, and plasmids were extracted for sequencing. Approximately 30 percent of the colonies were found to be wild-type. Nucleotide and amino acid sequences for TEAS 406/407 mutant genes and proteins are shown in SEQ ID NOS:11 and 12.

TEAS Y520F The tyrosine residue at position 520 of SEQ ID No:2 was changed to a phenylalanine residue by site-TEAS W273S. The TEAS W273S mutant was generated 35 directed mutagenesis with primers, in a manner similar to that described above. For Y520F the TAT codon was changed to TTC. The nucleotide sequence of the mutant gene is shown in SEQ ID No:5.

TEAS Y527F The tyrosine residue at position 527 of SEQ directed mutagenesis with primers, in a manner similar to that described above. For Y527F, the TAC codon at position 527 of the TEAS amino acid sequence was changed to TTC. The nucleotide sequence of the mutant TEAS Y527F gene is

TEAS W273E The tryptophan residue at position 273 of SEQ ID NO:2 was changed to a phenylalanine residue by site-directed mutagenesis with primers, in a manner similar to that described above. For W273E, the TGG codon at position 273 of the TEAS amino acid sequence was changed to GAG. The nucleotide sequence of the mutant gene is shown in, SEQ ID No.:3.

## EXAMPLE 2

Expression and Isolation of Synthase Polypeptides

Unless otherwise noted, mutated and non-mutated TEAS proteins were expressed in Escherichia coli, purified by metal chelation, anion exchange, and gel filtration chromatography.

Constructs of TEAS and mutant TEAS proteins in the vector pET-28b(+) (Novagen) were expressed in E. coli cells. For a typical protein preparation of any of these enzymes, E. coli strain BL21 (DE3) cells containing the plasmid construct were grown at 37° C. in 4×1 L terrific broth to an  $A_{600}$ =1.0. The temperature was dropped to 22° C., and protein expression was induced by adding IPTG to

a final concentration of 0.1 mM. After 15-20 h, the cells were harvested by centrifugation, resuspended in 5 mL buffer A (20 mM Tris, 500 mM NaCl, 20 mM imidazole, pH 7.9) per 1 g cells (wet weight), and stirred for 0.5 h at 4° C. The cells were then lysed by sonication, and the resulting lysate was centrifuged for 0.7 h at 82,000×g. The supernatant, containing the protein, was loaded over a 2-3 mL Ni<sup>2+</sup> chelating histidine affinity column (Qiagen) equilibrated in buffer A, and the column was washed with additional buffer A until the  $A_{280}$  of the eluent returned to 10 baseline. The protein was then eluted with a 20-200 mM imidazole gradient in buffer A. Protein-containing fractions were pooled and dialyzed against buffer B (50 mM HEPES, 5 mM MgCl2, 1 mM DTT), then loaded onto an 8 mL MonoQ cation-exchange column (Pharmacia). The column 15 was washed with 20 column volumes buffer B, and the protein was eluted with a 0-500 mM NaCl gradient in buffer B. The resulting protein was further purified by gel filtration on a Superdex-200 column (Pharmacia) in 50 mM Tris, 100 mM NaCl, 5 mM MgCl2, 1 mM DTT, pH 8.0. Purified 20 protein was then dialyzed against 5 mM Tris, 5 mM NaCl, 1 mM DTT, pH 8.0, concentrated to 18-22 mg/mL, and stored at -80° C. in 100° L aliquots until needed.

#### **EXAMPLE 3**

#### Crystallization and Structural Analysis of Synthase Polypeptides

Crystal Growth and Microseeding: All crystallization attempts were carried out by the hanging-drop vapor diffusion method. Concentrated protein was mixed with an equal volume (2-5 uL each) of reservoir solution on a plastic cover slip. The cover slip was then inverted over a well of a plastic 24-well tissue culture plate, containing 0.5-1.0 mL of reservoir solution, and sealed by a layer of vacuum grease between the well and cover slip. The plates were incubated at 4° C. while the protein concentration in the hanging drop slowly increased by vapor diffusion. Approximately 300 different reservoir solutions, ranging pH 4.5-9 with a variety of precipitants and added salts, were assayed for crystallization of TEAS (SEQ ID NO:2). TEAS crystallized with a reservoir solution of 15% PEG 8000, 100 mM MOPSO (3-[N-morpholino]-2-hydroxypropanesulfonic acid), 200 mM magnesium acetate, 1 mM DTT, pH 6.9-7.3. For precipitant solution, then diluted to 50 microliters. After initial centrifugation to remove large particles, the suspension was serially diluted with additional precipitant solution, and a small volume of a diluted seed stock was added to each new crystallization drop. For microseeding, crystals which 50 were no longer rapidly growing (usually 2 weeks after drops were set up), were "rinsed" by serially transferring them through two to three drops of reservoir solution. The crystal was then transferred to a fresh drop containing protein and tion as in the initial growth. Individual crystals varied in their degree of internal order. In some cases, several crystals were screened to identify a well-diffracting crystal with low

Data collection: Prior to data collection, crystals were 60 transferred to a drop of reservoir solution, or reservoir solution containing a compound to be soaked into the crystal. A small volume of cryoprotecant solution (15% PEG8000, 100 mM MOPSO, 200 mM Mg acetate, 20% ethylene glycol, 1 mM DT7, pH 7) was then added to the drop. After a short equilibration time (1-5 minutes), the crystal was transferred to a drop of cryoprotectant, or

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cryoprotectant with soaking compound added. After another short equilibration time, the crystal was picked up on a nylon loop, and quickly mounted for data collection in a stream of cold nitrogen gas (90-110K).

The TEAS crystals belonged to the tetragonal space group P4<sub>1</sub>2<sub>1</sub>2; the unit cell dimensions varied by a few angstroms between crystals, but on average a=126 Å, c=122 Å. The uncomplexed TEAS structure was initially refined to 2.8 Å (Table 11) against data collected from a crystal grown in the presence of 2 mM FHP (Table 10). Electron density at the active site allowed unambiguous modeling of FHP, the ARC and J-K loops, and nine additional residues at the NH<sub>2</sub> terminus. The refined TEAS-FHP model consisted of residues 17 to 548, three Mg<sup>2</sup>+ions. 150 water molecules, and one FHP molecule. The three-dimensional coordinates for TEAS in the presence of bound substrate is shown in Table 10. The three-dimensional coordinates for TEAS in the absence of FHP is shown in Table 11.

Crystals of TEAS complexed with trifluoro-farnesyl diphosphate (F3-FPP) were also prepared. In these crystals, a well-ordered diphosphate binding pocket was also observed. The A-C loop and the NH2-terminal segment exhibited well-defined electron density, the A-C loop was translated toward the active site, and there was strong electron density for the diphosphate moiety of F3-FPP. The hydrophobic pocket, however, remained flexible; the J-K loop and the farnesyl moiety of F3-FPP were disordered.

Homology models were created and energy-minimized using the Swiss PDB viewer interface of the SwissModel program (Peitsch M C (1996), Biochem. Soc. Trans., 24:274-279 and Guex N. and Peitsch M C, 1997, Electrophoresis., 18:2714–2723). Active site volumes were calculated with VOIDOO (Kleywegt, G. J., and Jones, T. A., CCP4/ESF-EACBM Newsletter on Protein Crystallography., 29, 26–28, 1993). To make closed active site cavities, the energy-minimized diphosphate moiety from the modeled TEAS cyclase reaction was appended to the residue equivalent to TEAS D301.

TEAS W273S crystal structures. Two TEAS W273S structures, in the presence of FHP, were determined from different crystals; both crystals appeared to be well ordered, as clear main-chain and side-chain density were apparent for residues throughout the protein, including the frequently microseeding, an existing crystal was crushed in a few uL of 45 mobile helices D1, D2, and E. Initial difference electron density maps from both crystals immediately revealed the W273S mutations. The two crystals were designated W273S-1 and W273S-2.

In each structure, the loops surrounding the active site were ordered, resulting in a closed active site pocket. The A/C loop in each structure was translated toward the active site, forming part of its outer rim, as observed in the wild-type TEAS/FHP complex. However, while the J/K loop of W273S-1 adopted the same conformation observed in the reservoir solution, and equilibrated against a reservoir solu- 55 wild-type TEAS/FHP complex, the same loop in W273S-2 adopted a different conformation. In this conformation of the J/K loop, Tyr527 moved away from the side chain of residue 273. In addition, Tyr520 and Asp525 were placed distal to the side chain of Asp444. Hydrogen bonds previously observed between the J/K loop, Arg266, and the N-terminal loop were also missing in the W273S-2 structure.

> The W273S-2 conformation does not appear to be an effect of the W273S mutation, as it was also observed in a wild-type TEAS crystal soaked with the epi-aristolochene mimic deoxycapsidiol, despite the fact that no electron density was readily apparent for the deoxycapsidiol molecule in that structure. Further, the TEAS active site loops

were distant from crystal contacts, and their conformations were not likely to be artifacts of crystal packing. It is possible that at different stages of the TEAS reaction, the enzyme's J/K loop exists in different, defined conformations, and that each of these crystal structures has captured an image of a different conformation. In both W273S structures, residues other than Arg266 and those on the J/K loop did not undergo significant rearrangement from the conformations observed in wild-type TEAS.

In each W273S crystal structure, electron density in the active site suggested that the substrate mimic FHP binds in multiple conformations. Some regions of this density possibly represented bound water molecules in the mutant active site. The presence of water molecules in the mutant active site is consistent with the observation that TEAS W273S gives rise to multiple hydroxylated terpenoid reaction products.

The FHP electron density in each W273S crystal structure was sufficient to suggest that FHP existed in a more extended conformation in the W273S structure, compared to the more tightly folded conformation of FHP in the wild-type TEAS/FHP complex. The observation that the active site of W273S binds multiple conformations of FHP is consistent with the fact that W273S converts FPP to multiple terpenoid hydrocarbon products.

TEAS C440W/W273S: TEAS C440W/W273S crystal- 25 lized under conditions identical to wild-type TEAS. A 0.3 mm crystal was soaked for 20 minutes in reservoir solution saturated in farnesyl hydroxy phosphate (FHP). After cryoprotection and flash freezing as described for wild-type TEAS, data were collected on a laboratory source with Cu-Ka radiation (MacScience Corp., Japan). A starting model of uncomplexed TEAS (Table 11) (Brookhaven Protein Database Code 5EAT (PDB 5EAT), with waters and magnesiums removed, was positioned against the mutant data with the rigid body module of the software program X-PLOR (A. T. Brunge, X-PLOR Version 3.1—A System for X-Ray Crystallography and NMR Yale University Press, New Haven, 1992, pp. 187–207). Rounds of positional and restrained b-factor refinement with bulk solvent modeling were also carried out in X-PLOR, with manual model building and adjustment carried out in the software program O (Jones, T A, Zou, J Y, Cowan, S W, and Kjeldgaard, M., Acta Cryst. D., 49:148-157, 1993). Additional rounds of refinement and map calculation using the CNS program suite resulted in significantly improved maps; this improvement was likely due to improved bulk solvent modeling.

TEAS C440W: TEAS C440W crystallized under conditions identical to wild-type TEAS, except that crystals nucleated less readily and were generally smaller. A mutant crystal was soaked for 6 hours in reservoir solution saturated in FHP before flash-freezing and data collection at SSRL beamline 7-1 (Stanford Synchrotoon Radiation Laboratory, Menlo Park, Calif.). A starting model of TEAS-FHP (Table 10), with water molecules, ligands, and residues 523–532 of SEQ ID NO:2 removed, was positioned against the data with 55 the rigid body module of X-PLOR. Rounds of positional and restrained b-factor refinement with bulk solvent and overall anisotropic temperature factor modeling were also carried out in X-PLOR, and manual model building and adjustment were carried out in the software program O. As with the double mutant, electron density maps were noticeably improved after refinement and map calculation in CNS.

## EXAMPLE 4

#### Terpene Synthase Enzyme Assays

Synthase activity assays were carried out based on the assay described in Vogeli and Chappell, Plant Physiol.

94:1860 (1990) and Vogeli, et al., Plant Physiol. 93:182 (1990). In general, radio-labeled (<sup>3</sup>H or <sup>14</sup>C) substrate was incubated with enzyme at room temperature in a buffered magnesium salt solution (200 mM Tris, pH 8, 50 mM Mg chloride, 1 mM DTT, unless otherwise noted); hydrocarbon products were then selectively extracted into an organic solvent such as hexane. The hexane extract generally was treated with silica gel to remove prenyl alcohols and other oxygenated compounds generated by non-enzymatic hydrolysis of substrate, which partition inefficiently into hexane. Hydrocarbon products present in the hexane phase were quantitated by scintillation counting.

A subsequent extraction with a more polar organic solvent such as ethyl acetate was sometimes carried out. Oxygenated compounds more efficiently partition into ethyl acetate-type solvents. Compounds present in the ethyl acetate phase were also quantitated by scintillation counting.

Substrate concentrations typically ranged from 0.1 nanomolar to 100 micromolar. In some assays, the substrate was not radiolabeled. Reactions generally were carried out in triplicate for each substrate concentration. Protein concentration was determined by the Bradford method. For determination of steady-state kinetic parameters, enzyme concentrations were chosen such that generation of products over time was linear throughout the course of the reaction.

Diterpene synthase assays typically were carried out using <sup>3</sup>H geranylgeranyl diphosphate (GGPP) and enzyme in 250 mM Tris, 10 mM Mg chloride, 1 mM DTT, pH 8.0. Sesquiterpene synthase assays typically were carried out using <sup>14</sup>C or <sup>3</sup>H FPP and enzyme in 100 mM Tris, 30 mM Mg chloride, 1 mM DTT, pH 8.0. Monoterpene synthase assays typically were carried out using <sup>3</sup>H GPP and enzyme. As a control for nonspecific binding of GPP by protein, identical reactions were set up which contained BSA, rather than enzyme.

Product analysis of wild type and mutant TEAS enzymes by Ag-TLC. Terpenoid hydrocarbon products are not readily separated by thin layer chromatography on normal or reverse-phase plates; however, some can be separated by argentation TLC (Ag-TLC), in which silica plates are first treated with silver nitrate. Ag-TLC described here generally followed the procedure described by Back et al., Arch. Biochem. Biophys. 315:527 (1994). A silica TLC plate was 45 dipped in 15% silver nitrate (aqueous), then dried for 3–5 hours at 110° C. After spotting of tritiated enzymatic products (solvent extract), the plate was developed in benzene:hexane, ethyl acetate (50:50:1, by volume), sprayed with En<sup>3</sup>Hance (NEN) fluorography spray, placed on film, and exposed for several days to several weeks. Long exposure times were generally necessary, as silver-nitrate treatment of the TLC plate appeared to cause quenching of the fluorography reagent's fluorescence. Alternatively, <sup>14</sup>C labelled products were detected after one to two days without the use of fluorography spray.

#### EXAMPLE 5

#### Activity of TEAS W273S

Diterpene Synthase Activity of TEAS W273S. The TEAS W273S enzyme and radiolabelled GGPP were incubated as described above and hydrocarbon products were extracted with hexane. Oxygenated products were then extracted with ethyl acetate. Reactions using wild-type TEAS gave counts
 lower than buffer alone. TEAS W273S, on the other hand, gave counts that were significantly higher for both the hexane and ethyl acetate extracts. Hydrocarbon products

formed from GGPP by W273S were distinct from the products made by acid-catalyzed loss of diphosphates from GGPP. See FIG. 3.

Sesquiterpene Synthase Activity of TEAS W273S. Products of FPP turnover by the purified TEAS W273S mutant were analyzed by argentation thin-layer chromatography (Ag-TLC). One major reaction product had an  $R_f$  of 0.7 by Ag-TLC, which was distinct from both 5-epi-aristolochene ( $R_f$ =0.78) and vetispiradiene ( $R_f$ =0.63). See FIG. 4. Preliminary GC/MS data showed that hexane extracts from FPP turnover by TEAS W273S contained at least four terpene hydrocarbons, with mass spectra distinct from either 5-epi-aristolochene or vetispiradiene. One of these products had a mass spectrum similar to germacrene A.

#### **EXAMPLE 6**

#### Activity of TEAS C440W/W273S

Diterpene Synthase Activity of TEAS C440W/W273S. The mutant TEAS C440W/W273S protein contains a tryp- 20 tophan residue at position 440 and a serine residue at position 273. Assays with GGPP were carried out using 0.5 micromolar <sup>3</sup>H GGPP, various concentrations of unlabelled GGPP (Echelon), and enzyme. Reactions were incubated for 60 minutes at room temperature. The TEAS C440W/W273S mutant protein converted GGPP to hexane-extractable products, whereas the wild-type enzyme did not. The results indicated that the product profile was altered compared to wild-type TEAS. Hexane-extractable products of GGPP turnover by the double mutant were analyzed by Ag-TLC. 30 The products included two species ( $R_z=0.11$  and 0.28) that were distinct from the hydrolysis product geranyl geraniol (R<sub>t</sub>=0.0). To verify that products generated by TEAS C440W/W273S from GGPP were not the hydrolysis product, geranylgeraniol, a sample was analyzed by 35 Ag-TLC. A reaction containing  ${}^{3}H$  GGPP (5  $\mu m$ ) and enzyme (40 µm) in 100 microliters buffer was incubated overnight at room temperature. As controls, <sup>3</sup>H GGPP was incubated in reaction buffer alone and in reaction buffer adjusted to pH 1.5. Both the enzymatic and control reactions 40 were extracted with hexane, which was spotted on an argentation TLC plate, and developed and exposed as described above. The results, shown in FIG. 3, demonstrated that the products formed by TEAS C440W/W273S were different from those generated by non-enzymatic degrada- 45 tion of geranylgeranyl diphosphate.

Sesquiterpene Synthase Activity of TEAS C440W/W273S. Reactions with FPP as substrate were carried out with  $^{14}$ C FPP (9  $\mu$ m) and enzyme (160  $\mu$ m) in reaction buffer (20  $\mu$ l). After incubating for 30 minutes at room temperature, products made by TEAS C440W/W273S were analyzed by Ag-TLC. The product profile of the double mutant was similar to that of TEAS W273S, with the addition of a major product having an  $R_f$  of 0.57. The new product was distinct from both 5-epi-aristolochene and vetispiradiene. Several other products were also formed, many of which migrated slowly upon argentation TLC. See FIG. 4.

## EXAMPLE 7

## Activity of TEAS C440W

Diterpene Synthase Activity of TEAS C440W. Enzyme assays with TEAS C440W were carried out as described in Example 6. As shown in FIG. 3, no hexane-extractable products were detectable by Ag-TLC after an overnight 65 incubation at room temperature with  $160 \, \mu \text{m}$  of enzyme and  $9 \, \mu \text{m}$  radiolabeled GGPP in  $20 \, \mu \text{l}$  volume.

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Sesquiterpene Synthase Activity of TEAS C440W. Ag-TLC analysis of the products made from radiolabelled by purified TEAS C440W detected the formation at least one major terpenoid hydrocarbon product ( $R_f$  0.63) that was distinct from 5-epi-aristolochene ( $R_f$  0.78) and vetispiradiene. The reactions product profile on Ag-TLC is shown in FIG. 4. Small amounts of slowly-migrating products ( $R_f$  0–0.09) were also formed.

GC/MS analysis of the hexane extract of TEAS C440W terpenoid hydrocarbon reaction products confirmed that this mutant formed a single major sesquiterpene hydrocarbon product as well as a small number of minor hydroxylated products. The mass spectrum of the major product closely matched the published mass spectrum of the spirocyclic compound hinesene. Hinesene differs from vetispiradiene in the stereochemistry at the C3 methyl group.

#### **EXAMPLE 8**

#### Activity of TEAS W273E

Sesquiterpene Synthase Activity of TEAS W273E. Reactions to determine the products made by TEAS W273E using FPP as substrate were carried out essentially as described above, using radiolabeled FPP. The results indicated that at least one product other than 5-epi-aristolochene was formed. The results also indicated that alkylation of TEAS by FPP had occurred. The alkylation was dependent upon the presence of MgCl<sub>2</sub> in the reaction mixture. In control experiments, boiled W273E-TEAS, as well as wild-type TEAS and BSA, were not alkylated. These results indicate that alkylation had occurred at position 273 and that the amino acid residue at position 273 is part of the active site.

### EXAMPLE 9

#### Activity of TEAS Y520F

Sesquiterpene Synthase Activity of TEAS Y520F. Reactions with radiolabeled FPP and TEAS Y520F enzyme were carried out essentially as described above. Reaction products were analysed by Ag-TLC and by GC/MS. A major product of the TEAS Y520F reaction had the same GC retention time as authentic germacrene A and the same mass spectrum as authentic germacrene A. The retention time and mass spectrum of this product were different from 5-epi-aristolochene.

#### EXAMPLE 10

### Activity of TEAS Y527F

Enzymatic Activity of TEAS Y527F. A crude extract of TEAS Y527F enzyme was made by inducing expression in *E. coli* cells, and sonicating the cells. The sonicate was clarified and the supernatant used for enzyme assays. No products were observed in assays using GPP as a substrate, indicating that TEAS Y527F does not have monoterpene synthase activity. Reaction products were obtained using FPP as a substrate. Analysis of these products by Ag-TLC indicated that products other than 5-epi-aristolochene were generated by the TEAS Y527F enzyme.

#### EXAMPLE 11

### Alignment of Terpene Synthase Sequences

Residues 265 to 535 of the TEAS primary amino acid sequence (SEQ ID NO: 2) were aligned with the full-length amino acid sequence of a limonene synthase (SEQ ID NO:

22), using the BLASTp program (NCBI) with a BLOSUM 62 scoring matrix, a gap open value of 11, a gap extension value of 1, an x\_dropoff value of 50, an expect value of 10, a wordsize of 3 and no filtering of low complexity sequences. The output of the alignment program, shown in 5 Table 12, included a gap between residues 527 and 528 of the TEAS sequence (numbered as 263 and 264 in the alignment output). Residues 321, 324, 345, 348, 349, 427, 452, 453, 454, 455, 458, 492, 496, 569, 572, 573, 577, 579 and 580 were selected as having the most suitable alignment with the 19 TEAS residues. Residue 580 of limonene cyclase instead of residue 583 was selected as aligning with residue 528 of TEAS, in order to maintain the spatial orientation of structural aspects found in TEAS, i.e.,  $\alpha$ -helices,  $\beta$ -sheets and loops shown in FIG. 1 and Table 10. 15

A region including residues 579 to 847 of the taxadiene primary amino acid sequence of SEQ ID NO: 44 was aligned with the full-length amino acid sequence of a bornyl diphosphate synthase (SEQ ID NO: 26), using the BLASTp program (NCBI) with a BLOSUM 62 scoring matrix, a gap open value of 11, a gap extension value of 1, an x\_dropoff value of 50, an expect value of 10, a wordsize of 3 and no filtering of low complexity sequences. The output of the alignment program, shown in Table 13, included a gap between residues 453 and 454 of the bornyl diphosphate synthase sequence. Residues 321, 324, 344, 347, 348, 426, 451, 452, 453, 454, 457, 492, 496, 568, 571, 572, 576, 578 and 579 of the bornyl diphosphate synthase were selected as having the most suitable alignment with residues 584, 587, 606, 609, 610, 688, 713, 714, 715, 716, 719, 753, 757, 831, 834, 835, 839, 841 and 842 of the query region sequence of SEQ ID NO: 44. Residues 453 and 454 of bornyl diphosphate synthase were selected to align with residues 715 and 716 of taxadiene synthase, in order to maintain the spatial orientation of structural aspects expected to be present in taxadiene synthase, i.e.,  $\alpha$ -helices,  $\beta$ -sheets and loops shown in FIG. 1 and Table 10.

Residues 265 to 535 of the TEAS primary amino acid sequence (SEQ ID NO: 2) were aligned with the full-length amino acid sequence of a δ-selinene synthase (SEQ ID NO: 48), using the BLASTp program (NCBI) with a BLOSUM 50 scoring matrix, a gap open value of 13, a gap extension 45 value of 2, an x\_dropoff value of 50, an expect value of 10, a wordsize of 3 and no filtering of low complexity sequences. The output of the alignment program is shown in Table 14. Residues 300, 303, 324, 327, 328, 406, 431, 432, 433, 434, 437, 471, 475, 548, 551, 552, 556, 558 and 559 of SEQ ID NO:48 were selected as having the most suitable alignment with residues 270, 273, 294, 297, 298, 376, 401, 402, 403, 404, 407, 440, 444, 516, 519, 520, 525, 527 and 528 of SEQ ID NO 2.

Residues 307 to 593 of the primary amino acid sequence of γ-humulene synthase (SEQ ID NO: 50) were aligned with the full-length amino acid sequence of abietadiene synthase (SEQ ID NO: 56), using the BLASTp program (NCBI) with a BLOSUM 62 scoring matrix, a gap open value of 11, a gap extension value of 1, an x\_dropoff value of 50, an expect value of 10, a wordsize of 3 and no filtering of low complexity sequences. The output of the alignment program is shown in Table 15. Residues 590, 593, 614, 617, 618, 696, 721, 722, 723, 724, 727, 761, 765, 837, 840, 841, 845, 847 and 848 of the diterpene synthase (SEQ ID NO: 56) were

selected as having the most suitable alignment with residues 312, 315, 336, 339, 340, 419, 444, 445, 446, 447, 450, 484, 488, 562, 565, 566, 570, 572 and 573 of the sesquiterpene synthase query sequence (SEQ ID NO: 50).

#### EXAMPLE 12

Generation of Novel Monoterpene Synthase Genes

A DNA sequence encoding a pinene synthase (SEQ ID NO:20) is used to construct a library of mutant pinene synthase genes. Random mutations are introduced at nucleotides encoding one or more of the following nine amino acid residues: L, C, C, G, H, S, L, G and Y, which correspond to positions 351, 372, 480, 481, 482, 485, 519, 600 and 601 of SEQ ID NO:20.

In some cases, the pinene synthase coding sequence is randomly mutated at nucleotides encoding one or more of amino acid residues 348, 375, 376, 597, 605, 607 and 608, which correspond to positions Y, I, T, F, D, Y and S of SEQ ID NO:20. The pinene synthase coding sequence is sometimes mutated at nucleotides encoding one or more of the following amino acid residues: Y, S and G, which correspond to positions 454, 479 and 523 of SEQ ID NO:20. In some cases, mutations at these ten positions are made in addition to mutations at nucleotides encoding the nine residues mentioned above. In other cases, mutations at these ten positions are made without introducing mutations at the nine residues mentioned above.

The pinene synthase coding sequence DNA is inserted in the pET28b(+) vector and mutagenized using the Quick-Change® method, following a protocol similar to that described in Example 1 for the TEAS 406/407 random library. The primers used to generate mutations are synthesized as indicated in Example 1, using N or S as nucleotides in the desired codons in order to generate random mutants.

Specific mutations at one or more of the above 19 pinene synthase amino acid residues are made by site-directed mutagenesis using a protocol similar to that described in Example 1 for TEAS. Primers are made that have specific A, T, C or G substitutions in the codons to be mutated, in order to generate the desired mutant(s).

Random and/or specific mutations are prepared in a manner similar to that described above to alter amino acid residues of other monoterpene synthases, e.g., limonene synthase, (SEQ ID NOS:22 or 58), myrcene synthase (SEQ ID NO:30), +sabinene synthase (SEQ ID NO:54), 1, 8 cincole synthase (SEQ ID NO:24) and +bornyl diphosphate synthase (SEQ ID NO:26), at residues whose  $\alpha$ -carbons have the interatomic distances and structural coordinates described in Tables 1–6.

#### **EXAMPLE 13**

Generation of Novel Sesquiterpene Synthase Genes

A DNA sequence encoding a cadinene synthase (SEQ ID NO:33) is used construct a library of mutant cadinene synthases. Random mutations are introduced at nucleotides encoding one or more of the following nine amino acid residues: W, I, S, G, Y, L, C, L and Y, which correspond to amino acid residues 280, 301, 409, 410, 411, 414, 448, 527 and 528 encoded by SEQ ID NO:33.

In some cases, the cadinene synthase coding sequence is mutated at nucleotides encoding one or more of amino acid residues G, A, S, M, D, Y and T, which correspond to amino acid residues 277, 304, 305, 524, 532, 534 and 535 encoded by SEQ ID NO:33. In addition, the cadinene synthase coding sequence is sometimes mutated at nucleotides encoding one or more of the following amino acid residues: 383, 408 and 452, which correspond to amino acids Y, T and D encoded by SEQ ID NO:33. In some cases, these mutations are made in addition to mutations at the nine residues mentioned above. In other cases, mutations at these ten residues are made without introducing mutations at the nine 10 residues mentioned above.

The cadinene synthase coding sequence is mutated using the QuickChange® method in the pET28b(+) vector, following a protocol similar to that described in Example 1 for the TEAS 406/407 random library. The primers used to generate mutations are synthesized as indicated in Example 11

Specific mutations at one or more of the above cadinene synthase amino acid residues are made by site-directed 20 mutagenesis using a protocol similar to that described in Example 1 for TEAS.

Random and/or specific mutations are prepared in a manner similar to that described above to alter amino acid residues of other sesquiterpene synthases, e.g., vetispiradiene synthase (SEQ ID NO:32), germacrene C synthase (SEQ ID NO:52), E-alpha-bisabolene synthase (SEQ ID NO:50),  $\delta$ -selinene synthase (SEQ ID NO:48), e-b-farnesene synthase (SEQ ID NO:28), at residues whose  $\alpha$ -carbons have the interatomic distances and structural coordinates described in Tables 1–6.

#### EXAMPLE 14

#### Generation of Novel Diterpene Synthase Genes

A DNA sequence encoding an abietadiene synthase (SEQ ID NO:56) is used construct a library of mutant abietadiene synthases. Random mutations are introduced at nucleotides encoding one or more of the following nine amino acid residues: S, S, I, A, L, V, G, F and Y, which correspond to positions 593, 614, 722, 723, 724, 727, 761, 840 and 841 of SEQ ID NO:56.

In some cases, the abietadiene synthase coding sequence is mutated at nucleotides encoding one or more of amino acid residues I, S, T, M, D, L and T, which correspond to positions 590, 617, 618, 837, 845, 847 and 848 of SEQ ID NO:56. The abietadiene synthase coding sequence is sometimes mutated at nucleotides encoding one or more of the following amino acid residues: Y, S and N, which correspond to positions 696, 721 and 765 of SEQ ID NO:56. In some caes, these mutations are made in addition to mutations at the nine residues mentioned above. In other cases, mutations are made at these ten residues without introducing mutations at the nine residues mentioned above.

The abietadiene synthase coding sequence is mutated using the QuickChange® method in the pET28b(+) vector, following a protocol similar to that described in Example 1 for the TEAS 406/407 random library. The primers used to generate mutations are synthesized as indicated in Example 60

Specific mutations at one or more of the above abietadiene synthase amino acid residues are made by site-directed mutagenesis using a protocol similar to that described in Example 1 for TEAS.

Random and/or specific mutations are prepared in a manner similar to that described above to alter amino acid residues of other diterpene synthases at amino acid residues whose  $\alpha$ -carbons have the interatomic distances and structural coordinates described in Tables 1–6, e.g., casbene synthase (SEQ ID NO:42) and taxadiene synthase (SEQ ID NO:44).

#### **EXAMPLE 15**

#### Expression of Mutant Synthases in Insect, Mammalian and Bacterial Cells

Constructs containing nucleic acids encoding mutant synthases of Examples 12, 13 and/or 14 are introduced into cultured cells of the insect *Spodoptera frugiperda* using a baculovirus expression vector. After expression of the gene, the mutant enzyme is isolated and purified from each clone.

Constructs containing nucleic acids encoding mutant synthases of Examples 12, 13 and/or 14 are introduced into cultured HeLa cells using an expression vector having an SV40 promoter. After expression of the gene, the mutant enzyme is isolated and purified from each clone.

Constructs containing nucleic acids encoding mutant synthases of Examples 12, 13 and/or 14 are introduced into *E. Coli* BL-21 on a plasmid vector as described in Example 1. The mutant synthase gene is expressed and the mutant enzyme is isolated and purified as described in Example 2.

#### Other Embodiments

To the extent not already indicated, it will be understood by those of ordinary skill in the art that any one of the various specific embodiments herein described and illustrated may be further modified to incorporate features shown in other of the specific embodiments.

It is to be understood that while the invention has been described in conjunction with the Detailed Description thereof, that the foregoing description is intended to illustrate, and not limit the scope of the invention, which is defined by the scope of the appended claims. Other aspects, advantages, and modifications are within the scope of the following claims.

TABLE 1

α-Carbon	X Position	Y Position	Z Position
1	119.144	43.487	44.133
2	120.203	38.695	43.506
3	114.058	43.884	41.015
4	109.327	46.145	41.743
5	110.682	46.410	45.284
6	99.381	42.920	45.148
7	103.445	38.054	44.605
8	106.807	36.336	45.151
9	107.629	38.010	41.804
10	109.375	34.842	40.617
11	111.944	37.854	37.602
12	110.233	31.098	47.361
13	109.178	33.314	52.875
14	115.915	32.218	48.369
15	118.846	34.443	51.796
16	116.461	32.848	54.290
17	114.100	38.006	55.620
18	116.617	41.285	51.702
19	114.855	43.486	54.238

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## TABLE 2

α-carbon	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19
1	0.0	5.0	6.0	10.5	9.0	19.8	16.6	14.3	13.0	13.5	11.2	15.6	16.7	12.5	11.8	15.0	13.7	8.3	11.0
2	5.0	0.0	8.4	13.3	12.4	21.3	16.8	13.7	12.7	11.9	10.2	13.1	15.4	9.2	9.4	12.8	13.6	9.3	12.9
3	6.0	8.4	0.0	5.3	6.0	15.3	12.6	11.3	8.7	10.2	7.2	14.8	16.6	13.9	15.1	17.4	15.7	11.3	13.3
4	10.5	13.3	5.3	0.0	3.8	11.0	10.4	10.7	8.3	11.4	9.6	16.1	17.0	16.8	18.1	19.6	16.8	13.3	13.9
5	9.0	12.4	6.0	3.8	0.0	11.8	11.1	10.8	9.6	12.5	11.6	15.5	15.2	15.4	15.9	17.3	13.8	10.1	10.3
6	19.8	21.3	15.3	11.0	11.8	0.0	6.4	9.9	10.2	13.6	15.5	16.2	15.8	20.0	22.2	21.8	18.7	18.5	18.0
7	16.6	16.8	12.6	10.4	11.1	6.4	0.0	3.8	5.0	7.8	11.0	10.1	11.1	14.3	17.4	17.0	15.3	15.3	15.9
8	14.3	13.7	11.3	10.7	10.8	9.9	3.8	0.0	3.8	5.4	9.3	6.6	8.6	10.5	13.9	13.7	12.9	12.8	14.1
9	13.0	12.7	8.7	8.3	9.6	10.2	5.0	3.8	0.0	3.8	6.0	9.2	12.1	12.1	15.4	16.1	15.3	13.8	15.4
10	13.5	11.9	10.2	11.4	12.5	13.6	7.8	5.4	3.8	0.0	5.0	7.8	12.4	10.5	14.6	15.5	16.0	14.7	17.0
11	11.2	10.2	7.2	9.6	11.6	15.5	11.0	9.3	6.0	5.0	0.0	12.0	16.2	12.8	16.1	18.0	18.2	15.3	17.8
12	15.6	13.1	14.8	16.1	15.5	16.2	10.1	6.6	9.2	7.8	12.0	0.0	6.0	5.9	10.2	9.5	11.4	12.8	14.9
13	16.7	15.4	16.6	17.0	15.2	15.8	11.1	8.6	12.1	12.4	16.2	6.0	0.0	8.2	9.8	7.4	7.3	11.0	11.7
14	12.5	9.2	13.9	16.8	15.4	20.0	14.3	10.5	12.1	10.5	12.8	5.9	8.2	0.0	5.0	6.0	9.5	9.7	12.8
15	11.8	9.4	15.1	18.1	15.9	22.2	17.4	13.9	15.4	14.6	16.1	10.2	9.8	5.0	0.0	3.8	7.1	7.2	10.2
16	15.0	12.8	17.4	19.6	17.3	21.8	17.0	13.7	16.1	15.5	18.0	9.5	7.4	6.0	3.8	0.0	5.8	8.8	10.8
17	13.7	13.6	15.7	16.8	13.8	18.7	15.3	12.9	15.3	16.0	18.2	11.4	7.3	9.5	7.1	5.8	0.0	5.7	5.7
18	8.3	9.3	11.3	13.3	10.1	18.5	15.3	12.8	13.8	14.7	15.3	12.8	11.0	9.7	7.2	8.8	5.7	0.0	3.8
19	11.0	12.9	13.3	13.9	10.3	18.0	15.9	14.1	15.4	17.0	17.8	14.9	11.7	12.8	10.2	10.8	5.7	13.8	0.0

TABLE 3 TABLE 5

α-Carbon	X Position	Y Position	Z Position	25 —				
1	119.144	43.487	44.133	_	α-Carbon	X Position	Y Position	Z Position
2	120.203	38.695	43.506					
3	114.058	43.884	41.015		1	120.203	38.695	43.506
4 5	109.327 110.682	46.145 46.410	41.743 45.284	30	2	114.058	43.884	41.015
6	106.807	36.336	45.151		3	106.807	36.336	45.151
7	107.629	38.010	41.804			107.620	20.010	44.004
8	109.375	34.842	40.617		4	107.629	38.010	41.804
9	111.944	37.854	37.602		5	109.375	34.842	40.617
10 11	110.233 115.915	31.098 32.218	47.361 48.369	35	6	111.944	37.854	37.602
12	118.846	34.443	51.796		7	110.233	31.098	47.361
13 14	116.461 114.100	32.848 38.006	51.796 54.290 55.620		8	118.846	34.443	51.796
15 16	116.617 114.855	41.285 43.486	51.702 54.238		9	116.461	32.848	54.290

TABLE 4

α-Carbon	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16
1	0.0	5.0	6.0	10.5	9.0	14.3	13.0	13.5	11.2	15.6	12.5	11.8	15.0	13.7	8.3	11.0
2	5.0	0.0	8.4	13.3	12.4	13.7	12.7	11.9	10.2	13.1	9.2	9.4	12.8	13.6	9.3	12.9
3	6.0	8.4	0.0	5.3	6.0	11.3	8.7	10.2	7.2	14.8	13.9	15.1	17.4	15.7	11.3	13.3
4	10.5	13.3	5.3	0.0	3.8	10.7	8.3	11.4	9.6	16.1	16.8	18.1	19.6	16.8	13.3	13.9
5	9.0	12.4	6.0	3.8	0.0	10.8	9.6	12.5	11.6	15.5	15.4	15.9	17.3	13.8	10.1	10.3
6	14.3	13.7	11.3	10.7	10.8	0.0	3.8	5.4	9.3	6.6	10.5	13.9	13.7	12.9	12.8	14.1
7	13.0	12.7	8.7	8.3	9.6	3.8	0.0	3.8	6.0	9.2	12.1	15.4	16.1	15.3	13.8	15.4
8	13.5	11.9	10.2	11.4	12.5	5.4	3.8	0.0	5.0	7.8	10.5	14.6	15.5	16.0	14.7	17.0
9	11.2	10.2	7.2	9.6	11.6	9.3	6.0	5.0	0.0	12.0	12.8	16.1	18.0	18.2	15.3	17.8
10	15.6	13.1	14.8	16.1	15.5	6.6	9.2	7.8	12.0	0.0	5.9	10.2	9.5	11.4	12.8	14.9
11	12.5	9.2	13.9	16.8	15.4	10.5	12.1	10.5	12.8	5.9	0.0	5.0	6.0	9.5	9.7	12.8
12	11.8	9.4	15.1	18.1	15.9	13.9	15.4	14.6	16.1	12.2	5.0	0.0	3.8	7.1	7.2	10.2
13	15.0	12.8	17.4	19.6	17.3	13.7	16.1	15.5	18.0	9.5	6.0	3.8	0.0	5.8	8.8	10.8
14	13.7	13.6	15.7	16.8	13.8	12.9	15.3	16.0	18.2	11.4	9.5	7.1	5.8	0.0	5.7	5.7
15	8.3	9.3	11.3	13.3	10.1	12.8	13.8	14.7	15.3	12.8	9.7	7.2	8.8	5.7	0.0	3.8
16	11.0	12.9	13.3	13.9	10.3	14.1	15.4	17.0	17.8	14.9	12.8	10.2	10.8	5.7	3.8	0.0

TABLE 6	TABLE 6-continued

α-											α-									
Carbon	1	2	3	4	5	6	7	8	9	5	Carbon	1	2	3	4	5	6	7	8	9
1	0	8.4	13.7	12.7	11.9	10.2	13.1	9.4	12.8		6	10.2	7.2	9.3	6	5	0	12	16.1	18
2	8.4	0	11.3	8.7	10.2	7.2	14.8	15.1	17.4		7	13.1	14.8	6.6	9.2	7.8	12	0	10.2	9.5
3	13.7	11.3	0	3.8	5.4	9.3	6.6	13.9	13.7		8	9.4	15.1	13.9	15.4	14.6	16.1	10.2	0	3.8
4	12.7	8.7	3.8	0	3.8	6	9.2	15.4	16.1		9	12.8	17.4	13.7	16.1	15.5	18	9.5	3.8	0
5	11 0	10.2	5.4	20	0	5	70	146	15.5											

TABLE 7

								17	JDL	Æ /									
					Orde	red A	rran	geme	nt of	R-G	roup	s at c	z-carl	ons	1–19	1			
	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19
A	С	W	I	I	S	Y	Т	Т	Т	Y	L	С	D	V	Т	Y	D	Y	Т
В	С	W	I	I	S	Y	T	S	T	Y	L	C	D	I	T	Y	D	Y	T
С	G	W	I	A	S	Y	T	C	G	Y	L	C	D	M	L	Y	D	Y	T
D E	G C	W	I L	A T	S S	Y	T S	S A	G G	Y	L I	C A	D N	M A	L L	V Y	D D	Y Y	T T
F	G	W	L	L	S	Y	S	T	V	Ĥ	L	G	D	A	V	Y	D	Y	T
Ğ	Č	w	Ĺ	T	S	Ŷ	Š	Ā	Ġ	Y	Ī	Ā	N	A	Ĺ	Ŷ	D	Ŷ	ŝ
H	L	W	I	T	T	Y	S	V	G	N	L	F	D	V	L	Y	D	F	T
I	P	W	I	V	D	Y	S	T	Α	G	L	S	D	Α	C	Y	D	Y	T
J	A	W	V	C	G	F	T	S	C	I	M	G	N	Ç	S	Y	D	Y	S
K L	N C	F W	F N	L I	G T	A Y	E S	I I	T S	A G	T M	G L	N D	I A	T M	Y	E D	F H	T Q
M	S	W	V	L	T	Y	S	S	S	Y	L	G	G	V	L	Y	D	F	T
N	N	F	F	L	v	Ñ	A	T	Ĺ	Ā	L	G	N	Ĺ	S	Ŷ	E	F	Ť
O	C	W	N	I	T	Y	I	S	G	P	L	L	D	Α	M	Y	D	Н	G
P	C	W	N	V	T	Y	I	G	G	I	L	L	D	Α	I	$\mathbf{Y}$	D	F	G
Q	С	Y	L	L	T	F	A	V	T	M	Т	G	N	Ι	Т	Y	D	Y	T
R S	C S	W	I F	I I	T V	Y F	S S	I S	S S	A V	I I	L L	D N	A V	I I	Y V	D	D	G G
S T	S	W	I	A	T	Y	S	V	A	S	I	L	D D	v A	I	Y	D D	H F	G
Ü	N	w	N	L	T	Y	S	Ĭ	S	S	I	F	N	S	M	Y	D	Н	G
v	F	L	A	Q	T	Ŷ	s	Ī	Ğ	Q	Ĺ	s	D	T	I	F	D	F	Ğ
W	I	S	S	T	V	Y	S	I	Α	L	V	G	N	M	F	Y	D	L	T
X	Y	L	C	I	T	Y	S	С	G	Η	S	L	G	F	G	Y	D	Y	S
Y	G	S W	F	I	T	F	S	S	S	V	I	L	N	A	V	Y	D	H	G
Z AA	Y A	W A	A N	C L	T T	Y N	S A	S L	G T	M S	L T	G C	D M	L L	I L	Y Y	D D	L Y	Y N
BB	F	L	C	v	T	Y	S	s	A	Y	V	L	G	L	L	Y	D	F	S
CC	F	w	Ã	M	Ť	Ŷ	Ñ	T	G	M	Ĺ	s	Ď	Ī	M	Ŷ	Ď	F	Š
DD	Y	M	С	V	T	F	V	S	S	G	I	L	G	F	V	Y	D	Y	T
EE	V	S	G	Q	V	Y	S	V	G	L	С	W	N	V	F	Y	D	Y	G
FF	С	S	G	T	T	M	F	A	L	G	V	G	N	L	F	Y	D	F	T
GG HH	C C	S A	G G	T T	T T	M M	S S	F F	A A	L L	I I	G G	N N	L V	F F	Y Y	D D	F Y	T T
П	I	W	V	I	S	Y	T	Т	G	L	V	I	N	T	S	Y	D	Y	T
JJ	Ý	w	Å	Ĉ	Т	Ŷ	S	S	G	M	Ĺ	Ġ	D	Ĺ	I	Ý	D	Ĺ	Ŷ
KK	Ċ	W	I	I	S	Y	T	S	T	Y	L	Ċ	D	V	Т	Y	D	Y	T
LL	С	W	I	I	S	Y	T	T	T	Y	L	С	D	I	T	Y	D	Y	T
MM	С	W	N	I	T	Y	S	I	S	G	M	L	D	Α	M	Y	D	Η	G
NN	F	Α	A	Q	T	Y	S	I	G	Q	L	S	D	T	I	F	D	F	G
00	F	A	I	A	Т	Y	S	V	A	S	I	L	D	A	I	Y	D	F	G

TABLE 8

				Orde	red A	rran	geme	nt of F	R-Gro	ups a	tα-c	arbons	1–16			
	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16
A	С	W	I	I	S	Т	Т	Y	L	С	V	Т	Y	D	Y	Т
В	С	W	I	I	S	S	T	Y	L	С	I	T	Y	D	Y	T
C	G	W	I	Α	S	С	G	Y	L	C	M	L	Y	D	Y	T
D	G	W	I	Α	S	S	G	$\mathbf{Y}$	L	С	M	L	Y	D	Y	T
E	C	W	L	T	S	Α	G	Y	I	Α	Α	L	Y	D	Y	T
F	G	W	L	L	S	T	V	Η	L	G	Α	V	Y	D	Y	T
G	С	W	L	T	S	Α	G	Y	I	Α	Α	L	Y	D	Y	S
Н	L	W	I	T	Т	V	G	N	L	F	V	L	Y	D	F	T
I	P	W	I	V	D	Т	Α	G	L	S	Α	С	Y	D	Y	T
J	Α	W	V	С	G	$\mathbf{S}$	С	I	M	G	С	S	Y	D	Y	S

TABLE 8-continued

	Ordered Arrangement of R-Groups at α-carbons 1–16															
	_			Oluc	ncu F	man	genie	nt OI I	<b>C</b> -O10	ирь а	i u-c	aroons	1-10			
	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16
K	N	F	F	L	G	I	Т	Α	Т	G	I	Т	Y	Е	F	Т
L	С	W	N	I	T	I	S	G	M	L	Α	M	Y	D	Η	Q
M	S	W	V	L	T	S	S	Y	L	G	V	L	Y	D	F	T
N	N	F	F	L	V	T	L	Α	L	G	L	S	Y	Е	F	T
O	С	W	N	I	T	S	G	P	L	L	Α	M	$\mathbf{Y}$	D	Н	G
P	C	W	N	V	T	G	G	I	L	L	A	I	Y	D	F	G
Q	C	Y	L	L	T	V	Т	M	Т	G	I	T	Y	D	Y	T
R	С	W	I	I	T	I	S	A	I	L	A	I	Y	D	D	G
S	S	W	F	I	V	S	S	V	I	L	V	I	Y	D	H	G
Т	S	W	I	A	Т	V	Α	S	I	L	A	I	Y	D	F	G
U	N	W	N	L	T	I	S	S	I	F	S	M	Y	D	H	G
V	F	L	A	Q	T	I	G	Q	L	S	Т	I	F	D	F	G
W	I	S	S	Т	V	I	A	L	V	G	M	F	Y	D	L	T
X	Y	L	С	I	Т	С	G	H	S	L	F	G	Y	D	Y	S
Y	G	S	F	I	Т	S	S	V	I	L	A	V	Y	D	Η	G
Z	Y	W	Α	С	T	S	G	M	L	G	L	I	Y	D	L	Y
AA	A	Α	N	L	Т	L	T	S	Т	С	L	L	Y	D	Y	N
BB	F	L	С	V	T	$\mathbf{S}$	A	Y	V	L	L	L	Y	D	F	S
CC	F	W	Α	M	T	T	G	M	L	$\mathbf{s}$	I	M	Y	D	F	S
DD	Y	M	C	V	Т	S	S	G	I	L	F	V	Y	D	Y	T
EE	V	S	G	Q	V	V	G	L	С	W	V	F	Y	D	Y	G
FF	С	S	G	T	T	Α	L	G	V	G	L	F	Y	D	F	T
GG	C	S	G	T	T	F	A	L	I	G	L	F	Y	D	F	T
$_{ m HH}$	С	Α	G	T	T	F	Α	L	I	G	V	F	Y	D	$\mathbf{Y}$	T
II	I	W	V	I	S	T	G	L	V	I	T	S	Y	D	Y	T
JJ	Y	W	A	C	T	S	G	M	L	G	L	I	Y	D	L	Y
KK	C	W	I	I	S	S	T	Y	L	С	V	T	Y	D	Y	T
LL	C	W	I	I	S	T	T	Y	L	С	I	T	Y	D	Y	T
MM	C	W	N	I	T	I	S	G	M	L	A	M	Y	D	H	G
NN	F	Α	A	Q	T	I	G	Q	L	S	Т	I	F	D	F	G
00	F	A	I	A	Т	V	Α	S	I	L	A	Ι	Y	D	F	G

				TAB	LE 9					35				TA	BLE 9-c	continu	ed		
_		0	rdered 2	Arrange	ments o	of α-Ca	rbons 1	<b>-</b> 9		_			(	Ordered	Arrangen	ents of c	α-Carbons	1–9	
	1	2	3	4	5	6	7	8	9	_		1	2	3	4	5	6 7	8	9
A	W	I	Т	T	Y	L	С	Т	Y	40	НН	A	G	F	A		I G	F	Y
В	W	I	S	T	Y	L	С	T	$\mathbf{Y}$		II	W	V	T	G		V I	S	Y
С	W	I	С	G	$\mathbf{Y}$	L	С	L	Y		JJ	W	Α	S	G	M	L G	I	Y
D	W	I	S	G	Y	L	С	L	Y		KK	W	I	S	T	Y	L C	T	Y
E	W	L	Α	G	Y	I	Α	L	Y		LL	W	I	T	T	Y	L C	T	Y
F	$\mathbf{W}$	L	T	$\mathbf{v}$	Н	L	G	V	Y		MM	W	N	I	S	G	M L	M	Y
G	W	L	Α	G	$\mathbf{Y}$	I	Α	L	Y	45	NN	Α	Α	I	G	Q	L S	I	F
H	W	I	V	G	N	L	F	L	Y		00	Α	I	V	Α		I L	I	Y
I	W	I	T	Α	G	L	S	C	Y							-			
J	W	V	S	C	I	M	G	S	Y										
K	F	F	I	T	Α	T	G	T	Y										
L	W	N	I	S	G	M	L	M	Y						TADI	E 10			
M	W	V	S	S	Y	L	G	L	Y	50					TABL	E 10			
N	F	F	T	L	Α	L	G	S	Y	50					e m i				
O	W	N	S	G	P	L	L	M	Y		S	tructura			of Tobacco				hase
P	W	N	G	G	I	L	L	I	Y		_		With	Farnesy	ıl Hydroxy	phospho	nate Boun	d	
Q	Y	L	V	T	M	T	G	T	Y					ъ.					-
R	W	I	I	S	Α	I	L	I	Y			Atom	Resi-	Resi-			_		В-
S	W	F	S	S	V	I	L	I	Y		Atom	Type	due	due #	X	Y	Z	OCC	factor
T	W	I	V	Α	S	I	L	I	Y	55		СВ	VAL	17	105.641	55.031	61.062	1.00	98.26
U	W	N	I	S	S	I	F	M	Y		1	CG1	VAL	17	105.641	56.123	61.269	1.00	98.26 97.24
V	L	Α	I	G	Q	L	S	I	F		2	CG1	VAL	17	104.598	53.957	62.133	1.00	94.24
W	S	S	I	Α	L	V	G	F	$\mathbf{Y}$		3								
X	L	С	C	G	H	S	L	G	Y		4	С	VAL	17	106.842	53.842	59.190	1.00	98.89
Y	S	F	S	S	V	I	L	V	Y		5	O N	VAL	17 17	107.108	52.650	59.359 59.594	1.00	96.64
Z	W	Α	S	G	M	L	G	I	Y	60	6 7		VAL	17	104.381 105.495	53.419		1.00	99.88
AA	Α	N	L	Т	S	T	С	L	Y			CA	VAL			54.412	59.646	1.00	99.06
BB	L	C	S	Ā	Y	v	Ĺ	L	Y		8	N	ALA	18	107.671	54.719	58.615	1.00	98.95
CC	W	A	Т	G	M	L	S	M	Y		9	CA	ALA	18	109.015	54.419	58.088	1.00	98.55
DD	M	C	S	S	G	I	L	V	Y		10	CB	ALA	18	110.007	55.478	58.572	1.00	97.57
EE	S	G	V	G	L	Ċ	W	F	Y		11	С	ALA	18	109.570	53.012	58.346	1.00	99.86
		_				v				65	12	0	ALA	18	109.580	52.170	57.447	1.00	100.00
FF	S	G	A	L	G		G	F	Y	03	13	N	ASP	19	110.068	52.793	59.562	1.00	99.07
GG	S	G	$\mathbf{F}$	Α	L	I	G	F	Y		14	CA	ASP	19	110.616	51.508	60.010	1.00	97.13

TABLE 10-continued

March   Marc	Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound									5	s	Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound							
16   CG   ASP   19   19,565   9,566   1,579   1,010   19,000   1,000	Atom				X	Y	z	OCC			Atom				x	Y	z	OCC	
19	15	СВ	ASP	19	109.507	50.447	60.064	1.00	96.62		88	CG	GLN	28	131.624	50.781	49.615	1.00	32.15
18																			
19										10									
20																			
22   CA   PHIE   20   112-925   501-97   571-90   1.09   584-17																			
CB   PHE   20																			
24 CG PHE 20 111.437 95.72 55.251 1.00 77.17 " 97 CG PHE 20 129.079 49.349 45.976 1.00 24.53   25 CD1 PHE 20 111.056 49.971 54.187 1.00 74.25   26 CD2 PHE 20 111.056 49.25 55.939 1.00 77.18   27 CE1 PHE 20 10.9581 49.25 55.939 1.00 77.18   28 CC2 PHE 20 110.9547 49.88   28 SS.35 1.00 72.55   29 CD2 PHE 20 110.9547 49.88   28 SS.35 1.00 78.19   29 CD2 PHE 20 110.9547 49.89   28 SS.35 1.00 78.19   29 CD2 PHE 20 110.9547 49.89   29 CD2 PHE 20 110.9547 49.99   29 CD2 PHE 20 110.9547 49.89   29 C																			
25										15									
Secondary   Cell   PHIE   20	25		PHE	20	110.691	49.971	54.147	1.00	74.72				PHE		128.241	49.241	47.089		26.35
28   CE2   PHE   20																			
20																			
Name										20									
32 CA SER 21 115.294 500/98 57.639 1.00 78.89										20									
33         CA         SER         21         116.65         505.50         73.95         100         75.96         106         CA         LEU         30         22.16         54.66         42.44         1,00         39.73           35         Ge         SER         21         117.499         48.250         57.731         1.00         80.91         1         108         CG         LEU         30         22.858         8.00         44.03         50.00         44.03         30         C         SER         21         117.30         51.00         50.00         1.00         CD         LEU         30         126.56         86.03         44.03         0         40.01           39         CD         PRO         22         118.737         35.80         55.50         1.00         60.39         111         C         LEU         30         13.03.84         54.74         42.09         1.00         43.00           41         CPRO         22         119.567         55.88         55.50         1.00         65.22         111         C         LEU         30         1.33.73         54.73         4.299         1.00         55.76         111         C         EER<																			
35         GB         SER         21         117,499         48,235         S8,515         100         75,81         000         96,73         25         109         GB         LEU         30         27,125         56,00         40,00         46,33           36         C         SER         21         117,305         51,00         56,00         10         69,67         25         100         CDL         LEU         30         127,508         83,033         44,05         10         54,37           38         N         PRO         22         118,111         52,134         56,691         10         63,25         111         CL         LEU         30         130,433         54,744         40,079         100         40,74           41         CS         PRO         22         118,505         55,803         10         63,92         111         C         LEU         30         13,438         54,744         42,009         100         40,74           41         CS         PRO         22         119,667         53,688         51,748         100         55,75         100         55,75         100         55,75         100         65,75         110																			
36   C   SIR   21   117.30   50.03   50.02   1.00   69.07   25   109   CD1   LEU   30   127.508   58.033   44.036   1.00   54.01   38.03   39. CD   PRO   22   118.11   52.124   56.99   1.00   63.25   111   C   LEU   30   127.508   56.738   44.29   1.00   54.01   39.00   40.75   40.00   40.00   40.00   40.75   40.00																			
No.   Sign   21   117,070   50,513   55,525   1,00   70,74   70,75   70,53										25									
38 N PRO 22 118.11 S2.134 56.90 1 .00 63.25										25									
18																			
41																			
42   CG   PRO   22   19.657   \$3.688   \$7.488   1.00   61.76   57.76   41.89   1.00   53.76   42.91   1.00   53.76   44.90   0.0   0.0   57.06   44.90   0.0   0.0   57.06   44.90   0.0   57.06   44.90   0.0   57.06   44.90   0.0   57.06   44.90   47.07   47.90																			
44										20									
44         O         PRO         22         120,236         50,771         55,859         1.00         52,85         118         O         SER         31         133,669         53,515         41,851         100         45,28           46         CA         SER         23         121,2030         52,208         53,724         100         51,757         51,660         100         53,579         118         0         SER         23         121,200         51,775         51,660         100         51,375         118         0         SER         23         121,200         51,775         51,660         100         51,375         100         45,40         35         121         CR         PHE         32         134,131         50,812         40,232         100         43,23           50         O         SER         23         122,160         51,210         53,878         1.00         61,762         122         CG         PHE         32         134,213         50,812         40,073         1.00         33,22           51         N         LEU         24         124,324         437,937         55,519         1.00         60,44         12         CC         PH										30									
46																			
48																			
48																			
Section   Color   Section   Color   Section										25									
51         N         LEU         24         123,101         50,004         54,168         1,00         58,09         124         CD2         PHE         32         135,266         48,948         8,858         1,00         22,49           53         CB         LEU         24         124,545         48,301         55,191         1,00         60,64         1,00         60,64         1,00         60,64         1,00         60,64         1,00         60,64         1,00         60,64         1,00         60,64         1,00         60,64         1,00         70,70         40         127         CZ         PHE         32         13,6261         48,023         38,647         1,00         28,23           56         CD2         LEU         24         123,539         55,851         1,00         70,77         129         0         PHE         32         134,698         52,256         37,829         1,00         40,51           57         C         LEU         24         126,529         50,754         5,808         1,00         50,75         1,30         8         8,808         1,00         50,523           59         N         TRP         25         126,563<								1.00		33			PHE						
55         CA         LEU         24         124,326         49,799         54,944         1,00         55,68         125         CE1         PHE         32         136,061         48,073         38,647         1,00         22,339           54         CG         LEU         24         123,413         47,379         55,651         1,00         67,70         40         128         CP         PHE         32         135,601         52,358         1,00         28,42           55         CDI         LEU         24         123,813         45,934         55,385         1,00         70.01         128         C         PHE         32         135,601         52,358         8,00         1,00         50,87           56         CDZ         LEU         24         126,555         50,313         51,988         1,00         50,23         131         CA         SER         33         137,855         52,816         3,889         1,00         55,26           58         O         LEU         24         126,563         50,635         51,688         1,00         50,23         131         CA         SER         33         138,687         4,04         1,00         41,38<																			
53         CB         LEU         24         124-548         48,301         55,191         1,00         60,64         126         CEZ         PHE         32         136,261         48,023         38,647         1,00         27,39           55         CD1         LEU         24         123,810         45,934         55,855         1,00         70,01         128         C         PHE         32         135,601         32,388         38,896         1,00         50,85           56         CD2         LEU         24         123,509         47,596         57,124         1,00         70,77         129         0         PHE         32         135,601         32,388         98,00         1,00         50,26         58         0         LEU         24         126,529         50,754         54,808         1,00         50,23         131         CA         SER         33         136,809         52,626         37,841         1,00         61,21           59         N         TRP         25         126,536         49,908         50,645         1,00         46,22         45         134         C         SER         33         136,641         51,583         30,731 <td< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></td<>																			
55         CD1         LEU         24         123,413         47,379         55,651         1.00         67,70         40         128         C         PHE         32         137,179         47,737         39,655         1.00         50,87           56         CD2         LEU         24         123,808         45,934         55,385         1.00         70,77         129         OP PHE         32         134,988         52,256         3,899         1.00         50,87           58         O         LEU         24         125,554         50,313         54,198         1.00         51,07         130         N         SER         33         136,899         52,266         38,999         1.00         55,26           58         O         LEU         24         125,554         50,036         51,977         1.00         44,22         4         133         06         SER         33         137,755         52,816         37,801         1.00         61,187           60         CA         TRP         25         126,563         50,636         51,977         1.00         44,22         45         132         CB         SER         33         138,648         51,25																			
56 CD2 LEU 24 123.810 45.934 55.365 1.00 70.77 129 O PHE 32 135.001 52.358 88.896 1.00 50.81 57 C LEU 24 123.598 47.596 57.124 1.00 70.77 129 O PHE 32 134.988 52.226 37.829 1.00 43.81 57 C LEU 24 126.553 50.754 54.808 1.00 50.23 131 CA SER 33 136.899 52.626 38.989 1.00 55.26 58 O LEU 24 126.529 50.754 54.808 1.00 45.50 132 CB SER 33 138.587 54.094 38.017 1.00 61.27 60 CA TRP 25 125.456 30.636 51.977 1.00 44.42 133 OG SER 33 138.587 54.094 38.017 1.00 61.87 61 CB TRP 25 126.536 49.908 50.645 1.00 40.22 45 134 CB SER 33 139.024 54.250 39.360 1.00 67.09 61 CB TRP 25 126.536 49.908 50.645 1.00 47.97 1.00 44.42 133 OG SER 33 139.045 54.250 39.360 1.00 67.09 62 CG TRP 25 125.583 48.510 50.867 1.00 47.97 1.00 45.60 1.00 67.99 62 CG TRP 25 125.504 47.407 51.384 1.00 50.67 1.00 45.60 1.00 67.09 62 CG TRP 25 125.704 47.219 51.729 1.00 45.66 1.00 67.09 61.00 67.09 61.00 67.00 61.00 67.00 61.00 67.00 61.00 67.00 61.00 67.00 61.00 67.00 61.00 67.00 61.00 67.0			LEU		123.413	47.379			67.70	40	127		PHE	32	137.179	47.737			28.24
57         C         LEU         24         125.554         50.313         54.198         1.00         51.07         130         N         SER         33         136.899         52.626         38.989         1.00         55.26         58         O         LEU         24         126.529         50.754         54.808         1.00         50.23         131         CA         SER         33         136.899         52.626         38.991         1.00         61.21           59         N         TRP         25         125.472         50.267         52.873         1.00         45.50         132         CB         SER         33         138.687         54.094         80.01         61.02         CB         TRP         25         126.563         50.636         51.977         1.00         44.22         45         133         OG         SER         33         138.641         1.583         37.31         1.00         69.07         60         CB         TRP         25         125.853         48.510         50.667         1.30         50.67         135         O. ER         33         138.641         50.93         80.00         50.94         63         CB         77.72         1.00										70									
58         O         LEU         24         126.529         50.754         54.808         1.00         50.23         131         CA         SER         33         137.755         52.816         37.841         1.00         61.21           59         N         TRP         25         125.472         50.267         52.873         1.00         44.52         132         CB         SER         33         137.755         52.816         37.841         1.00         61.21           60         CA         TRP         25         126.563         50.636         51.977         1.00         44.22         45         134         C         SER         33         139.488         51.329         39.500         1.00         67.09           62         CG         TRP         25         126.604         47.407         51.884         1.00         50.67         136         N         ILE         34         138.368         50.771         36.718         1.00         69.75           65         CE3         TRP         25         122.507         46.331         51.729         1.00         45.66         138         CB         ILE         34         138.408         8.69         35.442																			
60 CA TRP 25 126.563 50.636 51.977 1.00 44.42																			
61 CB TRP 25 126.556 49.908 50.645 1.00 46.22 45 134 C SER 33 138.641 51.583 37.731 1.00 59.75 62 CG TRP 25 125.853 48.510 50.867 1.00 47.97 135 O SER 33 139.488 51.329 38.589 1.00 59.49 63 CD2 TRP 25 126.604 47.407 51.384 1.00 50.67 136 N ILE 34 138.368 50.771 36.718 1.00 66.14 64 CE2 TRP 25 125.700 46.331 51.553 1.00 50.91 137 CA ILE 34 139.128 49.552 36.486 1.00 66.15 65 CE3 TRP 25 127.948 47.219 51.729 1.00 45.66 138 CB ILE 34 139.128 49.552 36.486 1.00 65.50 160 CD1 TRP 25 124.466 46.765 51.147 1.00 47.16 50 140 CG1 ILE 34 139.291 47.406 35.157 1.00 65.37 68 CZ2 TRP 25 126.101 45.088 52.053 1.00 52.99 141 CD1 ILE 34 139.291 47.406 35.157 1.00 65.37 69 CZ3 TRP 25 128.347 45.983 52.227 1.00 47.77 142 C ILE 34 140.544 49.875 36.013 1.00 70.13 70 CH2 TRP 25 127.423 44.934 52.384 1.00 51.93 141 CD1 ILE 34 140.544 49.875 36.013 1.00 70.13 70 CH2 TRP 25 127.945 52.663 1.00 47.77 142 C ILE 34 140.544 49.875 36.013 1.00 76.00 711 C TRP 25 127.957 52.550 52.063 1.00 43.75 72 O TRP 25 127.957 52.550 52.063 1.00 43.75 74 CA GIY 26 125.958 52.862 51.172 1.00 47.80 57 C GIY 26 126.210 54.267 50.894 1.00 39.84 77 N ASP 27 127.620 55.434 49.887 1.00 44.69 148 OD1 ASP 35 143.885 49.419 37.558 1.00 76.13 79 CB ASP 27 128.200 55.084 49.885 1.00 46.55 80 CG ASP 27 128.505 55.082 47.770 1.00 66.66 CB ASP 35 143.155 49.030 39.784 1.00 86.08 79 CB ASP 27 128.200 55.084 47.770 1.00 66.66 CB ASP 35 143.895 49.419 37.558 1.00 65.50 80 CG ASP 27 128.200 55.084 47.906 1.00 46.55 150 C ASP 35 143.895 49.419 37.558 1.00 65.50 80 CG ASP 27 128.200 55.084 47.906 1.00 46.55 150 C ASP 35 143.895 49.491 37.558 1.00 65.50 80 CG ASP 27 128.200 55.082 47.770 1.00 66.66 CB ASP 35 143.895 49.491 37.558 1.00 65.50 80 CG ASP 27 128.200 55.082 47.770 1.00 66.66 CB ASP 35 143.895 49.491 37.558 1.00 65.50 80 CG ASP 27 128.200 55.082 47.770 1.00 66.66 CB ASP 35 143.895 49.491 37.558 1.00 65.50 80 CG ASP 27 128.40 55.082 47.895 1.00 66.66 60 153 CA ASP 27 128.40 55.082 47.895 1.00 65.50 66.60 60 60 60 60 60 60 60 60 60 60 60 60 6																			
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67 NE1 TRP 25 124.466 46.765 51.147 1.00 47.16 50 140 CG1 ILE 34 139.291 47.406 35.157 1.00 65.37 68 CZ2 TRP 25 126.101 45.088 52.053 1.00 52.99 141 CD1 ILE 34 138.715 46.458 34.122 1.00 63.17 69 CZ3 TRP 25 128.347 45.983 52.227 1.00 47.77 142 C ILE 34 140.544 49.875 36.013 1.00 70.13 70 CH2 TRP 25 126.893 52.110 51.744 1.00 44.49 144 N ASP 35 141.554 94.543 36.782 1.00 73.05 72 O TRP 25 127.997 52.550 52.063 1.00 43.75 73 N GLY 26 125.988 52.862 51.172 1.00 47.80 75 145 CA ASP 35 142.935 49.673 36.388 1.00 70.88 73 N GLY 26 126.210 54.267 50.894 1.00 39.84 75 C GLY 26 126.375 53.696 48.580 1.00 46.55 149 OD2 ASP 35 143.155 49.030 39.784 1.00 92.06 76 O GLY 26 126.375 53.696 48.580 1.00 46.55 149 OD2 ASP 35 143.155 49.030 39.784 1.00 92.06 79 CB ASP 27 128.200 55.708 47.966 1.00 50.38 151 O ASP 35 143.155 49.030 39.784 1.00 92.06 79 CB ASP 27 127.307 58.091 47.770 1.00 66.06 79 CB ASP 27 127.307 58.091 47.770 1.00 66.06 81 OD1 ASP 27 127.482 59.318 47.597 1.00 64.78 153 CA ASN 36 142.949 49.430 31.577 1.00 72.78 82 OD2 ASP 27 129.441 54.857 47.686 1.00 46.14 156 OD1 ASN 36 142.949 49.430 31.577 1.00 72.78 84 O ASP 27 129.441 54.857 47.686 1.00 46.14 156 OD1 ASN 36 142.949 49.430 31.577 1.00 72.78 85 N GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 142.319 51.756 31.853 1.00 84.96 86 CA GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 144.361 46.704 31.906 1.00 65.06 86 CA GLN 28 130.759 52.921 48.461 1.00 28.69 65 159 O ASN 36 144.461 46.704 31.906 1.00 65.06																			
68 CZ2 TRP 25 126.101 45.088 52.053 1.00 52.99 141 CD1 ILE 34 138.715 46.458 34.122 1.00 63.17 69 CZ3 TRP 25 128.347 45.983 52.227 1.00 47.77 142 C ILE 34 140.544 49.875 36.013 1.00 70.13 70 CH2 TRP 25 127.423 44.934 52.384 1.00 51.93 143 O ILE 34 140.725 50.551 35.001 1.00 76.00 76.00 71 C TRP 25 127.6289 52.110 51.744 1.00 44.49 144 N ASP 35 141.545 49.454 36.782 1.00 73.05 72 O TRP 25 127.997 52.550 52.063 1.00 43.75 145 CA ASP 35 142.935 49.673 36.388 1.00 70.88 73 N GLY 26 126.210 54.267 50.894 1.00 39.84 147 CG ASP 35 143.895 49.419 37.558 1.00 76.13 74 CA GLY 26 126.210 54.267 50.894 1.00 39.84 146 CB ASP 35 143.288 48.547 38.638 1.00 84.32 75 C GLY 26 126.375 53.696 48.580 1.00 46.55 149 OD2 ASP 35 143.195 49.030 39.784 1.00 86.08 77 N ASP 27 127.620 55.434 49.287 1.00 46.92 150 C ASP 35 143.195 49.030 39.784 1.00 86.08 79 CB ASP 27 128.200 55.708 47.966 1.00 50.38 1 OD ASP 35 143.555 47.552 35.425 1.00 65.76 79 CB ASP 27 128.544 57.196 47.827 1.00 66.06 81 OD ASP 35 143.98 48.714 35.227 1.00 66.53 81 ODD ASP 27 128.546 57.196 47.827 1.00 66.478 153 CA ASN 36 142.940 49.241 34.019 1.00 66.55 81 ODD ASP 27 127.482 59.318 47.597 1.00 66.478 153 CA ASN 36 142.949 49.430 31.577 1.00 72.78 82 OD2 ASP 27 129.441 54.857 47.686 1.00 46.14 156 OD1 ASN 36 142.949 49.430 31.577 1.00 72.78 85 N GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 142.319 51.756 31.853 1.00 84.96 86 CA GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 144.461 46.704 31.906 1.00 65.06 86 CA GLN 28 130.759 52.921 48.461 1.00 28.69 65 159 O ASN 36 144.461 46.704 31.906 1.00 65.06										50									
70         CH2         TRP         25         127,423         44,934         52,384         1.00         51,93         143         O         ILE         34         140,725         50,551         35,001         1.00         76,00           71         C         TRP         25         126,893         52,110         51,744         1.00         44.49         144         N         ASP         35         141,545         49,454         36,782         1.00         73.05           72         O         TRP         25         127,997         52,550         52,063         1.00         43,75         145         CA         ASP         35         141,545         49,454         36,782         1.00         70.88           73         N         GLY         26         126,210         54,267         50,894         1.00         39,84         147         CG         ASP         35         143,895         49,419         37,558         1.00         76,13           75         C         GLY         26         126,210         54,467         50,894         1.00         44,69         148         OD1         ASP         35         142,931         47,387         38,638         1	68	CZ2	TRP	25	126.101	45.088	52.053	1.00	52.99		141	CD1	ILE	34	138.715	46.458	34.122	1.00	63.17
71 C TRP 25 126.893 52.110 51.744 1.00 44.49 144 N ASP 35 141.545 49.454 36.782 1.00 73.05 72 O TRP 25 127.997 52.550 52.063 1.00 43.75 145 CA ASP 35 142.935 49.673 36.388 1.00 70.88 73 N GLY 26 125.958 52.862 51.172 1.00 47.80 55 146 CB ASP 35 143.895 49.419 37.558 1.00 76.13 74 CA GLY 26 126.210 54.267 50.894 1.00 39.84 147 CG ASP 35 143.895 49.419 37.558 1.00 76.13 75 C GLY 26 126.744 54.449 49.483 1.00 44.69 148 OD1 ASP 35 143.288 48.547 38.638 1.00 92.06 76 O GLY 26 126.375 53.696 48.580 1.00 46.55 149 OD2 ASP 35 143.155 49.030 39.784 1.00 92.06 77 N ASP 27 127.620 55.434 49.287 1.00 46.92 150 C ASP 35 143.198 48.714 35.227 1.00 68.52 78 CA ASP 27 128.200 55.708 47.966 1.00 50.38 151 O ASP 35 143.555 47.552 35.425 1.00 65.76 79 CB ASP 27 128.544 57.196 47.827 1.00 57.61 80 CG ASP 27 127.307 58.091 47.770 1.00 66.06 61.53 80 CG ASP 27 127.482 59.318 47.597 1.00 64.78 153 CA ASN 36 142.949 49.430 31.577 1.00 72.78 82 OD2 ASP 27 129.441 54.857 47.686 1.00 46.14 156 OD1 ASN 36 142.949 49.430 31.577 1.00 79.82 83 C ASP 27 129.441 54.857 47.686 1.00 46.14 156 OD1 ASN 36 142.949 49.430 31.577 1.00 79.82 85 N GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 142.319 51.756 31.853 1.00 84.96 86 CA GLN 28 130.759 52.921 48.461 1.00 28.69 65 159 O ASN 36 144.461 46.704 31.906 1.00 65.06																			
72 O TRP 25 127.997 52.550 52.063 1.00 43.75																			
73 N GLY 26 125.958 52.862 51.172 1.00 47.80 55 146 CB ASP 35 143.895 49.419 37.558 1.00 76.13 74 CA GLY 26 126.210 54.267 50.894 1.00 39.84 147 CG ASP 35 143.288 48.547 38.638 1.00 84.32 75 C GLY 26 126.744 54.494 94.83 1.00 44.69 148 OD1 ASP 35 142.931 47.387 38.344 1.00 92.06 16.00 1.00 1.00 1.00 1.00 1.00 1.00																			
75 C GLY 26 126.744 54.49 49.483 1.00 44.69 148 OD1 ASP 35 142.931 47.387 38.344 1.00 92.06  76 O GLY 26 126.375 53.696 48.580 1.00 46.55 149 OD2 ASP 35 143.155 49.030 39.784 1.00 86.08  77 N ASP 27 127.620 55.434 49.287 1.00 46.92 150 C ASP 35 143.198 48.714 35.227 1.00 68.52  78 CA ASP 27 128.200 55.708 47.966 1.00 50.38 151 O ASP 35 143.555 47.552 35.425 1.00 66.53  80 CG ASP 27 128.544 57.196 47.827 1.00 57.61 60 152 N ASN 36 142.940 49.214 34.019 1.00 66.53  80 CG ASP 27 127.307 58.091 47.770 1.00 66.06 60 153 CA ASN 36 142.940 49.214 34.019 1.00 66.53  81 OD1 ASP 27 126.168 57.582 47.895 1.00 64.78 154 CB ASN 36 142.949 49.430 31.577 1.00 72.78  82 OD2 ASP 27 127.482 59.318 47.597 1.00 67.46 155 CG ASN 36 142.949 49.430 31.577 1.00 72.78  83 C ASP 27 129.441 54.857 47.686 1.00 46.14 156 OD1 ASN 36 140.708 50.194 31.962 1.00 78.85  84 O ASP 27 130.165 55.082 46.711 1.00 47.50 157 ND2 ASN 36 142.319 51.756 31.853 1.00 84.96  85 N GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 144.381 47.686 32.646 1.00 68.26  86 CA GLN 28 130.759 52.921 48.461 1.00 28.69 65 159 O ASN 36 144.461 46.704 31.906 1.00 65.02	73	N	GLY	26	125.958	52.862	51.172	1.00	47.80	55	146	CB	ASP	35	143.895	49.419	37.558	1.00	76.13
76 O GLY 26 126.375 53.696 48.580 1.00 46.55 149 OD2 ASP 35 143.155 49.030 39.784 1.00 86.08 77 N ASP 27 127.620 55.434 49.287 1.00 46.92 150 C ASP 35 143.198 48.714 35.227 1.00 68.52 78 CA ASP 27 128.200 55.708 47.966 1.00 50.38 151 O ASP 35 143.555 47.552 35.425 1.00 65.76 80 CG ASP 27 127.307 58.091 47.770 1.00 66.06 153 CA ASN 36 142.940 49.214 34.019 1.00 66.53 81 OD1 ASP 27 127.307 58.091 47.770 1.00 66.06 153 CA ASN 36 142.949 49.430 31.577 1.00 72.78 82 OD2 ASP 27 127.482 59.318 47.597 1.00 67.46 155 CG ASN 36 142.949 49.430 31.577 1.00 72.78 82 OD2 ASP 27 129.441 54.857 47.686 1.00 46.14 156 OD1 ASN 36 140.708 50.194 31.962 1.00 78.35 84 O ASP 27 130.165 55.082 46.711 1.00 47.50 157 ND2 ASN 36 142.319 51.756 31.853 1.00 84.96 85 N GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 144.383 47.686 32.646 1.00 68.26 86 CA GLN 28 130.759 52.921 48.461 1.00 28.69 65 159 O ASN 36 144.461 46.704 31.906 1.00 65.02																			
77         N         ASP         27         127.620         55.434         49.287         1.00         46.92         150         C         ASP         35         143.198         48.714         35.227         1.00         68.52           78         CA         ASP         27         128.200         55.708         47.966         1.00         50.38         151         O         ASP         35         143.198         48.714         35.227         1.00         68.52           79         CB         ASP         27         128.544         57.196         47.827         1.00         57.61         60         152         N         ASN         36         142.940         49.214         34.019         1.00         66.53           80         CG         ASP         27         127.307         58.091         47.770         1.00         66.06         153         CA         ASN         36         142.949         49.214         34.019         1.00         66.53           81         OD1         ASP         27         126.168         57.582         47.597         1.00         64.78         154         CB         ASN         36         142.949         49.430         31.5																			
78         CA         ASP         27         128.200         55.708         47.966         1.00         50.38         151         O         ASP         35         143.555         47.552         35.425         1.00         65.76           79         CB         ASP         27         128.544         57.196         47.827         1.00         57.61         60         152         N         ASN         36         142.940         49.214         34.019         1.00         66.53           80         CG         ASP         27         127.307         58.091         47.770         1.00         66.06         153         CA         ASN         36         143.083         48.471         32.765         1.00         66.53           81         OD1         ASP         27         126.168         57.582         47.895         1.00         64.78         154         CB         ASN         36         142.949         49.430         31.577         1.00         72.78           82         OD2         ASP         27         129.441         54.857         47.686         1.00         46.14         156         OD1         ASN         36         141.889         50.497																			
80 CG ASP 27 127.307 58.091 47.770 1.00 66.06 60 153 CA ASN 36 143.083 48.471 32.765 1.00 67.50 81 OD1 ASP 27 126.168 57.582 47.895 1.00 64.78 154 CB ASN 36 142.949 49.430 31.577 1.00 72.78 82 OD2 ASP 27 127.482 59.318 47.597 1.00 67.46 155 CG ASN 36 141.889 50.497 31.804 1.00 79.82 83 C ASP 27 129.441 54.857 47.686 1.00 46.14 156 OD1 ASN 36 140.708 50.194 31.962 1.00 78.35 84 O ASP 27 130.165 55.082 46.711 1.00 47.50 157 ND2 ASN 36 142.319 51.756 31.853 1.00 84.96 85 N GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 144.383 47.686 32.646 1.00 68.26 86 CA GLN 28 130.759 52.921 48.461 1.00 28.69 65 159 O ASN 36 144.461 46.704 31.906 1.00 65.02	78	CA	ASP	27	128.200	55.708		1.00	50.38		151	O	ASP	35	143.555	47.552	35.425	1.00	65.76
81 OD1 ASP 27 126.168 57.582 47.895 1.00 64.78 154 CB ASN 36 142.949 49.430 31.577 1.00 72.78 82 OD2 ASP 27 126.148 59.318 47.597 1.00 67.46 155 CG ASN 36 141.889 50.497 31.804 1.00 79.82 83 C ASP 27 129.441 54.857 47.686 1.00 46.14 156 OD1 ASN 36 140.708 50.194 31.962 1.00 78.35 84 O ASP 27 130.165 55.082 46.711 1.00 47.50 157 ND2 ASN 36 142.319 51.756 31.853 1.00 84.96 85 N GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 144.833 47.686 32.646 1.00 68.26 86 CA GLN 28 130.759 52.921 48.461 1.00 28.69 65 159 O ASN 36 144.461 46.704 31.906 1.00 65.02										60									
82 OD2 ASP 27 127.482 59.318 47.597 1.00 67.46 155 CG ASN 36 141.889 50.497 31.804 1.00 79.82 83 C ASP 27 129.441 54.857 47.686 1.00 46.14 156 OD1 ASN 36 140.708 50.194 31.962 1.00 78.35 84 O ASP 27 130.165 55.082 46.711 1.00 47.50 157 ND2 ASN 36 142.319 51.756 31.853 1.00 84.96 85 N GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 144.383 47.686 32.646 1.00 68.26 86 CA GLN 28 130.759 52.921 48.461 1.00 28.69 65 159 O ASN 36 144.461 46.704 31.906 1.00 65.02																			
83 C ASP 27 129.441 54.857 47.686 1.00 46.14 156 OD1 ASN 36 140.708 50.194 31.962 1.00 78.35 84 O ASP 27 130.165 55.082 46.711 1.00 47.50 157 ND2 ASN 36 142.319 51.756 31.853 1.00 84.96 85 N GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 144.383 47.686 32.646 1.00 68.26 86 CA GLN 28 130.759 52.921 48.461 1.00 28.69 65 159 O ASN 36 144.461 46.704 31.906 1.00 65.02																			
85 N GLN 28 129.642 53.855 48.536 1.00 40.05 158 C ASN 36 144.383 47.686 32.646 1.00 68.26 86 CA GLN 28 130.759 52.921 48.461 1.00 28.69 65 159 O ASN 36 144.461 46.704 31.906 1.00 65.02	83	С	ASP	27	129.441	54.857	47.686	1.00	46.14		156	OD1	ASN	36	140.708	50.194	31.962	1.00	78.35
86 CA GLN 28 130.759 52.921 48.461 1.00 28.69 <sup>65</sup> 159 O ASN 36 144.461 46.704 31.906 1.00 65.02																			
										65									

TABLE 10-continued

Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound									5	s	Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound							
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
161	CA	GLN	37	146.709	47.500	33.370	1.00	71.18		234	N	ILE	46	145.415	34.225	32.299	1.00	34.27
162	CB	GLN	37	147.721	48.431	34.048	1.00	78.38		235	CA	ILE	46	144.239	33.358	32.373	1.00	30.12
163 164	CG CD	GLN GLN	37 37	149.005 149.198	47.761 47.904	34.524 36.027	$\frac{1.00}{1.00}$	90.52 100.00	10	236 237	CB CG2	ILE ILE	46 46	142.942 141.706	34.181 33.420	32.608 32.123	1.00 1.00	33.74 32.63
165	OE1	GLN	37	148.538	48.718	36.673	1.00	100.00		238	CG2	ILE	46	142.812	34.534	34.093	1.00	34.26
166	NE2	GLN	37	150.106	47.105	36.592	1.00	100.00		239	CD1	ILE	46	141.644	35.444	34.407	1.00	30.01
167	C	GLN	37	146.651	46.131	34.069	1.00	65.44		240	C	ILE	46	144.099	32.518	31.110	1.00	29.77
168 169	O N	GLN VAL	37 38	147.138 146.023	45.138 46.086	33.533 35.244	1.00 $1.00$	63.18 57.92		241 242	O N	ILE GLU	46 47	143.850 144.283	31.315 33.156	31.186 29.956	1.00 1.00	30.24 31.42
170	CA	VAL	38	145.883	44.849	36.021	1.00	51.76	15	243	CA	GLU	47	144.185	32.482	28.666	1.00	37.34
171	CB	VAL	38	145.388	45.152	37.461	1.00	50.39		244	CB	GLU	47	144.460	33.476	27.537	1.00	45.72
172 173	CG1 CG2	VAL VAL	38 38	145.198 146.371	43.862 46.071	38.251 38.166	1.00 $1.00$	44.02 43.36		245 246	CG CD	GLU GLU	47 47	144.290 144.808	32.896 33.813	26.137 25.035	1.00 1.00	68.61 79.62
173	C	VAL	38	144.916	43.870	35.349	1.00	52.33		247	OE1	GLU	47	145.302	34.922	25.339	1.00	89.55
175	O	VAL	38	145.142	42.656	35.348	1.00	48.70	20	248	OE2	GLU	47	144.728	33.417	23.852	1.00	86.41
176	N	ALA	39	143.858	44.412	34.752	1.00	49.41	20	249	С	GLU	47	145.169	31.314	28.580	1.00	37.12
177 178	CA CB	ALA ALA	39 39	142.848 141.722	43.610 44.502	34.068 33.584	1.00 $1.00$	48.03 56.98		250 251	O N	GLU ALA	47 48	144.860 146.348	30.275 31.492	27.997 29.171	1.00 1.00	45.60 33.83
179	C	ALA	39	143.434	42.823	32.900	1.00	47.68		252	CA	ALA	48	147.378	30.459	29.170	1.00	30.76
180	0	ALA	39	143.178	41.627	32.759	1.00	52.03		253	CB	ALA	48	148.720	31.054	29.575	1.00	33.78
181 182	N CA	GLU GLU	40 40	144.219 144.855	43.501 42.881	32.068 30.908	1.00 $1.00$	46.51 40.96	25	254 255	C	ALA ALA	48 48	146.986 147.071	29.323 28.150	30.110 29.743	1.00 1.00	30.08 30.80
183	CB	GLU	40	145.507	43.952	30.036	1.00	49.36		256	N	LEU	49	143.542	29.685	31.312	1.00	27.01
184	CG	GLU	40	144.507	44.896	29.383	1.00	62.86		257	CA	LEU	49	146.110	28.720	32.321	1.00	23.68
185	CD	GLU	40	145.161	46.109	28.745	1.00	67.78		258	CB	LEU	49	145.793	29.445	33.628	1.00	19.74
186 187	OE1 OE2	GLU GLU	40 40	146.229 144.601	45.957 47.218	28.112 28.880	$\frac{1.00}{1.00}$	67.66 70.01		259 260	CG CD1	LEU LEU	49 49	146.936 146.368	30.167 31.161	34.337 35.328	$\frac{1.00}{1.00}$	15.12 8.67
188	C	GLU	40	145.893	41.852	31.337	1.00	38.90	30	261	CD2	LEU	49	147.844	29.164	35.033	1.00	12.64
189	O	GLU	40	146.076	40.832	30.678	1.00	39.36		262	C	LEU	49	144.862	27.985	31.836	1.00	24.27
190 191	N CA	LYS LYS	41 41	146.569 147.584	42.135 41.243	32.447 32.998	$\frac{1.00}{1.00}$	41.55 38.43		263 264	O N	LEU LYS	49 50	144.610 144.101	26.842 28.663	32.214 30.983	$\frac{1.00}{1.00}$	31.26 27.77
192	CB	LYS	41	148.219	41.884	34.238	1.00	43.42		265	CA	LYS	50	142.863	28.154	30.394	1.00	30.88
193	CG	LYS	41	149.304	41.056	34.903	1.00	55.00		266	CB	LYS	50	142.247	29.263	29.548	1.00	31.45
194 195	CD	LYS	41	149.864	41.780	36.119	1.00	61.88	35	267	CG	LYS	50	140.775	29.153	29.242	1.00	31.65
193	CE NZ	LYS LYS	41 41	151.040 151.665	41.028 41.794	36.721 37.835	1.00 $1.00$	62.99 69.92		268 269	CD CE	LYS LYS	50 50	140.333 138.871	30.468 30.468	28.621 28.250	1.00 1.00	33.98 42.53
197	C	LYS	41	146.914	39.926	33.373	1.00	36.30		270	NZ	LYS	50	138.455	31.817	27.773	1.00	47.24
198	0	LYS	41	147.362	38.855	32.966	1.00	34.80		271	C	LYS	50	143.120	26.925	29.527	1.00	32.84
199 200	N CA	TYR TYR	42 42	145.823 145.051	40.027 38.868	34.132 34.572	1.00 $1.00$	35.61 29.43		272 273	O N	LYS GLU	50 51	142.449 144.092	25.901 27.033	29.675 28.625	1.00 1.00	31.46 33.57
201	CB	TYR	42	143.880	39.307	35.457	1.00	29.64	40	274	CA	GLU	51	144.439	25.927	27.741	1.00	38.44
202	CG	TYR	42	144.229	39.658	36.890	1.00	30.55		275	CB	GLU	51	145.286	26.416	26.566	1.00	45.31
203 204	CD1 CE1	TYR TYR	42 42	145.556 145.866	39.697 40.002	37.330 38.660	1.00 $1.00$	37.21 37.30		276 277	CG CD	GLU GLU	51 51	145.241 143.953	25.501 25.633	25.339 24.532	1.00 1.00	54.99 61.58
205	CD2	TYR	42	143.222	39.937	37.814	1.00	28.62		278	OE1	GLU	51	143.086	26.463	24.893	1.00	63.41
206	CE2	TYR	42	143.519	40.241	39.139	1.00	37.07		279	OE2	GLU	51	143.815	24.912	23.519	1.00	63.99
207	CZ	TYR	42	144.839	40.272	39.556	1.00	40.82	45	280	С	GLU	51	145.179	24.824	28.501	1.00	37.86
208 209	OH C	TYR TYR	42 42	145.121 144.499	40.567 38.097	40.869 33.377	1.00 $1.00$	43.81 30.40		281 282	O N	GLU GLN	51 52	145.145 145.867	23.662 25.192	28.097 29.582	1.00 1.00	43.06 36.20
210	O	TYR	42	144.603	36.872	33.318	1.00	29.10		283	CA	GLN	52	146.592	24.212	30.397	1.00	38.10
211	N	ALA	43	143.920	38.827	32.426	1.00	24.33		284	CB	GLN	52	147.453	24.897	31.465	1.00	41.72
212 213	CA CB	ALA ALA	43 43	143.340 142.713	38.227 39.308	31.227 30.356	$\frac{1.00}{1.00}$	29.09 19.04	50	285 286	CG CD	GLN GLN	52 52	148.691 149.505	25.615 26.249	30.943 32.061	$\frac{1.00}{1.00}$	47.34 49.46
214	C	ALA	43	144.358	37.423	30.421	1.00	29.23		287	OE1	GLN	52	149.640	25.683	33.145	1.00	43.98
215	O	ALA	43	144.074	36.308	29.984	1.00	29.14		288	NE2	GLN	52	150.049	27.438	31.799	1.00	53.90
216 217	N CA	LYS LYS	44 44	145.559 146.637	37.972 37.371	30.260 29.491	$\frac{1.00}{1.00}$	37.81 40.64		289 290	C O	GLN GLN	52 52	145.563 145.732	23.339 22.122	31.093 31.219	$\frac{1.00}{1.00}$	38.87 40.47
218	C	LYS	44	147.069	36.041	30.095	1.00	34.80		291	N	THR	53	144.501	23.991	31.554	1.00	37.87
219	O	LYS	44	147.221	35.048	29.357	1.00	35.25	55	292	CA	THR	53	143.407	23.323	32.236	1.00	32.77
220	CB	LYS	44	147.824	38.329	29.396	1.00	54.18		293	CB OG1	THR	53 53	142.541	24.347	32.992	1.00	31.08
221 222	CG CD	LYS LYS	44 44	149.001 150.141	37.784 38.787	28.605 28.552	1.00 $1.00$	67.03 79.04		294 295	OG1 CG2	THR THR	53 53	143.315 141.296	24.933 23.685	34.050 33.569	1.00 1.00	28.04 32.90
223	CE	LYS	44	151.313	38.247	27.750	1.00	20.00		296	C	THR	53	142.570	22.522	31.233	1.00	32.05
224	NZ	LYS	44	152.431	39.227	27.673	1.00	20.00		297	O	THR	53	142.013	21.476	31.573	1.00	29.73
225 226	N CA	GLU GLU	45 45	147.332 147.771	36.000 34.779	31.397 32.070	1.00 $1.00$	31.47 30.36	60	298 299	N CA	ARG ARG	54 54	142.529 141.785	22.992 22.312	29.988 28.933	1.00 1.00	28.01 23.69
227	CB	GLU	45	148.288	35.080	33.480	1.00	26.58		300	СВ	ARG	54	141.723	23.176	27.673	1.00	23.31
228	CG	GLU	45	149.071	33.920	34.105	1.00	19.97		301	CG	ARG	54	140.724	22.682	26.633	1.00	24.23
229 230	CD OE1	GLU GLU	45 45	149.394 149.791	34.128 33.146	35.580 36.246	$\frac{1.00}{1.00}$	33.35 31.26		302 303	CD NE	ARG ARG	54 54	140.755 140.674	23.527 24.969	25.360 25.619	1.00 1.00	30.78 45.26
231	OE2	GLU	45	149.249	35.264	36.080	1.00	37.21		304	CZ	ARG	54	139.564	25.633	25.942	1.00	46.57
232	C	GLU	45	146.649	33.747	32.142	1.00	31.64	65	305	NH1	ARG	54	138.405	24.999	26.058	1.00	51.75
233	О	GLU	45	146.902	32.545	32.058	1.00	38.67		306	NH2	ARG	54	139.608	26.946	26.140	1.00	43.00

TABLE 10-continued

s	Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound								5	Structural Coordinates of Tobacco 5-Epi-Aristolochen With Farnesyl Hydroxyphosphonate Bound								hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
307	С	ARG	54	142.487	20.998	28.617	1.00	30.27		380	С	LEU	64	137.184	8.363	37.778	1.00	32.66
308	O	ARG	54	141.842	19.955	28.479	1.00	33.41		381	O	LEU	64	136.773	9.515	37.650	1.00	32.91
309	N	ASN	55	143.821	21.050	28.526	1.00	33.72	10	382	N	ALA	65	137.664	7.881	38.923	1.00	26.30
310	CA	ASN	55	144.648	19.899	28.240 29.348	1.00	33.22 35.28		383	CA	ALA	65 65	137.721	8.683	40.141 41.265	1.00	27.29 26.12
311 312	C O	ASN ASN	55 55	144.538 144.679	18.872 17.660	29.348	1.00 $1.00$	35.20 35.19		384 385	CB C	ALA ALA	65 65	138.362 138.482	7.885 9.988	39.919	1.00 1.00	33.96
313	CB	ASN	55	146.080	20.341	27.963	1.00	36.29		386	o	ALA	65	138.019	11.057	40.318	1.00	35.01
314	CG	ASN	55	146.150	21.264	26.761	1.00	20.00		387	N	ASP	66	139.630	9.897	39.250	1.00	35.46
315	OD1	ASN	55	145.473	21.038	25.754	1.00	20.00	15	388	CA	ASP	66	140.459	11.064	38.961	1.00	35.10
316	ND2	ASN	55 56	146.963	22.307 19.330	26.857	1.00 $1.00$	20.00		389 390	CB	ASP	66	141.776 142.685	10.646	38.298 39.229	1.00	36.60
317 318	N CA	MET MET	56 56	144.309 144.150	18.442	30.581 31.734	1.00	34.89 34.60		391	CG OD1	ASP ASP	66 66	142.633	9.867 10.067	40.461	1.00 1.00	34.65 25.86
319	СВ	MET	56	144.058	19.241	33.039	1.00	27.26		392	OD2	ASP	66	143.488	9.057	38.717	1.00	44.77
320	CG	MET	56	145.378	19.792	33.544	1.00	38.81		393	C	ASP	66	139.746	12.065	38.059	1.00	31.14
321	SD	MET	56	145.237	20.594	35.159	1.00	40.35	20	394	O	ASP	66	139.846	13.276	38.266	1.00	31.51
322 323	CE C	MET MET	56 56	145.790 142.880	22.242 17.606	34.734 31.560	1.00 $1.00$	41.02 38.38		395 396	N CA	THR THR	67 67	139.045 138.316	11.552 12.392	37.051 36.105	1.00 1.00	26.69 26.40
323	Ö	MET	56	142.871	16.406	31.847	1.00	36.10		397	CB	THR	67	137.793	11.571	34.918	1.00	25.70
325	N	LEU	57	141.816	18.253	31.084	1.00	33.14		398	OG1	THR	67	138.891	10.917	34.270	1.00	27.33
326	CA	LEU	57	140.535	17.593	30.852	1.00	33.61		399	CG2	THR	67	137.095	12.474	33.918	1.00	28.90
327	CB	LEU	57	139.444	18.633	30.566	1.00	24.13	25	400	С	THR	67	137.146	13.113	36.769	1.00	24.73
328 329	CG CD1	LEU LEU	57 57	138.939 138.092	19.472 20.624	31.742 31.235	$\frac{1.00}{1.00}$	25.14 18.69	23	401 402	O N	THR LEU	61 68	136.899 136.425	14.290 12.401	36.502 37.629	1.00 1.00	27.41 23.13
330	CD2	LEU	57	138.143	18.604	32.703	1.00	10.16		403	CA	LEU	68	135.295	12.985	38.333	1.00	18.00
331	С	LEU	57	140.610	16.611	29.686	1.00	37.09		404	СВ	LEU	68	134.504	11.909	39.078	1.00	13.23
332	O	LEU	57	139.922	15.588	29.679	1.00	34.95		405	CG	LEU	68	133.804	10.871	38.201	1.00	16.24
333	N	LEU	58	141.453	16.924	28.703	1.00	35.34	20	406	CD1	LEU	68	133.109	9.843	39.078	1.00	17.55
334 335	CA CB	LEU LEU	58 58	141.605 141.930	16.071 16.926	27.533 26.304	$\frac{1.00}{1.00}$	35.75 33.37	30	407 408	CD2 C	LEU LEU	68 68	132.811 135.787	11.552 14.047	37.272 39.305	1.00 1.00	7.96 19.40
336	CG	LEU	58	140.886	17.987	25.951	1.00	36.57		409	ŏ	LEU	68	135.181	15.113	39.420	1.00	21.96
337	CD1	LEU	58	141.334	18.779	24.736	1.00	34.54		410	N	ASN	69	136.899	13.766	39.980	1.00	17.72
338	CD2	LEU	58	139.540	17.333	25.691	1.00	38.57		411	CA	ASN	69	137.471	14.714	40.931	1.00	23.29
339	С	LEU	58	142.628	14.946	27.688	1.00	40.17		412	CB	ASN	69	138.608	14.071	41.728	1.00	27.45
340 341	O N	LEU ALA	58 59	143.001 143.066	14.298 14.697	26.710 28.922	$\frac{1.00}{1.00}$	38.69 45.53	35	413 414	CG OD1	ASN ASN	69 69	138.102 137.171	13.102 13.413	42.783 43.530	1.00 1.00	44.15 45.05
342	CA	ALA	59	144.038	13.637	29.198	1.00	52.73		415	ND2	ASN	69	138.709	11.921	42.846	1.00	48.60
343	CB	ALA	59	144.562	13.754	30.626	1.00	52.29		416	С	ASN	69	137.954	15.985	40.240	1.00	21.73
344	С	ALA	59	143.402	12.263	28.950	1.00	60.58		417	O	ASN	69	137.784	17.083	40.764	1.00	19.56
345 346	O N	ALA THR	59 60	142.320 144.084	11.962 11.432	29.450 28.168	1.00 $1.00$	62.17 63.55		418 419	N CA	LEU LEU	70 70	138.526 139.012	15.834 16.979	39.050 38.293	1.00 1.00	19.22 18.06
347	CA	THR	60	143.575	10.109	27.794	1.00	63.50	40	420	CB	LEU	70	139.736	16.522	37.025	1.00	14.83
348	CB	THR	60	144.405	9.528	26.641	1.00	63.61		421	CG	LEU	70	140.274	17.650	36.141	1.00	18.22
349	OG1	THR	60	145.776	9.434	27.039	1.00	67.85		422	CD1	LEU	70	141.330	18.446	36.889	1.00	15.05
350	CG2	THR	60	144.302	10.426	25.420	1.00	59.57		423	CD2	LEU	70	140.845	17.078	34.862	1.00	20.94
351 352	C	THR THR	60 60	143.372 142.237	9.004 8.681	28.844 29.198	1.00 $1.00$	64.51 69.94		424 425	C	LEU LEU	70 70	137.835 137.844	17.871 19.069	37.925 38.212	1.00 1.00	20.53 23.73
353	N	GLY	61	144.470	8.435	29.337	1.00	59.72	45	426	Ň	ILE	71	136.817	17.269	37.312	1.00	19.75
354	CA	GLY	61	144.394	7.339	30.294	1.00	59.70		427	CA	ILE	71	135.613	17.986	36.901	1.00	16.33
355	С	GLY	61	144.087	7.572	31.767	1.00	60.15		428	CB	ILE	71	134.617	17.043	36.184	1.00	19.20
356 357	O N	GLY MET	61 62	144.627 143.233	6.862 8.546	32.620 32.098	$\frac{1.00}{1.00}$	62.75 62.09		429 430	CG2 CG1	ILE ILE	71 71	133.278 135.216	17.744 16.574	35.950 34.856	$\frac{1.00}{1.00}$	17.41 25.29
358	CA	MET	62	142.904	8.860	33.482	1.00	62.20		431	CD1	ILE	71	134.273	15.748	34.009	1.00	23.29
359	C	MET	62	141.787	7.962	34.001	1.00	57.87	50	432	С	ILE	71	134.927	18.661	38.088	1.00	15.45
360	O	MET	62	140.858	7.592	33.304	1.00	60.93		433	0	ILE	71	134.507	19.813	37.991	1.00	10.60
361	CB	MET	62	142.512	10.333	33.619	1.00	65.78		434	N	ASP	72 72	134.846	17.952	39.212	1.00	14.16
362 363	CG SD	MET MET	62 62	142.374 141.924	10.806 12.547	35.057 35.177	$\frac{1.00}{1.00}$	71.62 20.00		435 436	CA CB	ASP ASP	72 72	134.222 134.206	18.477 17.400	40.425 41.516	1.00 1.00	13.13 9.28
364	CE	MET	62	141.630	12.940	33.455	1.00	20.00		437	CG	ASP	72	133.456	17.828	42.766	1.00	8.19
365	N	LYS	63	141.877	7.621	35.308	1.00	53.67	55	438	OD1	ASP	72	132.472	18.591	42.660	1.00	19.56
366	CA	LYS	63	140.886	6.787	35.975	1.00	51.44		439	OD2	ASP	72	133.842	17.381	43.865	1.00	24.87
367 368	OB	LYS	63	141.401	6.323	37.342	1.00	55.26 65.11		440 441	C	ASP	72 72	134.969	19.709	40.926	1.00	20.93
368 369	CG CD	LYS LYS	63 63	142.607 142.968	5.402 4.915	37.287 38.682	$\frac{1.00}{1.00}$	65.11 69.24		441 442	O N	ASP THR	72 73	134.357 136.298	20.734 19.618	41.230 40.980	1.00 1.00	31.87 26.21
370	CE	LYS	63	144.127	3.936	38.635	1.00	76.58		443	CA	THR	73	137.162	20.689	41.452	1.00	17.72
371	NZ	LYS	63	144.434	3.376	39.980	1.00	78.96	60	444	C	THR	73	137.051	21.912	40.558	1.00	16.99
372	С	LYS	63	139.576	7.543	36.173	1.00	48.71	00	445	O	THR	73	136.913	23.046	41.054	1.00	15.04
373 374	O N	LYS LEU	63 64	139.559 138.490	8.778 6.802	36.167 36.386	$\frac{1.00}{1.00}$	48.30 44.07		446 447	CB OG1	THR THR	73 73	138.627 138.771	20.258 19.152	41.508 42.406	1.00 1.00	18.68 20.00
375	CA	LEU	64	137.182	7.413	36.586	1.00	38.53		448	CG2	THR	73 73	139.503	21.413	41.971	1.00	23.27
376	CB	LEU	64	136.100	6.343	36.778	1.00	40.14		449	N	ILE	74	137.124	21.732	39.245	1.00	13.35
377	CG	LEU	64	134.671	6.886	36.899	1.00	35.60	<i>c=</i>	450	CA	ILE	74	137.024	22.837	38.298	1.00	15.24
378 379	CD1	LEU	64	134.283	7.589	35.606	1.00	32.53	65	451	CB	ILE	74 74	137.214	22.342	36.844	1.00	17.48
319	CD2	LEU	64	133.689	5.773	37.203	1.00	30.12		452	CG2	ILE	74	136.841	23.434	35.843	1.00	10.21

TABLE 10-continued

Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound									. 5	s	Structural Coordinates of Tobacco 5-Epi-Aristolochene Syntha With Farnesyl Hydroxyphosphonate Bound							
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	occ	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
453	CG1	ILE	74 138.	.658	21.881	36.637	1.00	16.00		526	CA	PHE	83	134.774	24.241	29.475	1.00	32.99
454	CD1	ILE	74 138.		21.335	35.253	1.00	19.68	10	527	CB	PHE	83	135.829	23.389	30.191	1.00	38.74
455 456	C O	ILE ILE	74 135. 74 135.		23.554 24.774	38.431 38.285	$\frac{1.00}{1.00}$	21.97 37.27	10	528 529	CG CD1	PHE PHE	83 83	137.052 137.204	24.157 24.583	30.603 31.921	1.00 1.00	40.58 40.06
457	N	GLU	75 134.		22.793	38.712	1.00	24.94		530	CD2	PHE	83	138.041	24.476	29.675	1.00	41.50
458	CA	GLU	75 133.		23.362	38.869	1.00	17.86		531	CE1	PHE	83	138.320	25.318	32.309	1.00	43.27
459 460	CB CG	GLU GLU	75 132. 75 131.		22.266 21.565	38.893 37.557	1.00 $1.00$	22.45 23.19		532 533	CE2 CZ	PHE PHE	83 83	139.163 139.303	25.211 25.634	30.050 31.371	1.00 1.00	38.18 46.92
461	CD	GLU	75 131. 75 130.		20.685	37.539	1.00	24.24	15	534	C	PHE	83	133.444	23.496	29.471	1.00	32.87
462	OE1	GLU	75 130.		19.984	38.540	1.00	11.68	13	535	O	PHE	83	133.378	22.340	29.886	1.00	31.33
463	OE2	GLU	75 130.		20.699	36.513	1.00	19.29		536	N	GLU	84	132.397	24.133	28.960	1.00	33.76
464 465	C O	GLU GLU	75 133. 75 132.		24.181 25.323	40.142 40.119	1.00 $1.00$	16.49 18.93		537 538	CA CB	GLU GLU	84 84	131.086 129.991	23.496 24.514	28.929 28.601	1.00 1.00	38.03 47.83
466	N	ARG	76 133.		23.590	41.248	1.00	11.73		539	CG	GLU	84	129.901	25.690	29.578	1.00	58.57
467	CA	ARG	76 133.		24.248	42.552	1.00	15.45	20	540	CD	GLU	84	129.403	25.312	30.975	1.00	69.66
468 469	CB CG	ARG ARG	76 134. 76 133.		23.282 22.097	43.636 43.899	1.00 $1.00$	7.10 15.61		541 542	OE1 OE2	GLU GLU	84 84	129.719 128.695	24.208 26.146	31.479 31.586	1.00 1.00	74.44 66.21
470	CD	ARG	76 133.		21.197	44.975	1.00	12.16		543	C	GLU	84	131.030	22.314	27.968	1.00	39.30
471	NE	ARG	76 132.	.824	20.231	45.508	1.00	16.00		544	O	GLU	84	130.339	21.328	28.228	1.00	37.93
472	CZ	ARG	76 132.		20.165	46.789	1.00	19.23		545 546	N	LYS	85	131.780	22.401	26.872	1.00	37.65
473 474	NH1 NH2	ARG ARG	76 132. 76 131.		21.010 19.234	47.670 47.202	$\frac{1.00}{1.00}$	26.80 29.06	25	546 547	CA CB	LYS LYS	85 85	131.815 132.367	21.329 21.839	25.886 24.551	1.00 1.00	40.19 49.51
475	C	ARG	76 134.		25.519	42.564	1.00	20.45		548	CG	LYS	85	132.443	20.770	23.469	1.00	57.11
476	0	ARG	76 134.		26.454	43.319	1.00	19.47		549	CD	LYS	85	133.176	21.261	22.237	1.00	73.34
477 478	N CA	LEU LEU	77 135. 77 136.		25.539 26.692	41.732 41.634	$\frac{1.00}{1.00}$	20.12 19.40		550 551	CE NZ	LYS LYS	85 85	133.280 134.029	20.160 20.532	21.180 19.952	1.00 1.00	81.58 94.03
479	CB	LEU	77 137.		26.281	41.014	1.00	12.91		552	C	LYS	85	132.661	20.161	26.381	1.00	37.98
480	CG	LEU	77 138.	.678	25.382	41.843	1.00	12.12	30	553	O	LYS	85	132.200	19.018	26.404	1.00	43.86
481	CD1	LEU	77 139.		24.903	40.973 43.070	1.00	2.00		554	N	GLU	86	133.894	20.461	26.784	1.00	36.06
482 483	CD2 C	LEU LEU	77 139. 77 135.		26.125 27.823	40.818	$\frac{1.00}{1.00}$	4.07 22.31		555 556	CA CB	GLU GLU	86 86	134.825 136.122	19.448 20.102	27.277 27.774	1.00 1.00	32.72 36.71
484	O	LEU	77 136.		28.906	40.702	1.00	30.09		557	CG	GLU	86	136.998	20.743	26.698	1.00	42.45
485	N	GLY	78 134.		27.551	40.238	1.00	26.16		558	CD	GLU	86	136.500	22.103	26.219	1.00	44.54
486 487	CA C	GLY GLY	78 133. 78 134.		28.542 28.855	39.447 38.082	$\frac{1.00}{1.00}$	20.26 16.66	35	559 560	OE1 OE2	GLU GLU	86 86	135.646 136.977	22.720 22.566	26.891 25.162	1.00 1.00	49.68 47.77
488	o	GLY	78 134.		29.898	37.504	1.00	19.22		561	C	GLU	86	134.213	18.618	28.402	1.00	29.96
489	N	ILE	79 135.		27.959	37.553	1.00	13.23		562	0	GLU	86	134.254	17.389	28.370	1.00	32.06
490 491	CA CB	ILE ILE	79 135. 79 137.		28.179 28.083	36.247 36.321	1.00 $1.00$	16.00 14.32		563 564	N CA	ILE ILE	87 87	133.638 133.013	19.303 18.648	29.388 30.534	1.00 1.00	28.21 27.11
492	CG2	ILE	79 138.		29.257	37.111	1.00	9.65	40	565	CB	ILE	87	132.618	19.672	31.617	1.00	28.37
493	CG1	ILE	79 137.		26.751	36.944	1.00	15.84	40	566	CG2	ILE	87	131.813	18.996	32.729	1.00	28.34
494 495	CD1 C	ILE ILE	79 139. 79 135.		26.574	37.082 35.185	1.00 $1.00$	20.69 17.13		567 568	CG1 CD1	ILE	87 87	133.880 133.613	20.338 21.386	32.179 33.241	1.00 1.00	22.12 21.16
493	Ö	ILE	79 135.		27.216 27.256	34.033	1.00	20.55		569	CD1	ILE ILE	87	133.013	17.815	30.150	1.00	27.00
497	N	SER	80 134.		26.377	35.567	1.00	21.41		570	Ö	ILE	87	131.581	16.735	30.700	1.00	29.31
498	CA	SER	80 133.		25.392	34.654	1.00	23.76	45	571	N	ASP	88	131.007	18.309	29.200	1.00	31.52
499 500	CB OG	SER SER	80 133. 80 131		24.393 25.047	35.419 36.170	$\frac{1.00}{1.00}$	20.88 23.54	73	572 573	CA CB	ASP ASP	88 88	129.815 129.009	17.593 18.445	28.751 27.764	1.00 1.00	39.20 40.43
501	C	SER	80 133.		25.977	33.485	1.00	20.44		574	CG	ASP	88	127.717		27.330	1.00	36.63
502	0	SER	80 132.		25.280	32.505	1.00	28.56		575	OD1	ASP	88	126.845		28.194	1.00	36.19
503 504	N CA	TYR TYR	81 132. 81 131.		27.252 27.907	33.577 32.507	$\frac{1.00}{1.00}$	18.39 19.66		576 577	OD2 C	ASP ASP	88 88	127.577 130.173	17.470 16.253	26.125 28.107	1.00 1.00	42.80 40.12
505	CB	TYR	81 131.		29.244	32.986	1.00	10.58	50	578	Ö	ASP	88	129.660	15.210	28.513	1.00	41.11
506	CG	TYR	81 132.	.396	30.362	33.170	1.00	19.55		579	N	ASP	89	131.088	16.251	27.130	1.00	37.03
507 508	CD1 CE1	TYR TYR	81 132. 81 133.		31.285 32.331	32.151 32.320	$\frac{1.00}{1.00}$	26.56 23.18		580 581	CA C	ASP ASP	89 89	131.496 132.107	15.096 14.047	26.397 27.304	$\frac{1.00}{1.00}$	33.87 33.85
509	CD2	TYR	81 133.		30.513	34.367	1.00	12.72		582	Ö	ASP	89	132.107	12.832	26.978	1.00	33.37
510	CE2	TYR	81 133.	.998	31.555	34.546	1.00	18.75		583	CB	ASP	89	132.464	15.574	25.325	1.00	27.32
511 512	CZ	TYR	81 134.		32.460	33.519	1.00	21.59	55	584 585	CG OD1	ASP	89 80	131.779	16.561	24.391	1.00	34.13
512 513	OH C	TYR TYR	81 135. 81 132.		33.487 28.103	33.688 31.240	$\frac{1.00}{1.00}$	23.90 23.33		585 586	OD1 OD2	ASP ASP	89 89	130.528 132.481	16.553 17.345	24.341 23.721	$\frac{1.00}{1.00}$	20.00 20.00
514	O	TYR	81 132.	.294	28.498	30.195	1.00	26.36		587	N	ILE	90	132.765	14.453	28.372	1.00	31.52
515	N	HIS	82 134.		27.835	31.352	1.00	27.33		588	CA	ILE	90	133.385	13.527	29.308	1.00	25.50
516 517	CA CB	HIS HIS	82 135. 82 136.		27.955 28.227	30.229 30.724	$\frac{1.00}{1.00}$	28.77 20.70		589 590	CB CG2	ILE ILE	90 90	134.370 134.861	14.245 13.285	30.258 31.328	1.00 1.00	22.28 22.91
518	CG	HIS	82 136.		29.592	31.301	1.00	18.33	60	591	CG1	ILE	90	135.549	14.818	29.465	1.00	31.26
519	CD2	HIS	82 137.	.002	29.989	32.553	1.00	7.90		592	CD1	ILE	90	136.543	15.620	30.301	1.00	31.23
520 521	ND1 CE1	HIS HIS	82 136. 82 136.		30.742 31.788	30.548 31.312	1.00 $1.00$	14.62 11.77		593 594	C	ILE ILE	90 90	132.297 132.331	12.836 11.620	30.124 30.316	1.00 1.00	23.61 27.20
522	NE2	HIS	82 130.		31.766	32.533	1.00	13.85		595	N	LEU	91	131.331	13.622	30.593	1.00	23.72
523	C	HIS	82 135.	.085	26.654	29.440	1.00	28.78	65	596	CA	LEU	91	130.218	13.099	31.379	1.00	20.80
524 525	O	HIS	82 135.		26.643	28.265	1.00	31.36	65	597 508	CB	LEU	91	129.469	14.238	32.071	1.00	20.19
525	N	PHE	83 134.	.719	25.557	30.098	1.00	30.57		598	CG	LEU	91	130.232	14.888	33.225	1.00	17.15

TABLE 10-continued

s	Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound									s	Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound							
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor	5	Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
599	CD1	LEU	91	129.464	16.078	33.772	1.00	13.18		672	CA	ASN	100	131.294	0.473	37.995	1.00	100.00
600	CD2	LEU	91	130.479	13.852	34.318	1.00	9.86	4.0	673	CB	ASN	100	130.733	-0.892	38.382	1.00	97.11
601 602	C O	LEU LEU	91 91	129.270 128.649	12.281 11.334	30.510 30.988	1.00 1.00	22.03 22.93	10	674 675	CG OD1	ASN ASN	100 100	129.297 128.429	-1.056 -0.279	37.956 38.360	1.00 1.00	95.75 86.86
603	N	ASP	92	129.183	12.640	29.231	1.00	21.86		676	ND2	ASN	100	129.028	-2.069	37.139	1.00	94.51
604	CA	ASP	92	128.337	11.930	28.276	1.00	23.86		677	С	ASN	100	132.513	0.784	38.857	1.00	100.00
605 606	CB CG	ASP ASP	92 92	128.314 127.282	12.668 12.105	26.933 25.973	1.00 1.00	30.84 37.16		678 679	O N	ASN CYS	100 101	133.196 132.829	-0.133 2.068	39.303 39.047	1.00 1.00	100.00 98.98
607	OD1	ASP	92	126.182	12.690	25.879	1.00	34.52	15	680	CA	CYS	101	133.942	2.429	39.953	1.00	94.29
608	OD2	ASP	92	127.568	11.083	25.309	1.00	45.02	15	681	CB	CYS	101	134.350	3.905	39.872	1.00	96.36
609 610	C O	ASP ASP	92 92	128.928 128.208	10.539 9.542	28.090 28.106	1.00 1.00	30.87 38.64		682 683	SG C	CYS CYS	101 101	135.708 133.151	4.382 2.160	41.017 41.226	1.00 1.00	100.00 90.03
611	N	GLN	93	130.247	10.490	27.914	1.00	33.31		684	Ö	CYS	101	132.261	2.914	41.595	1.00	89.78
612	CA	GLN	93	130.974	9.239	27.738	1.00	34.79		685	N	ASN	102	133.483	1.057	41.870	1.00	85.97
613 614	CB CG	GLN GLN	93 93	132.454 133.345	9.531 8.300	27.466 27.331	1.00 1.00	46.61 60.12	20	686 687	CA CB	ASN ASN	102 102	132.753 133.072	0.573 -0.902	43.043 43.238	1.00 1.00	81.73 86.41
615	CD	GLN	93	134.831	8.640	27.354	1.00	75.57		688	CG	ASN	102	132.971	-1.688	41.962	1.00	88.71
616	OE1	GLN	93	135.217	9.801	27.510	1.00	79.60		689	OD1	ASN	102	133.978	-2.107	41.412	1.00	90.82
617 618	NE2	GLN GLN	93 93	135.672 130.833	7.621 8.380	27.208 28.994	$\frac{1.00}{1.00}$	81.92 35.74		690 691	ND2	ASN	102 102	131.750 132.652	-1.869 1.257	41.462	1.00 1.00	81.79 74.07
619	C O	GLN	93	130.620	7.171	28.906	1.00	39.97		692	C	ASN ASN	102	132.032	0.881	44.413 45.187	1.00	77.40
620	N	ILE	94	130.933	9.019	30.159	1.00	32.85	25	693	N	ASP	103	133.474	2.260	44.713	1.00	58.82
621 622	CA CB	ILE ILE	94 94	130.817 131.191	8.326 9.266	31.441 32.625	1.00 $1.00$	35.57 33.17		694 695	CA CB	ASP ASP	103 103	133.377 134.746	2.904 3.418	46.037 46.524	1.00 1.00	48.91 50.06
623	CG2	ILE	94	130.909	8.588	33.969	1.00	25.21		696	CG	ASP	103	135.346	4.487	45.622	1.00	54.31
624	CG1	ILE	94	132.671	9.652	32.538	1.00	32.16		697	OD1	ASP	103	135.589	4.210	44.429	1.00	68.60
625 626	CD1 C	ILE ILE	94 94	133.120 129.407	10.631 7.770	33.603 31.645	1.00 $1.00$	32.74 38.37	30	698 699	OD2 C	ASP ASP	103 103	135.616 132.290	5.599 3.974	46.128 46.178	1.00 1.00	47.23 38.98
627	Ö	ILE	94	129.407	6.716	32.260	1.00	45.31	30	700	Ö	ASP	103	132.290	4.585	45.198	1.00	30.42
628	N	TYR	95	128.421	8.477	31.102	1.00	38.86		701	N	LEU	104	131.920	4.168	47.408	1.00	25.79
629 630	CA CB	TYR TYR	95 95	127.021 126.122	8.082 9.249	31.212 30.784	1.00 1.00	39.68 34.17		702 703	CA CB	LEU LEU	104 104	130.764 130.414	5.139 5.114	47.702 49.195	1.00 1.00	26.09 14.86
631	CG	TYR	95	124.637	8.974	30.877	1.00	27.88		704	CG	LEU	104	129.294	6.042	49.674	1.00	13.82
632	CD1	TYR	95	124.060	8.539	32.070	1.00	26.18	35	705	CD1	LEU	104	127.971	5.654	49.031	1.00	10.36
633 634	CE1 CD2	TYR TYR	95 95	122.697 123.810	8.279 9.144	32.155 29.770	1.00 1.00	24.69 21.67		706 707	CD2 C	LEU LEU	104 104	129.171 131.082	5.996 6.567	51.191 47.274	1.00 1.00	7.52 26.59
635	CE2	TYR	95	122.447	8.888	29.845	1.00	22.88		708	Ö	LEU	104	130.232	7.240	46.696	1.00	27.23
636	CZ	TYR	95	121.896	8.454	31.039	1.00	23.25		709	N	CYS	105	132.297	7.021	47.574	1.00	28.20
637 638	OH C	TYR TYR	95 95	120.546 126.715	8.185 6.846	31.112 30.369	1.00 1.00	32.19 40.10		710 711	CA CB	CYS CYS	105 105	132.735 134.164	8.370 8.606	47.225 47.721	1.00 1.00	24.21 29.54
639	O	TYR	95	125.987	5.953	30.803	1.00	41.67	40	712	SG	CYS	105	134.889	10.178	47.188	1.00	32.14
640	N	ASN	96	127.291	6.796	29.173	1.00	40.40		713	С	CYS	105	132.659	8.634	45.724	1.00	20.98
641 642	CA CB	ASN ASN	96 96	127.073 127.273	5.682 6.146	28.261 26.815	1.00 1.00	50.10 50.46		714 715	O N	CYS THR	105 106	132.062 133.258	9.618 7.744	45.285 44.941	1.00 1.00	24.94 22.03
643	CG	ASN	96	126.252	7.189	26.392	1.00	53.74		716	CA	THR	106	133.261	7.890	43.489	1.00	23.52
644	OD1	ASN	96	125.093	7.141	26.806	1.00	52.90	45	717	CB	THR	106	134.197	6.858	42.823	1.00	18.21
645 646	ND2 C	ASN ASN	96 96	126.679 127.911	8.138 4.434	25.567 28.545	1.00 1.00	56.55 54.46	43	718 719	OG1 CG2	THR THR	106 106	135.481 134.372	6.893 7.183	43.461 41.349	1.00 1.00	20.18 18.22
647	Ö	ASN	96	127.502	3.324	28.202	1.00	58.60		720	C	THR	106	131.858	7.739	42.906	1.00	22.72
648	N	GLN	97	129.067	4.606	29.183	1.00	57.00		721	O	THR	106	131.481	8.465	41.984	1.00	21.14
649 650	CA CB	GLN GLN	97 97	129.933 131.385	3.469 3.924	29.494 29.690	$\frac{1.00}{1.00}$	62.35 63.17		722 723	N CA	SER SER	107 107	131.092 129.730	6.802 6.532	43.461 43.009	$\frac{1.00}{1.00}$	26.53 24.15
651	CG	GLN	97	131.622	4.834	30.885	1.00	68.59	50	724	CB	SER	107	129.158	5.312	43.735	1.00	24.72
652	CD	GLN	97	133.052	5.351	30.970	1.00	68.54		725	OG	SER	107	129.913	4.147	43.453	1.00	34.14
653 654	OE1 NE2	GLN GLN	97 97	133.659 133.594	5.357 5.798	32.040 29.836	$\frac{1.00}{1.00}$	66.32 59.07		726 727	C	SER SER	107 107	128.818 128.128	7.731 8.170	43.228 42.306	1.00 1.00	20.49 15.26
655	C	GLN	97	129.458	2.654	30.698	1.00	66.41		728	Ň	ALA	108	128.822	8.254	44.453	1.00	18.95
656	O	GLN	97	129.682	1.442	30.754	1.00	66.29		729	CA	ALA	108	128.002	9.408	44.810	1.00	16.79
657 658	N CA	ASN ASN	98 98	128.790 128.274	3.317 2.676	31.642 32.854	1.00 $1.00$	75.07 85.32	55	730 731	CB C	ALA ALA	108 108	128.168 128.349	9.732 10.623	46.282 43.953	1.00 1.00	13.84 15.33
659	СВ	ASN	98	127.000	1.878	32.554	1.00	92.56		732	Ö	ALA	108	127.455	11.340	43.499	1.00	19.47
660	CG	ASN	98	125.798	2.771	32.325	1.00	97.44		733	N	LEU	109	129.644	10.836	43.722	1.00	10.72
661 662	OD1 ND2	ASN ASN	98 98	125.273 125.357	3.375 2.865	33.262 31.074	1.00 $1.00$	97.63 98.68		734 735	CA CB	LEU LEU	109 109	130.106 131.627	11.954 12.093	42.907 42.993	1.00 1.00	11.35 15.00
663	C	ASN	98	129.314	1.791	33.535	1.00	89.40	60	736	CG	LEU	109	132.277	13.237	42.203	1.00	19.48
664	O	ASN	98	129.073	0.612	33.812	1.00	88.19	UU	737	CD1	LEU	109	131.670	14.577	42.596	1.00	15.00
665 666	N CA	SER SER	99 99	130.486 131.560	2.369 1.640	33.779 34.435	1.00 1.00	94.53 98.60		738 739	CD2 C	LEU LEU	109 109	133.778 129.673	13.239 11.754	42.442 41.459	1.00 1.00	17.22 15.76
667	СВ	SER	99	132.918	2.248	34.106	1.00	99.63		740	O	LEU	109	129.216	12.692	40.807	1.00	28.55
668	OG	SER	99	132.996	3.591	34.559	1.00	100.00		741	N	GLN	110	129.813	10.526	40.966	1.00	19.50
669 670	C O	SER SER	99 99	131.332 131.030	1.673 2.717	35.926 36.500	1.00 1.00	99.96 98.72	65	742 743	CA CB	GLN GLN	110 110	129.417 129.679	10.184 8.699	39.600 39.339	1.00 1.00	20.50 24.06
671	N	ASN	100	131.508	0.532	36.566	1.00	100.00		744	CG	GLN	110	129.287	8.221	37.949	1.00	31.51

TABLE 10-continued

s	Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound								5	s	tructura			of Tobacco				le Synthase		
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	occ	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		
745	CD	GLN	110	129.373	6.711	37.797	1.00	31.57		818	N	GLY	118	121.872	16.813	33.613	1.00	17.98		
746 747	OE1 NE2	GLN GLN	110 110	129.661 129.117	5.990 6.225	38.756 36.586	$\frac{1.00}{1.00}$	35.16 34.19	10	819 820	CA C	GLY GLY	118 118	120.756 120.761	16.292 14.808	32.839 32.521	$\frac{1.00}{1.00}$	21.36 $21.11$		
748	C	GLN	110	127.926	10.478	39.414	1.00	21.12	10	821	Ö	GLY	118	119.760	14.284	32.032	1.00	23.89		
749	O	GLN	110	127.508	11.009	38.389	1.00	20.13		822	N	PHE	119	121.880	14.134	32.773	1.00	20.97		
750 751	N CA	PHE PHE	111 111	127.139 125.699	10.140 10.356	40.431 40.409	$\frac{1.00}{1.00}$	23.63 21.06		823 824	CA CB	PHE PHE	119 119	121.994 123.465	12.702 12.282	32.510 32.477	1.00 1.00	15.97 13.26		
752	СВ	PHE	111	125.065	9.729	41.655	1.00	21.44		825	CG	PHE	119	124.281	13.007	31.439	1.00	19.34		
753 754	CG CD1	PHE	111	123.565	9.806	41.685	1.00	18.29	15	826	CD1	PHE	119	125.155	14.025	31.808	1.00	18.10		
754 755	CD1 CD2	PHE PHE	111 111	122.795 122.921	8.933 10.744	40.924 42.483	1.00 $1.00$	18.70 14.43		827 828	CD2 CE1	PHE PHE	119 119	124.160 125.896	12.686 14.714	30.090 30.850	1.00 1.00	17.78 14.61		
756	CE1	PHE	111	121.404	8.990	40.959	1.00	18.31		829	CE2	PHE	119	124.896	13.370	29.122	1.00	16.11		
757 758	CE2 CZ	PHE PHE	111 111	121.533 120.773	10.810 9.929	42.523 41.758	1.00 $1.00$	21.91 17.47		830 831	CZ C	PHE PHE	119 119	125.765 121.238	14.386 11.917	29.503 33.576	1.00 1.00	22.05 20.90		
759	C	PHE	111	125.373	11.846	40.345	1.00	19.49	20	832	Ö	PHE	119	121.230	11.917	34.749	1.00	17.99		
760	0	PHE	111	124.731	12.305	39.399	1.00	17.88	20	833	N	ASN	120	120.157	11.263	33.161	1.00	20.78		
761 762	N CA	ARG ARG	112 112	125.857 125.606	12.598 14.033	41.332 41.407	1.00 $1.00$	16.34 8.21		834 835	CA CB	ASN ASN	120 120	119.326 117.928	10.494 10.307	34.078 33.477	1.00 1.00	23.46 23.75		
763	CB	ARG	112	126.326	14.651	42.608	1.00	7.94		836	CG	ASN	120	116.919	9.766	34.481	1.00	23.50		
764	CG	ARG	112	126.081	16.153	42.745	1.00	14.61		837	OD1	ASN	120	117.147	9.782	35.695	1.00	18.94		
765 766	CD NE	ARG ARG	112 112	126.507 127.955	16.703 16.745	44.100 44.291	1.00 $1.00$	22.36 19.69	25	838 839	ND2 C	ASN ASN	120 120	115.786 119.940	9.295 9.145	33.973 34.447	1.00 1.00	24.72 29.37		
767	CZ	ARG	112	128.777	17.561	43.639	1.00	22.41		840	Õ	ASN	120	119.467	8.092	34.011	1.00	37.56		
768	NH1	ARG	112	128.300	18.407	42.737	1.00	29.76		841	N	ILE	121	120.999	9.183	35.251	1.00	30.57		
769 770	NH2 C	ARG ARG	112 112	130.073 125.961	17.555 14.808	43.915 40.145	$\frac{1.00}{1.00}$	26.92 13.57		842 843	CA CB	ILE ILE	121 121	121.674 123.118	7.965 8.250	35.691 36.202	1.00 1.00	28.30 31.51		
771	O	ARG	112	125.113	15.505	39.588	1.00	17.92		844	CG2	ILE	121	123.116	9.395	37.212	1.00	18.33		
772 773	N CA	LEU LEU	113 113	127.205 127.671	14.676 15.385	39.693 38.504	1.00 $1.00$	11.94 14.83	30	845 846	CG1 CD1	ILE ILE	121 121	123.734 125.160	6.970 7.119	36.784 37.270	1.00 1.00	34.25 35.18		
774	CB	LEU	113	127.671	15.088	38.239	1.00	20.65		847	CD1	ILE	121	125.160	7.119	36.774	1.00	24.20		
775	CG	LEU	113	130.149	15.516	39.322	1.00	16.72		848	O	ILE	121	120.435	7.888	37.746	1.00	29.58		
776 777	CD1 CD2	LEU LEU	113 113	131.568 129.970	15.259 16.985	38.847 39.651	1.00 $1.00$	15.77 21.06		849 850	N CA	SER SER	122 122	120.654 119.886	5.963 5.158	36.594 37.538	1.00 1.00	28.31 31.22		
778	CD2	LEU	113	126.840	15.108	37.256	1.00	19.17	35	851	CB	SER	122	119.782	3.711	37.040	1.00	37.94		
779	O	LEU	113	126.484	16.034	36.532	1.00	26.79	55	852	OG	SER	122	119.046	2.907	37.948	1.00	37.31		
780 781	N CA	LEU LEU	114 114	126.516 125.717	13.841 13.464	37.014 35.849	$\frac{1.00}{1.00}$	23.83 14.66		853 854	C O	SER SER	122 122	120.471 121.690	5.193 5.086	38.942 39.121	1.00 1.00	26.25 36.49		
782	CB	LEU	114	125.668	11.943	35.703	1.00	18.52		855	N	PRO	123	119.606	5.367	39.956	1.00	25.72		
783	CG	LEU	114	126.969	11.251	35.287	1.00	19.65		856	CD	PRO	123	118.162	5.626	39.787	1.00	24.10		
784 785	CD1 CD2	LEU LEU	114 114	126.800 127.352	9.746 11.666	35.362 33.882	1.00 $1.00$	22.06 21.06	40	857 858	CA CB	PRO PRO	123 123	119.995 118.807	5.427 6.139	41.367 42.015	1.00 1.00	24.12 14.52		
786	С	LEU	114	124.300	14.030	35.939	1.00	15.67		859	CG	PRO	123	117.635	5.620	41.222	1.00	18.26		
787	O	LEU	114	123.787	14.596	34.972	1.00	18.67		860	С	PRO	123 123	120.266	4.057	41.978	1.00	26.29		
788 789	N CA	ARG ARG	115 115	123.678 122.328	13.883 14.387	37.104 37.328	1.00 $1.00$	9.26 7.86		861 862	O N	PRO GLU	123	120.649 120.106	3.957 3.007	43.143 41.176	1.00 1.00	26.93 31.58		
790	CB	ARG	115	121.849	14.021	38.736	1.00	10.49	15	863	CA	GLU	124	120.362	1.656	41.665	1.00	42.38		
791 792	CG CD	ARG ARG	115 115	121.450 120.323	12.562 12.202	38.901 37.949	1.00 $1.00$	12.35 17.12	45	864 865	CB CG	GLU GLU	124 124	119.734 118.661	0.614 -0.182	40.749 41.472	1.00 1.00	52.63 66.87		
793	NE	ARG	115	119.807	10.857	38.179	1.00	21.26		866	CD	GLU	124	117.857	-1.078	40.558	1.00	84.70		
794	CZ	ARG	115		10.570	39.000	1.00	15.94		867	OE1	GLU	124	118.072		39.323	1.00	92.92		
795 796	NH1 NH2	ARG ARG	115 115	118.199 118.406	11.537 9.314	39.676 39.152	$\frac{1.00}{1.00}$	8.20 17.38		868 869	OE2 C	GLU GLU	124 124	116.995 121.850	-1.820 1.396	41.075 41.860	$\frac{1.00}{1.00}$	95.72 40.43		
797	С	ARG	115	122.250	15.899	37.126	1.00	13.44	50	870	O	GLU	124	122.243	0.345	42.359	1.00	40.69		
798 799	O	ARG	115	121.379	16.390	36.402	1.00	8.27		871	N	ILE	125	122.665	2.383	41.493	1.00	40.56		
800	N CA	GLN GLN	$\frac{116}{116}$	123.180 123.225	16.629 18.086	37.743 37.629	$\frac{1.00}{1.00}$	14.35 11.93		872 873	CA CB	ILE ILE	125 125	124.113 124.796	2.311 3.532	41.658 40.995	$\frac{1.00}{1.00}$	33.74 34.47		
801	CB	GLN	116	124.364	18.664	38.471	1.00	4.12		874	CG2	ILE	125	124.231	4.828	41.567	1.00	35.94		
802 803	CG CD	GLN GLN	$\frac{116}{116}$	124.165 125.303	18.534 19.142	39.968 40.768	1.00 $1.00$	5.13 11.78	~ ~	875 876	CG1 CD1	ILE ILE	125 125	126.317 127.051	3.471 4.648	41.176 40.560	1.00 1.00	33.47 33.54		
804	OE1	GLN	116	125.080	19.142	41.669	1.00	21.37	55	877	CD1	ILE	125	124.397	2.311	43.166	1.00	27.96		
805	NE2	GLN	116	126.530	18.747	40.451	1.00	11.47		878	O	ILE	125	125.450	1.867	43.612	1.00	32.46		
806 807	C O	GLN GLN	116 116	123.392 123.126	18.530 19.682	36.183 35.851	$\frac{1.00}{1.00}$	15.48 19.88		879 880	N CA	PHE PHE	126 126	123.422 123.518	2.783 2.850	43.938 45.393	1.00 1.00	24.55 31.56		
808	N	HIS	117	123.827	17.607	35.328	1.00	19.55		881	CB	PHE	126	122.701	4.034	45.925	1.00	31.55		
809	CA	HIS	117	124.031	17.893	33.912	1.00	15.02	60	882	CG CD1	PHE	126	123.245	5.377	45.536	1.00	36.38		
810 811	CB CG	HIS HIS	117 117	125.405 126.538	17.392 18.253	33.460 33.925	1.00 $1.00$	13.78 17.44		883 884	CD1 CD2	PHE PHE	126 126	122.701 124.300	6.079 5.946	44.465 46.245	1.00 1.00	32.78 35.39		
812	CD2	HIS	117	126.999	18.525	35.169	1.00	18.38		885	CE1	PHE	126	123.197	7.328	44.105	1.00	32.25		
813 814	ND1 CE1	HIS HIS	117 117	127.322 128.216	18.983 19.668	33.059 33.748	1.00 $1.00$	19.26 20.36		886 887	CE2 CZ	PHE PHE	126 126	124.805 124.252	7.194 7.889	45.894 44.820	1.00 1.00	31.56 28.07		
815	NE2	HIS	117	128.210	19.408	35.031	1.00	17.37		888	C	PHE	126	124.232	1.568	46.079	1.00	37.75		
816	C	HIS	117	122.930	17.349	33.006	1.00	16.42	65	889	O	PHE	126	122.939	1.520	47.308	1.00	36.32		
817	O	HIS	117	123.036	17.419	31.780	1.00	15.29		890	N	SER	127	122.730	0.542	45.289	1.00	42.49		

TABLE 10-continued

891 CA SER 127 122.268 -0.732 45.836 1.00 43.38   892 CB SER 127 121.659 -1.601 44.733 1.00 48.30   965 CB PHE 136 122.619 2.066 53 893 OG SER 127 120.465 -1.025 44.233 1.00 59.77 10 966 CG PHE 136 123.082 3.525 53 893 OG SER 127 123.228 -2.001 47.632 1.00 39.67   896 CD PHE 136 122.848 4.238 52 895 O SER 127 123.228 -2.001 47.632 1.00 35.01 968 CD2 PHE 136 123.708 4.044 51 886 N 1XYS 128 124.567 -1.503 45.886 1.00 35.16 969 CE1 PHE 136 122.3708 4.044 51 887 CA 1XYS 128 125.743 -2.179 46.426 1.00 36.47 970 CE2 PHE 136 123.3708 4.044 51 889 CG 1XYS 128 125.743 -2.179 46.426 1.00 36.47 970 CE2 PHE 136 123.3708 4.044 51 889 CG 1XYS 128 125.743 -2.179 46.426 1.00 37.85 15 971 CZ PHE 136 123.3708 4.044 51 900 CD 1XYS 128 128.170 -0.947 43.606 1.00 37.35 15 973 O PHE 136 123.308 1.280 54 900 CD 1XYS 128 128.353 0.388 42.892 1.00 50.12 974 N 1XYS 137 125.323 -0.232 55 903 C 1XYS 128 129.338 0.328 41.776 1.00 54.36 975 CA 1XYS 137 125.232 -0.232 55 903 C 1XYS 128 129.338 0.328 41.776 1.00 54.36 975 CA 1XYS 137 125.845 -2.039 53 905 N PHE 129 125.656 -0.501 48.199 1.00 38.61 977 CG 1XYS 137 125.585 -3.745 52 906 CA PHE 129 125.656 -0.501 48.199 1.00 38.63 979 CE 1XYS 137 125.585 -3.745 52 900 CD PHE 129 125.656 -0.501 48.199 1.00 38.69 900 CD PHE 129 125.656 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.650 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.650 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 38.69 900 CD PHE 129 125.556 0.0501 48.199 1.00 50.00 900 000 PHE 129 125.650 0.0501 48.199 1.00 900 000 PHE 129 125.650 0.0501 48.19	lochene Synthase Bound
892         CB         SER         127         121.659         -1.601         44.733         1.00         48.30         965         CB         PHE         136         123.082         3.525         53           893         G         SER         127         123.401         -1.482         46.527         1.00         39.67         967         CDI         PHE         136         121.752         5.079         52           895         O         SER         127         123.228         -2.001         47.632         1.00         35.01         968         CD2         PHE         136         121.752         5.719         52           896         N         IYS         128         124.567         -1.503         45.886         1.00         35.16         969         CEI         PHE         136         123.478         4.044         51           897         CA         IYS         128         126.877         -2.180         45.889         1.00         35.16         969         CEI         PHE         136         123.478         4.04         58           899         CG         IYS         128         122.180         44.732         1.00         35.25	Z OCC facto
893 OG SER 127 120.465 -1.025 44.233 1.00 59.77 10 966 CG PHE 136 122.848 4.238 52 894 C SER 127 123.401 -1.482 46.527 1.00 39.67 967 CD1 PHE 136 121.752 5.079 52 895 O SER 127 123.228 -2.001 47.632 1.00 35.01 968 CD2 PHE 136 121.752 5.079 52 896 N LYS 128 124.567 -1.503 45.886 1.00 35.16 969 CE1 PHE 136 121.512 5.714 51 897 CA LYS 128 125.743 -2.179 46.426 1.00 36.47 970 CE2 PHE 136 122.378 4.674 50 898 CB LYS 128 126.77 -2.180 45.389 1.00 33.52 971 CZ PHE 136 122.376 5.510 50 899 CG LYS 128 127.146 -0.834 44.732 1.00 37.85 15 972 C PHE 136 122.376 5.510 51 900 CD LYS 128 128.170 -0.947 43.606 1.00 37.35 973 O PHE 136 122.376 5.510 54 901 CE LYS 128 128.353 0.388 42.892 1.00 50.12 974 N LYS 137 124.404 0.564 54 902 NZ LYS 128 126.233 -1.623 47.769 1.00 54.36 975 CA LYS 137 125.232 -0.232 55 903 C LYS 128 127.102 -2.217 48.412 1.00 46.49 977 CG LYS 137 125.845 -2.039 53 905 N PHE 129 126.030 1.625 49.466 1.00 38.63 979 CE LYS 137 125.845 -2.039 53 906 CA PHE 129 126.609 1.626 49.256 1.00 28.66 981 C LYS 137 125.872 0.698 56 909 CD1 PHE 129 126.640 2.506 46.997 1.00 19.33 982 O LYS 137 127.009 -4.340 51 910 CD2 PHE 129 128.653 1.537 48.368 1.00 17.79 983 N GLU 138 125.569 0.472 57 911 CE1 PHE 129 126.946 2.506 46.997 1.00 19.33 982 O LYS 137 125.872 0.698 56 912 CE2 PHE 129 129.550 1.760 47.368 1.00 17.79 983 N GLU 138 123.907 1.846 0.914 C PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.907 1.846 0.914 C PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.907 1.846 0.915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.907 3.180 61 916 CB GLN 130 121.456 -0.310 50.403 1.00 51.16 30 991 O GLU 138 123.907 5.135 57 917 CA GLN 130 122.384 -0.700 50.123 1.00 40.45 989 OE2 GLU 138 123.907 3.180 61 918 CB GLN 130 121.456 -0.310 50.403 1.00 50.70 992 N SER 139 128.210 0.242 58 919 CG GLN 130 122.486 -2.405 51.263 1.00 50.70 992 N SER 139 129.597 -1.354 57 922 NE2 GLN 130 122.496 -2.405 51.263 1.00 51.99 995 OG SER 139 129.518 -1.477 56 924 O GLN 130 122.486 -2.405 51.263 1.00 51.99 995	3.929 1.00 37.7
894         C         SER         127         123.401         -1.482         46.527         1.00         39.67         967         CD1         PHE         136         121.752         5.079         52           895         O         SER         127         123.228         -2.001         47.632         1.00         35.01         968         CD2         PHE         136         123.708         4.044         5.71         51         898         CB         LYS         128         124.567         -1.503         45.886         1.00         35.16         969         CE1         PHE         136         123.708         4.044         50         898         CB         LYS         128         125.743         -2.179         46.426         1.00         36.47         970         CE2         PHE         136         123.478         4.674         50           899         CB         LYS         128         127.146         -0.834         44.732         1.00         37.35         15         971         CZ         PHE         136         123.368         12.80         49         10         137         124.404         0.564         54         40         123.70         124.404         0.5	3.941 1.00 30.3 2.644 1.00 18.9
896         N         LYS         128         124.567         -1.503         45.886         1.00         35.16         969         CE1         PHE         136         121.512         5.714         51           897         CA         LYS         128         125.743         -2.179         46.426         1.00         36.47         970         CE2         PHE         136         122.378         4.674         50           899         CG         LYS         128         127.146         -0.834         44.732         1.00         37.85         15         972         C         PHE         136         122.376         5.510         50           900         CD         LYS         128         128.170         -0.947         43.606         1.00         37.35         973         O         PHE         136         123.368         1.28         129.338         0.328         41.776         1.00         54.36         975         CA         LYS         137         124.040         0.564         54           903         C         LYS         128         129.338         0.328         41.776         1.00         38.71         976         CB         LYS         137	2.485 1.00 16.8
897         CA         LYS         128         125.743         -2.179         46.426         1.00         36.47         970         CE2         PHE         136         123.478         4.674         5.00           898         CB         LYS         128         125.743         -2.180         45.389         1.00         33.52         971         CZ         PHE         136         123.376         5.510         50           900         CD         LYS         128         128.170         -0.947         43.606         1.00         37.85         15         973         O         PHE         136         123.307         1.310         56           901         CE         LYS         128         128.353         0.388         42.892         1.00         50.12         974         N         LYS         137         125.232         -0.232         55           903         C         LYS         128         126.233         -1.623         47.769         1.00         54.36         975         CA         LYS         137         125.333         -0.957         54           904         O         LYS         128         127.102         -2.217         48.412	.569 1.00 2.9
898         CB         LYS         128         126.877         -2.180         45.389         1.00         33.52         971         CZ         PHE         136         122.376         5.510         50           899         CG         LYS         128         127.146         -0.834         44.732         1.00         37.85         15         972         C         PHE         136         123.368         1.280         54           900         CD         LYS         128         128.170         -0.947         43.606         1.00         37.35         973         O         PHE         136         123.307         1.310         56           901         CE         LYS         128         128.353         0.388         42.892         1.00         50.12         974         N         LYS         137         125.232         -0.232         55           903         C         LYS         128         126.233         -1.623         47.769         1.00         38.63         975         CA         LYS         137         126.333         -0.957         54           904         O         LYS         129         126.023         -1.510         48.412	269 1.00 9.3 ).350 1.00 8.3
900 CD LYS 128 128.170	0.200 1.00 11.6
901 CE LYS 128 128.353 0.388 42.892 1.00 50.12 974 N LYS 137 124.404 0.564 54 902 NZ LYS 128 129.338 0.328 41.776 1.00 54.36 975 CA LYS 137 125.232 -0.232 55 903 C LYS 128 126.233 -1.623 47.769 1.00 38.71 976 CB LYS 137 125.332 -0.232 55 905 N PHE 129 125.656 -0.501 48.199 1.00 38.63 905 N PHE 129 126.028 0.135 49.466 1.00 31.98 977 CG LYS 137 125.845 -2.039 53 905 CB PHE 129 126.309 1.626 49.256 1.00 24.98 980 NZ LYS 137 127.016 -2.672 52 906 CA PHE 129 126.309 1.626 49.256 1.00 24.98 980 NZ LYS 137 125.872 0.698 56 909 CD1 PHE 129 126.946 2.506 46.997 1.00 19.33 982 O LYS 137 125.872 0.698 56 909 CD1 PHE 129 128.653 1.537 48.368 1.00 17.79 983 N GLU 138 125.569 0.472 57 911 CE1 PHE 129 127.877 2.735 45.988 1.00 22.73 911 CE1 PHE 129 129.590 1.760 47.368 1.00 19.77 25 985 CB GLU 138 125.482 0.895 60 913 CZ PHE 129 124.929 -0.024 50.509 1.00 31.84 987 CD GLU 138 123.997 1.184 60 915 O PHE 129 124.929 -0.024 50.509 1.00 31.84 987 CD GLU 138 123.040 3.272 59 917 CA LYS 137 126.333 -0.957 54 9.90 CD GLN 130 121.456 -0.310 50.403 1.00 51.76 999 CB GLU 138 123.040 3.272 59 919 CG GLN 130 121.456 -0.310 50.403 1.00 50.70 992 N SER 139 129.653 0.027 58 921 OG LN 130 120.394 3.017 49.105 5.049 1.00 55.44 35 997 O SER 139 131.606 1.247 57 924 O GLN 130 122.846 -3.245 50.449 1.00 55.44 35 997 O SER 139 131.606 1.247 57 924 O GLN 130 122.846 -3.245 50.449 1.00 55.44 35 997 O SER 139 131.606 1.247 57	.992 1.00 40.0
902 NZ LYS 128 129.338 0.328 41.776 1.00 54.36 975 CA LYS 137 125.232 -0.232 55 903 C LYS 128 126.233 -1.623 47.769 1.00 38.71 976 CB LYS 137 125.333 -0.957 54 904 O LYS 128 127.102 -2.217 48.412 1.00 46.49 977 CG LYS 137 125.845 -2.039 53 905 N PHE 129 126.028 0.135 49.466 1.00 31.98 978 CD LYS 137 127.016 -2.672 52 906 CA PHE 129 126.309 1.626 49.256 1.00 24.98 980 NZ LYS 137 127.016 -2.672 52 907 CB PHE 129 127.324 1.904 48.191 1.00 20.86 981 C LYS 137 125.872 0.698 56 909 CD1 PHE 129 128.653 1.537 48.368 1.00 17.79 983 N GLU 138 125.569 0.472 57 911 CE1 PHE 129 127.877 2.735 45.988 1.00 22.73 984 CA GLU 138 125.569 0.472 57 913 CZ PHE 129 129.201 2.361 46.174 1.00 17.69 986 CG GLU 138 123.997 1.184 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.997 1.184 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.997 1.184 60 916 N GLN 130 122.720 -0.922 51.010 1.00 47.58 990 CC GLU 138 123.040 3.272 59 917 CA GLN 130 122.720 -0.922 51.010 1.00 47.58 990 CC GLU 138 123.040 3.272 59 917 CA GLN 130 122.720 -0.922 51.010 1.00 47.58 990 CC GLU 138 123.040 3.272 59 919 CG GLN 130 121.515 1.197 50.231 1.00 50.70 992 N SER 139 129.653 0.027 58 921 OE1 GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.975 -1.354 57 922 NE2 GLN 130 122.496 -2.405 51.263 1.00 51.99 996 C SER 139 131.606 1.247 57 924 O GLN 130 122.496 -2.405 51.263 1.00 55.44 35 997 O SER 139 131.606 1.247 57	5.173 1.00 36.0 6.554 1.00 35.2
904 O LYS 128 127.102 -2.217 48.412 1.00 46.49 977 CG LYS 137 125.845 -2.039 53 905 N PHE 129 125.656 -0.501 48.199 1.00 38.63 979 CB LYS 137 125.845 -2.039 53 906 CA PHE 129 126.028 0.135 49.466 1.00 31.98 979 CE LYS 137 125.558 -3.745 52 907 CB PHE 129 126.309 1.626 49.256 1.00 24.98 980 NZ LYS 137 125.872 0.698 56 909 CD1 PHE 129 126.946 2.506 46.997 1.00 19.33 982 O LYS 137 125.872 0.698 56 909 CD1 PHE 129 128.653 1.537 48.368 1.00 17.79 983 N GLU 138 125.569 0.472 57 911 CE1 PHE 129 127.877 2.735 45.988 1.00 22.73 984 CA GLU 138 125.482 0.895 60 913 CZ PHE 129 129.590 1.760 47.368 1.00 17.79 985 CB GLU 138 125.482 0.895 60 913 CZ PHE 129 129.201 2.361 46.174 1.00 17.69 986 CG GLU 138 123.097 1.184 60 914 C PHE 129 124.929 -0.024 50.509 1.00 31.84 987 CD GLU 138 123.097 1.184 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.00 3.272 59 917 CA GLN 130 122.8270 -0.922 51.010 1.00 40.45 989 OE2 GLU 138 123.040 3.272 59 919 CG GLN 130 121.456 -0.310 50.403 1.00 51.46 30 991 O GLU 138 123.040 3.272 59 919 CG GLN 130 121.351 1.197 50.231 1.00 50.70 992 N SER 139 129.975 -1.354 57 922 NE2 GLN 130 120.394 3.017 49.105 1.00 51.40 3 1.00 51.40 3 51.40 30 991 O GLU 138 129.518 -1.477 56 923 C GLN 130 122.394 3.017 49.105 51.263 1.00 55.44 35 997 O SER 139 131.606 1.247 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	i.451 1.00 37.6
905 N PHE 129 125.656 -0.501 48.199 1.00 38.63 20 978 CD LYS 137 127.016 -2.672 52 906 CA PHE 129 126.028 0.135 49.466 1.00 31.98 979 CE LYS 137 126.558 -3.745 52 907 CB PHE 129 126.309 1.626 49.256 1.00 24.98 980 NZ LYS 137 127.709 -4.340 51 908 CG PHE 129 127.324 1.904 48.191 1.00 20.86 981 C LYS 137 125.872 0.698 56 909 CD1 PHE 129 126.640 2.506 46.997 1.00 19.33 982 O LYS 137 125.872 0.698 56 910 CD2 PHE 129 128.653 1.537 48.368 1.00 17.79 983 N GLU 138 125.569 0.472 57 911 CE1 PHE 129 127.877 2.735 45.988 1.00 22.73 985 CB GLU 138 125.482 0.895 60 913 CZ PHE 129 129.590 1.760 47.368 1.00 17.79 985 CB GLU 138 123.997 1.184 60 914 C PHE 129 124.929 -0.024 50.509 1.00 31.84 987 CD GLU 138 123.997 1.184 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.070 2.650 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.040 3.272 59 917 CA GLN 130 122.720 -0.922 51.010 1.00 47.58 989 OE2 GLU 138 123.040 3.272 59 919 CG GLN 130 121.456 -0.310 50.403 1.00 51.16 30 991 O GLU 138 122.270 0.242 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.975 -1.354 57 922 NE2 GLN 130 122.394 3.017 49.105 51.263 1.00 55.44 35 997 O SER 139 131.606 1.247 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	.670 1.00 36.0
906 CA PHE 129 126.028 0.135 49.466 1.00 31.98 20 979 CE LYS 137 126.558 -3.745 52 907 CB PHE 129 126.309 1.626 49.256 1.00 24.98 980 NZ LYS 137 127.709 -4.340 51 908 CG PHE 129 127.324 1.904 48.191 1.00 20.86 981 C LYS 137 125.872 0.698 56 909 CD1 PHE 129 126.946 2.506 46.997 1.00 19.33 982 O LYS 137 126.612 1.614 56 910 CD2 PHE 129 128.653 1.537 48.368 1.00 17.79 983 N GLU 138 125.569 0.472 57 911 CE1 PHE 129 127.877 2.735 45.988 1.00 22.73 984 CA GLU 138 125.569 0.472 57 911 CE2 PHE 129 129.590 1.760 47.368 1.00 19.77 25 985 CB GLU 138 123.997 1.184 60 913 CZ PHE 129 129.201 2.361 46.174 1.00 17.69 986 CG GLU 138 123.997 1.184 60 915 O PHE 129 124.929 -0.024 50.509 1.00 31.84 987 CD GLU 138 123.703 2.650 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.703 2.650 60 916 N GLN 130 123.854 -0.700 50.123 1.00 40.45 989 OE2 GLU 138 123.040 3.272 59 917 CA GLN 130 122.720 -0.922 51.010 1.00 47.58 990 C GLU 138 123.040 3.272 59 919 CG GLN 130 121.515 1.197 50.231 1.00 50.70 992 N SER 139 128.231 0.024 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.975 -1.354 57 922 NE2 GLN 130 120.394 3.017 49.105 1.00 55.44 35 997 O SER 139 131.606 1.247 57 924 O GLN 130 122.496 -2.405 51.263 1.00 55.44 35 997 O SER 139 131.606 1.247 57 924 O GLN 130 122.886 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	3.721 1.00 43.9 2.985 1.00 45.6
908 CG PHE 129 127.324 1.904 48.191 1.00 20.86 981 C LYS 137 125.872 0.698 56 909 CD1 PHE 129 126.946 2.506 46.997 1.00 19.33 982 O LYS 137 126.612 1.614 56 910 CD2 PHE 129 128.653 1.537 48.368 1.00 17.79 983 N GLU 138 125.569 0.472 57 911 CE1 PHE 129 127.877 2.735 45.988 1.00 22.73 984 CA GLU 138 125.569 0.472 57 912 CE2 PHE 129 129.590 1.760 47.368 1.00 19.77 25 985 CB GLU 138 125.482 0.895 60 913 CZ PHE 129 129.201 2.361 46.174 1.00 17.69 986 CG GLU 138 123.997 1.184 60 914 C PHE 129 124.929 -0.024 50.509 1.00 31.84 987 CD GLU 138 123.703 2.650 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.040 3.272 59 917 CA GLN 130 123.854 -0.700 50.123 1.00 40.45 989 OE2 GLU 138 123.040 3.272 59 918 CB GLN 130 122.720 -0.922 51.010 1.00 47.58 990 CC GLU 138 127.641 1.172 58 918 CB GLN 130 121.515 1.197 50.231 1.00 50.70 992 N SER 139 128.210 0.242 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.975 -1.354 57 922 NE2 GLN 130 120.394 3.017 49.105 1.00 55.44 35 997 O SER 139 131.606 1.247 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	2.011 1.00 46.9
909 CD1 PHE 129 126.946 2.506 46.997 1.00 19.33 982 O LYS 137 126.612 1.614 56 910 CD2 PHE 129 128.653 1.537 48.368 1.00 17.79 983 N GLU 138 125.569 0.472 57 911 CE1 PHE 129 127.877 2.735 45.988 1.00 19.77 25 985 CB GLU 138 125.482 0.895 60 913 CZ PHE 129 129.590 1.760 47.368 1.00 19.77 25 985 CB GLU 138 125.482 0.895 60 913 CZ PHE 129 129.201 2.361 46.174 1.00 17.69 986 CG GLU 138 123.997 1.184 60 914 C PHE 129 124.929 -0.024 50.509 1.00 31.84 987 CD GLU 138 123.703 2.650 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.040 3.272 59 917 CA GLN 130 123.854 -0.700 50.123 1.00 40.45 989 OE2 GLU 138 123.040 3.272 59 918 CB GLN 130 121.456 -0.310 50.403 1.00 51.16 30 991 O GLU 138 123.040 3.272 59 919 CG GLN 130 121.515 1.197 50.231 1.00 50.70 992 N SER 139 128.210 0.242 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.975 -1.354 57 922 NE2 GLN 130 122.394 3.017 49.105 1.00 51.94 35 995 OG SER 139 129.518 -1.477 56 923 C GLN 130 122.846 -2.405 51.263 1.00 55.44 35 997 O SER 139 131.606 1.247 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	.276 1.00 45.4
910 CD2 PHE 129 128.653 1.537 48.368 1.00 17.79 983 N GLU 138 125.569 0.472 57 911 CE1 PHE 129 127.877 2.735 45.988 1.00 22.73 984 CA GLU 138 125.161 1.290 58 913 CZ PHE 129 129.590 1.760 47.368 1.00 19.77 25 985 CB GLU 138 125.482 0.895 60 913 CZ PHE 129 129.201 2.361 46.174 1.00 17.69 986 CG GLU 138 123.997 1.184 60 914 C PHE 129 124.929 -0.024 50.509 1.00 31.84 987 CD GLU 138 123.703 2.650 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.703 2.650 60 916 N GLN 130 123.854 -0.700 50.123 1.00 40.45 989 OE2 GLU 138 123.040 3.272 59 917 CA GLN 130 122.720 -0.922 51.010 1.00 47.58 990 C GLU 138 123.040 3.272 59 919 CG GLN 130 121.456 -0.310 50.403 1.00 51.16 30 991 O GLU 138 128.283 1.909 59 919 CG GLN 130 121.456 -0.310 50.403 1.00 50.70 992 N SER 139 128.210 0.242 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.975 -1.354 57 922 NE2 GLN 130 120.394 3.017 49.105 1.00 55.44 35 997 O SER 139 131.606 1.247 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	5.472 1.00 42.7 5.108 1.00 49.7
912 CE2 PHE 129 129.590 1.760 47.368 1.00 19.77 25 985 CB GLU 138 125.482 0.895 60 913 CZ PHE 129 129.201 2.361 46.174 1.00 17.69 986 CG GLU 138 123.997 1.184 60 914 C PHE 129 124.929 -0.024 50.509 1.00 31.84 987 CD GLU 138 123.703 2.650 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 123.040 3.272 59 917 CA GLN 130 123.854 -0.700 50.123 1.00 40.45 989 OE2 GLU 138 123.040 3.272 59 917 CA GLN 130 122.720 -0.922 51.010 1.00 47.58 990 C GLU 138 123.040 3.272 59 918 CB GLN 130 121.456 -0.310 50.403 1.00 51.16 30 991 O GLU 138 123.040 3.272 59 919 CG GLN 130 121.515 1.197 50.231 1.00 50.70 992 N SER 139 128.210 0.242 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.653 0.027 58 921 OE1 GLN 130 119.310 1.063 49.303 1.00 62.26 994 CB SER 139 129.975 -1.354 57 922 NE2 GLN 130 122.496 -2.405 51.263 1.00 51.99 996 C SER 139 130.384 1.114 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	7.747 1.00 44.3
913 CZ PHE 129 129.201 2.361 46.174 1.00 17.69 986 CG GLU 138 123.997 1.184 60 914 C PHE 129 124.929 -0.024 50.509 1.00 31.84 987 CD GLU 138 123.703 2.650 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 124.127 3.180 61 916 N GLN 130 123.854 -0.700 50.123 1.00 40.45 989 OE2 GLU 138 123.040 3.272 59 917 CA GLN 130 122.720 -0.922 51.010 1.00 47.58 990 CC GLU 138 127.641 1.172 58 918 CB GLN 130 121.456 -0.310 50.403 1.00 51.16 30 991 O GLU 138 128.283 1.909 59 919 CG GLN 130 121.515 1.197 50.231 1.00 50.70 992 N SER 139 128.210 0.242 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.655 0.027 58 921 OE1 GLN 130 119.310 1.063 49.303 1.00 62.26 994 CB SER 139 129.975 -1.354 57 922 NE2 GLN 130 122.496 -2.405 51.263 1.00 51.99 996 C SER 139 130.384 1.114 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	3.824 1.00 43.3
914 C PHE 129 124.929 -0.024 50.509 1.00 31.84 987 CD GLU 138 123.703 2.650 60 915 O PHE 129 125.051 0.462 51.635 1.00 32.60 988 OE1 GLU 138 124.127 3.180 61 916 N GLN 130 123.854 -0.700 50.123 1.00 40.45 989 OE2 GLU 138 123.040 3.272 59 917 CA GLN 130 122.720 -0.922 51.010 1.00 47.58 990 CC GLU 138 127.641 1.172 58 918 CB GLN 130 121.456 -0.310 50.403 1.00 51.16 30 991 O GLU 138 128.283 1.909 59 919 CG GLN 130 121.515 1.197 50.231 1.00 50.70 992 N SER 139 128.210 0.242 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.653 0.027 58 921 OE1 GLN 130 119.310 1.063 49.303 1.00 62.26 994 CB SER 139 129.975 -1.354 57 922 NE2 GLN 130 120.394 3.017 49.105 1.00 58.79 995 OG SER 139 129.518 -1.477 56 923 C GLN 130 122.496 -2.405 51.263 1.00 51.99 996 C SER 139 130.384 1.114 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	0.157 1.00 48.2 0.285 1.00 55.5
916 N GLN 130 123.854 -0.700 50.123 1.00 40.45 989 OE2 GLU 138 123.040 3.272 59 917 CA GLN 130 122.720 -0.922 51.010 1.00 47.58 990 C GLU 138 123.040 3.272 59 918 CB GLN 130 121.456 -0.310 50.403 1.00 51.16 30 991 O GLU 138 128.283 1.909 59 919 CG GLN 130 121.515 1.197 50.231 1.00 50.70 992 N SER 139 128.210 0.242 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.653 0.027 58 921 OE1 GLN 130 119.310 1.063 49.303 1.00 62.26 994 CB SER 139 129.975 -1.354 57 922 NE2 GLN 130 120.394 3.017 49.105 1.00 58.79 995 OG SER 139 129.518 -1.477 56 923 C GLN 130 122.496 -2.405 51.263 1.00 51.99 996 C SER 139 130.384 1.114 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	0.528 1.00 59.8
917 CA GLN 130 122.720 -0.922 51.010 1.00 47.58 990 C GLU 138 127.641 1.172 58 918 CB GLN 130 121.456 -0.310 50.403 1.00 51.16 30 991 O GLU 138 128.283 1.909 59 919 CG GLN 130 121.515 1.197 50.231 1.00 50.70 992 N SER 139 128.210 0.242 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.653 0.027 58 921 OE1 GLN 130 119.310 1.063 49.303 1.00 62.26 994 CB SER 139 129.975 -1.354 57 922 NE2 GLN 130 120.394 3.017 49.105 1.00 58.79 995 OG SER 139 129.518 -1.477 56 923 C GLN 130 122.496 -2.405 51.263 1.00 51.99 996 C SER 139 130.384 1.114 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	.577 1.00 58.0
918 CB GLN 130 121.456 -0.310 50.403 1.00 51.16 30 991 O GLU 138 128.283 1.909 59 919 CG GLN 130 121.515 1.197 50.231 1.00 50.70 992 N SER 139 128.210 0.242 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.653 0.027 58 921 OE1 GLN 130 119.310 1.063 49.303 1.00 62.26 994 CB SER 139 129.975 -1.354 57 922 NE2 GLN 130 120.394 3.017 49.105 1.00 58.79 995 OG SER 139 129.915 -1.477 56 923 C GLN 130 122.496 -2.405 51.263 1.00 51.99 996 C SER 139 130.384 1.114 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	9.674 1.00 70.0 3.913 1.00 46.1
919 CG GLN 130 121.515 1.197 50.231 1.00 50.70 992 N SER 139 128.210 0.242 58 920 CD GLN 130 120.308 1.755 49.505 1.00 54.25 993 CA SER 139 129.655 0.027 58 921 OE1 GLN 130 119.310 1.063 49.303 1.00 62.26 994 CB SER 139 129.975 -1.354 57 922 NE2 GLN 130 120.394 3.017 49.105 1.00 58.79 995 OG SER 139 129.518 -1.477 56 923 C GLN 130 122.496 -2.405 51.263 1.00 51.99 996 C SER 139 130.384 1.114 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	0.662 1.00 40.10 0.662 1.00 51.0
921         OE1         GLN         130         119.310         1.063         49.303         1.00         62.26         994         CB         SER         139         129.975         -1.354         57           922         NE2         GLN         130         120.394         3.017         49.105         1.00         58.79         995         OG         SER         139         129.518         -1.477         56           923         C         GLN         130         122.496         -2.405         51.263         1.00         51.99         996         C         SER         139         130.384         1.114         57           924         O         GLN         130         122.818         -3.245         50.419         1.00         55.44         35         997         O         SER         139         131.606         1.247         57	3.149 1.00 40.3
922 NE2 GLN 130 120.394 3.017 49.105 1.00 58.79 995 OG SER 139 129.518 -1.477 56 923 C GLN 130 122.496 -2.405 51.263 1.00 51.99 996 C SER 139 130.384 1.114 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	3.122 1.00 37.2
923 C GLN 130 122.496 -2.405 51.263 1.00 51.99 996 C SER 139 130.384 1.114 57 924 O GLN 130 122.818 -3.245 50.419 1.00 55.44 35 997 O SER 139 131.606 1.247 57	7.541 1.00 42.9 5.204 1.00 42.4
	7.326 1.00 38.8
	7.423 1.00 44.3
	5.531 1.00 35.6 5.721 1.00 26.8
927 OB ASP 131 121.556 -4.258 54.314 1.00 58.61 1000 CB LEU 140 129.289 3.244 54	1.514 1.00 27.1
	3.476 1.00 27.1 2.511 1.00 23.2
930 OD2 ASP 131 119.893 _2.532 54.391 1.00 65.85 10.03 CD2 LEU 140 130.317 1.814 52	2.511 1.00 23.2 2.726 1.00 21.3
931 C ASP 131 120.382 4.583 52.103 1.00 64.98 <sup>4U</sup> 1004 C LEU 140 130.325 4.241 56	5.547 1.00 23.4
	5.054 1.00 18.1
	7.803 1.00 21.1 3.725 1.00 23.0
935 CB GLU 132 118.735 -7.913 52.120 1.00 78.41 1008 CB ALA 141 129.182 4.951 60	0.007 1.00 13.9
	0.053 1.00 29.6 0.614 1.00 34.0
70, 00 020 200 200 0000 0000 2000 0 1222 212 201100 002	3.710 1.00 34.0 3.710 1.00 31.9
939 OE2 GLU 132 118.331 -9.837 55.165 1.00 100.00 1012 CA SER 142 133.723 5.413 58	3.963 1.00 25.7
	9.534 1.00 27.7 3.731 1.00 23.3
	7.719 1.00 25.3
943 CA ASN 133 116.408 4.124 53.726 1.00 71.81 50 1016 O SER 142 135.629 6.255 57	7.761 1.00 36.0
	5.617 1.00 19.4 5.361 1.00 13.4
	1.194 1.00 12.4
947 ND2 ASN 133 115.288 -2.303 56.583 1.00 85.57 1020 CG ASP 143 134.284 6.196 52	2.835 1.00 22.6
	.811 1.00 21.9 2.778 1.00 35.1
	5.162 1.00 19.6
951 CA GLY 134 117.120 -1.177 51.416 100 63.86 1024 O ASP 143 132.675 8.238 54	1.559 1.00 21.7
	5.664 1.00 21.8 5.566 1.00 19.3
	5.376 1.00 19.5 5.376 1.00 16.6
955 CA LYS 135 118.989 0.921 53.993 1.00 52.54 60 1028 CG1 VAL 144 134.789 12.769 56	5.155 1.00 22.1
930 CB E13 133 116.026 0.803 33.462 1.00 34.30 1029 CG2 VAL 144 134.976 10.901 37	7.857 1.00 20.4 1.136 1.00 18.9
	3.801 1.00 18.9.
959 CE LYS 135 114.795 1.428 55.666 1.00 66.94 1032 N LEU 145 134.984 10.657 53	3.293 1.00 15.1
	.912 1.00 18.5 .181 1.00 22.9
	.722 1.00 23.2
	0.715 1.00 24.4

TABLE 10-continued

S	tructura			of Tobacco l Hydroxy				hase	5	s _	tructura			of Tobacco yl Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	Z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
1037	CD2	LEU	145	137.327	12.898	51.968	1.00	18.75		1110	CB	HIS	155	120.920	18.177	47.928	1.00	2.00
1038	C	LEU	145	133.744	10.535	51.168	1.00	23.90		1111	CG	HIS	155	120.932	18.340	49.415	1.00	2.00
1039	O	LEU	145	133.236	11.135	50.219	1.00	26.81	10	1112	CD2	HIS	155	120.036	18.914	50.255	1.00	3.97
1040 1041	N CA	GLY GLY	146 146	133.303 132.159	9.358 8.705	51.610 50.999	1.00 $1.00$	24.92 21.44		1113 1114	ND1 CE1	HIS HIS	155 155	121.946 121.676	17.856 18.123	50.211 51.476	1.00 1.00	3.74 12.80
1042	C	GLY	146	130.868	9.313	51.512	1.00	19.58		1115	NE2	HIS	155	120.522	18.764	51.529	1.00	11.17
1043	O	GLY	146	129.953	9.591	50.740	1.00	22.50		1116	C	HIS	155	119.742	17.092	45.997	1.00	10.18
1044	N	LEU	147	130.805	9.524	52.823	1.00	9.20		1117	O	HIS	155	119.025	18.030	45.645	1.00	16.07
1045	CA	LEU	147	129.643	10.116	53.467	1.00	7.33	15	1118	N	VAL	156	120.182	16.163	45.152	1.00	6.86
1046 1047	CB CG	LEU LEU	147 147	129.849 129.927	10.163 8.831	54.980 55.721	1.00 $1.00$	7.87 12.02		1119 1120	CA CB	VAL VAL	156 156	119.843 121.109	16.202 16.099	43.733 42.823	1.00 1.00	7.09 5.97
1047	CD1	LEU	147	130.341	9.066	57.157	1.00	8.20		1121	CG1	VAL	156	122.161	17.113	43.248	1.00	2.00
1049	CD2	LEU	147	128.583	8.122	55.656	1.00	15.08		1122	CG2	VAL	156	121.678	14.682	42.841	1.00	4.04
1050	C	LEU	147	129.388	11.527	52.945	1.00	20.24		1123	С	VAL	156	118.866	15.087	43.354	1.00	11.84
1051	O	LEU	147	128.244	11.900	52.680	1.00	27.36	20	1124	O	VAL	156	118.644	14.827	42.170	1.00	13.76
1052 1053	N CA	LEU LEU	148 148	130.462 130.371	12.303 13.676	52.795 52.304	1.00 $1.00$	20.47 20.11		1125 1126	N CA	ARG ARG	157 157	118.264 117.329	14.443 13.357	44.351 44.074	1.00 1.00	12.59 21.43
1054	CB	LEU	148	131.751	14.347	52.330	1.00	16.79		1127	CB	ARG	157	117.224	12.398	45.271	1.00	17.56
1055	CG	LEU	148	131.829	15.805	51.857	1.00	13.41		1128	OG	ARG	157	116.482	12.908	46.491	1.00	22.45
1056	CD1	LEU	148	130.897	16.683	52.683	1.00	6.32		1129	CD	ARG	157	116.525	11.846	47.583	1.00	26.00
1057	CD2	LEU	148	133.256	16.306	51.961	1.00	9.98	25	1130	NE	ARG	157	115.512	12.037	48.620	1.00	35.19
1058	С	LEU	148	129.777	13.758	50.895	1.00	16.22	25	1131	CZ	ARG	157	114.360 114.064	11.370	48.676	1.00	40.41
1059 1060	O N	LEU ASN	148 149	128.838 130.332	14.520 12.985	50.657 49.965	$\frac{1.00}{1.00}$	19.05 14.19		1132 1133	NH1 NH2	ARG ARG	157 157	113.505	10.465 11.598	47.753 49.664	$\frac{1.00}{1.00}$	42.25 42.66
1061	CA	ASN	149	129.840	12.986	48.592	1.00	19.68		1134	C	ARG	157	115.945	13.815	43.609	1.00	22.46
1062	CB	ASN	149	130.776	12.199	47.678	1.00	17.57		1135	O	ARG	157	115.473	14.885	43.985	1.00	28.62
1063	CG	ASN	149	132.009	12.987	47.306	1.00	21.68		1136	N	THR	158	115.334	13.012	42.740	1.00	30.57
1064	OD1	ASN	149	132.904	13.181	48.129	1.00	27.23	30	1137	CA	THR	158	114.003	13.287	42.200	1.00	23.48
1065 1066	ND2 C	ASN ASN	149 149	132.055 128.414	13.469 12.461	46.067 48.486	$\frac{1.00}{1.00}$	17.93 24.02		1138 1139	CB OG1	THR THR	158 158	113.951 114.132	13.012 11.613	40.675 40.424	1.00 1.00	18.85 23.14
1067	Õ	ASN	149	127.676	12.401	47.571	1.00	25.30		1140	CG2	THR	158	115.044	13.781	39.959	1.00	5.29
1068	N	LEU	150	128.033	11.596	49.424	1.00	23.15		1141	С	THR	158	112.962	12.409	42.911	1.00	26.07
1069	CA	LEU	150	126.685	11.049	49.449	1.00	19.85		1142	O	THR	158	113.258	11.786	43.936	1.00	29.73
1070	CB	LEU	150	126.606	9.844	50.391	1.00	15.00	35	1143	N	HIS	159	111.745	12.362	42.373	1.00	25.85
1071 1072	CG CD1	LEU LEU	150 150	125.224 124.735	9.198 8.634	50.548 49.215	1.00 $1.00$	14.74 8.02		1144 1145	CA CB	HIS HIS	159 159	110.681 109.312	11.551 11.987	42.967 42.435	1.00 1.00	24.71 24.02
1072	CD2	LEU	150	125.287	8.115	51.600	1.00	2.00		1146	CG	HIS	159	108.903	13.358	42.872	1.00	20.05
1074	С	LEU	150	125.745	12.153	49.925	1.00	20.18		1147	CD2	HIS	159	108.888	14.538	42.209	1.00	14.66
1075	O	LEU	150	124.640	12.304	49.404	1.00	22.47		1148	ND1	HIS	159	108.453	13.629	44.147	1.00	22.06
1076	N	TYR	151	126.209	12.930	50.904	1.00	18.20	40	1149	CE1	HIS	159	108.179	14.917	44.250	1.00	20.70
1077 1078	CA CB	TYR TYR	151 151	125.440 126.226	14.041 14.739	51.455 52.569	1.00 $1.00$	18.43 10.57		1150 1151	NE2 C	HIS HIS	159 159	108.434 110.893	15.492 10.054	43.088 42.723	1.00 1.00	18.69 28.82
1079	CG	TYR	151	125.598	16.032	53.044	1.00	11.49		1151	Ö	HIS	159	110.377	9.211	43.464	1.00	29.90
1080	CD1	TYR	151	124.759	16.056	54.156	1.00	5.47		1153	N	ALA	160	111.674	9.733	41.695	1.00	22.36
1081	CE1	TYR	151	124.171	17.242	54.586	1.00	8.73		1154	CA	ALA	160	111.966	8.351	41.341	1.00	16.69
1082	CD2	TYR	151	125.835	17.234	52.372	1.00	10.72	45	1155	CB	ALA	160	112.118	8.233	39.835	1.00	12.78
1083 1084	CE2	TYR	151 151	125.250 124.421	18.421 18.420	52.791 53.898	$\frac{1.00}{1.00}$	9.08	73	1156 1157	C O	ALA	160 160	113.218 113.748	7.821 6.775	42.038 41.655	1.00 1.00	24.25 30.58
1084	CZ OH	TYR TYR	151	123.845	19.598	54.316	1.00	11.39 11.19		1157	N	ALA ASP	161	113.748	8.531	43.066	1.00	27.69
1086	C	TYR	151	125.117	15.041	50.355	1.00	17.39		1159	ĊA	ASP	161	114.880	8.134	43.800	1.00	24.25
1087	O	TYR	151	123.990	15.521	50.256	1.00	26.93		1160	CB	ASP	161	115.968	9.212	43.669	1.00	22.91
1088	N	GLU	152	126.121	15.374	49.552	1.00	15.23		1161	CG	ASP	161	116.356	9.495	42.222	1.00	29.35
1089	CA	GLU	152	125.937 127.282	16.316	48.455	1.00	18.45	50	1162	OD1	ASP	161	116.405	8.550	41.404	1.00	29.54
1090 1091	CB CG	GLU GLU	152 152	127.282	16.649 17.293	47.798 48.727	$\frac{1.00}{1.00}$	14.61 17.73		1163 1164	OD2 C	ASP ASP	$\frac{161}{161}$	116.623 114.626	10.672 7.840	41.906 45.281	$\frac{1.00}{1.00}$	19.16 $26.01$
1092	CD	GLU	152	127.962	18.712	49.169	1.00	16.74		1165	Õ	ASP	161	115.308	8.380	46.154	1.00	34.22
1093	OE1	GLU	152	126.980	19.292	48.662	1.00	16.63		1166	N	ASP	162	113.670	6.957	45.561	1.00	28.71
1094	OE2	GLU	152	128.681	19.252	50.034	1.00	22.78		1167	CA	ASP	162	113.339	6.590	46.939	1.00	28.70
1095	С	GLU	152	124.977	15.750	47.413	1.00	13.63	55	1168	CB	ASP	162	111.999	5.859	46.993	1.00	34.90
1096 1097	O N	GLU ALA	152 153	124.114 125.115	16.463 14.458	46.904 47.125	$\frac{1.00}{1.00}$	20.19 18.04		1169 1170	CG CD1	ASP ASP	162 162	110.851 110.426	6.726 7.600	46.536 47.322	$\frac{1.00}{1.00}$	42.80 36.44
1097	CA	ALA	153	123.113	13.778	46.143	1.00	14.37		1170	OD2	ASP	162	110.426	6.543	45.389	1.00	43.54
1099	CB	ALA	153	124.859	12.417	45.794	1.00	13.44		1172	C	ASP	162	114.423	5.728	47.573	1.00	27.14
1100	C	ALA	153	122.815	13.624	46.580	1.00	13.36		1173	O	ASP	162	114.386	5.441	48.769	1.00	28.39
1101	0	ALA	153	121.921	13.577	45.738	1.00	11.20	60	1174	N	ILE	163	115.378	5.302	46.756	1.00	26.67
1102	N	SER	154	122.574	13.568	47.889	1.00	16.02	50	1175	CA	ILE	163	116.485	4.486	47.229	1.00	27.39
1103 1104	CA CB	SER SER	154 154	121.218 121.250	13.413 13.157	48.416 49.928	1.00 $1.00$	13.22 8.73		1176 1177	CB CG2	ILE ILE	163 163	117.250 118.201	3.866 4.881	46.030 45.412	1.00 1.00	24.84 29.37
1104	ОG	SER	154	121.230	14.330	50.651	1.00	16.49		1177	OG1	ILE	163	118.201	2.623	46.471	1.00	30.57
1106	C	SER	154	120.312	14.607	48.118	1.00	14.26		1179	CD1	ILE	163	118.639	1.863	45.323	1.00	43.90
1107	O	SER	154	119.087	14.504	48.207	1.00	26.54	<i>c=</i>	1180	C	ILE	163	117.407	5.372	48.078	1.00	28.12
1108	N	HIS	155	120.915	15.735	47.757	1.00	14.00	65	1181	O	ILE	163	118.169	4.881	48.912	1.00	29.38
1109	CA	HIS	155	120.154	16.942	47.457	1.00	8.22		1182	N	LEU	164	117.277	6.686	47.890	1.00	30.46

TABLE 10-continued

s	tructura			of Tobacco				hase	· 5	s	tructura			of Tobacco l Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
1183	CA	LEU	164	118.070	7.684	48.607	1.00	24.33		1256	О	ILE	173	126.481	11.801	62.754	1.00	30.04
1184	CB	LEU	164	118.646	8.695	47.612	1.00	16.04	10	1257	N	HIS	174	126.404 127.757	10.972	60.662	1.00	18.99
1185 1186	CG CD1	LEU LEU	164 164	119.602 119.864	8.181 9.271	46.538 45.514	$\frac{1.00}{1.00}$	20.33 22.74	10	1258 1259	CA CB	HIS HIS	174 174	127.737	10.441 9.228	60.721 59.799	1.00 1.00	26.30 36.54
1187	CD2	LEU	164	120.894	7.720	47.184	1.00	17.92		1260	CG	HIS	174	127.114	8.034	60.257	1.00	41.37
1188	C	LEU LEU	164 164	117.259	8.441 9.518	49.658 50.101	$\frac{1.00}{1.00}$	23.88		1261	CD2 ND1	HIS HIS	174 174	126.355	7.147 7.644	59.571 61.579	1.00 1.00	35.65 39.38
1189 1190	N	GLU	165	117.667 116.120	7.882	50.101	1.00	32.16 21.56		1262 1263	CE1	HIS	174	127.057 126.295	6.569	61.687	1.00	33.20
1191	CA	GLU	165	115.256	8.529	51.043	1.00	18.89	15	1264	NE2	HIS	174	125.857	6.248	60.483	1.00	35.00
1192 1193	CB CG	GLU GLU	165 165	113.947 114.127	7.755 6.324	51.202 51.689	1.00 $1.00$	24.79 48.18		1265 1266	C O	HIS HIS	174 174	128.804 129.945	11.504 11.419	60.407 60.872	1.00 1.00	27.93 25.80
1193	CD	GLU	165	112.819	5.561	51.814	1.00	57.24		1267	N	LEU	175	128.410	12.508	59.626	1.00	25.88
1195	OE1	GLU	165	111.765	6.076	51.375	1.00	63.77		1268	CA	LEU	175	129.312	13.600	59.280	1.00	17.20
1196 1197	OE2 C	GLU	165 165	112.850	4.434	52.353 52.405	1.00 $1.00$	62.32		1269	CB CG	LEU LEU	175 175	128.804 129.069	14.376 13.747	58.066 56.696	1.00 1.00	12.92 4.37
1197	Ö	GLU GLU	165	115.907 115.598	8.727 9.687	53.106	1.00	16.81 18.04	20	1270 1271	CD1	LEU	175	128.472	14.624	55.606	1.00	2.00
1199	N	ASP	166	116.817	7.828	52.771	1.00	23.03		1272	CD2	LEU	175	130.566	13.572	56.482	1.00	6.69
1200 1201	CA CS	ASP ASP	166 166	117.497 117.383	7.914 6.579	54.061 54.811	1.00 $1.00$	27.94 35.23		1273 1274	C	LEU LEU	175 175	129.459 130.534	14.530 15.074	60.470 60.705	1.00 1.00	17.96 30.10
1201	OG	ASP	166	117.365	6.177	55.082	1.00	50.03		1274	N	GLU	176	128.375	14.699	61.225	1.00	17.67
1203	OD1	ASP	166	115.565	5.024	54.771	1.00	57.17	25	1276	CA	GLU	176	128.386	15.550	62.412	1.00	27.17
1204 1205	OD2 C	ASP	166 166	115.169	7.013 8.330	55.606 53.943	$\frac{1.00}{1.00}$	53.93 24.13	25	1277 1278	CB CG	GLU GLU	176 176	126.969 125.997	15.740 16.452	62.959 62.037	$\frac{1.00}{1.00}$	26.19 39.91
1205	Ö	ASP ASP	166	118.966 119.674	8.409	54.950	1.00	23.24		1279	CD	GLU	176	123.997	16.432	62.645	1.00	51.08
1207	N	ALA	167	119.401	8.638	52.721	1.00	14.78		1280	OE1	GLU	176	124.184	15.676	63.398	1.00	49.19
1208 1209	CA CB	ALA ALA	167 167	120.780 120.993	9.044 9.169	52.443 50.948	$\frac{1.00}{1.00}$	16.72 12.70		1281 1282	OE2 C	GLU GLU	176 176	123.932 129.241	17.600 14.913	62.368 63.505	$\frac{1.00}{1.00}$	52.29 29.58
1210	С	ALA	167	120.993	10.333	53.136	1.00	24.13	30	1283	Ö	GLU	176	129.241	15.604	64.237	1.00	29.36 37.29
1211	O	ALA	167	122.355	10.443	53.590	1.00	29.67		1284	N	SER	177	129.156	13.589	63.604	1.00	31.69
1212 1213	N CA	LEU LEU	168 168	120.317 120.614	11.313 12.590	53.193 53.831	$\frac{1.00}{1.00}$	27.80 19.27		1285 1286	CA CB	SER SER	177 177	129.883 129.310	12.816 11.395	64.607 64.678	$\frac{1.00}{1.00}$	28.71 24.94
1213	CB	LEU	168	119.540	13.623	53.487	1.00	23.80		1287	OG	SER	177	129.868	10.660	65.755	1.00	27.40
1215	CG	LEU	168	119.706	15.016	54.099	1.00	18.12		1288	C	SER	177	131.392	12.758	64.370	1.00	25.55
1216 1217	CD1 CD2	LEU LEU	168 168	121.006 118.524	15.642 15.890	53.626 53.719	1.00 $1.00$	19.21 17.36	35	1289 1290	O N	SER ALA	177 178	132.177 131.787	12.795 12.682	65.324 63.102	1.00 1.00	17.66 17.70
1217	CD2	LEU	168	120.730	12.450	55.343	1.00	20.39		1290	N CA	ALA	178	133.195	12.600	62.723	1.00	19.27
1219	O	LEU	168	121.663	12.973	55.943	1.00	26.94		1292	CB	ALA	178	133.330	11.789	61.441	1.00	22.41
1220 1221	N CA	ALA ALA	169 169	119.776 119.784	11.755 11.555	55.954 57.400	1.00 $1.00$	22.95 28.99		1293 1294	C O	ALA ALA	178 178	133.897 135.107	13.948 14.054	62.558 62.769	1.00 1.00	24.31 24.74
1222	CB	ALA	169	118.472	10.934	57.856	1.00	26.34	40	1295	N	ALA	179	133.107	14.975	62.214	1.00	25.32
1223	C	ALA	169	120.959	10.676	7.816	1.00	31.11	40	1296	CA	ALA	179	133.630	16.326	61.971	1.00	25.36
1224 1225	O N	ALA PHE	169 170	121.529 121.319	10.855 9.736	58.895 56.944	1.00 $1.00$	32.24 26.27		1297 1298	CB C	ALA ALA	179 179	132.460 134.658	17.289 16.943	61.779 62.928	1.00 1.00	32.63 23.67
1226	CA	PHE	170	122.423	8.819	57.197	1.00	22.01		1299	Ö	ALA	179	135.706	17.420	62.487	1.00	23.47
1227	CB	PHE	170	122.448	7.714	56.135	1.00	20.64		1300	N	PRO	180	134.384	16.929	64.244	1.00	21.48
1228 1229	CG CD1	PHE PHE	170 170	123.592 123.622	6.747 5.837	56.284 57.338	$\frac{1.00}{1.00}$	28.63 28.29	45	1301 1302	CD CA	PRO PRO	180 180	133.196 135.291	16.355 17.505	64.900 65.247	$\frac{1.00}{1.00}$	20.79 20.21
1230	OD2	PHE	170	124.642	6.745	55.368	1.00	25.46		1303	GB	PRO	180	134.601	17.155	66.568	1.00	10.68
1231	CE1	PHE	170	124.683	4.938	57.479	1.00	27.46		1304	CG	PRO	180	133.162		66.203	1.00	16.34
1232 1233	CE2 CZ	PHE PHE	170 170	125.706 125.726	5.850 4.945	55.500 56.558	$\frac{1.00}{1.00}$	24.24 22.35		1305 1306	C O	PRO PRO	180 180	136.747 137.623	17.040 17.772	65.264 65.722	$\frac{1.00}{1.00}$	21.05 30.38
1234	C	PHE	170	123.752	9.564	57.205	1.00	20.63		1307	N	HIS	181	137.015	15.846	64.750	1.00	24.41
1235	O	PHE	170	124.440	9.610	58.224	1.00	23.76	50	1308	CA	HIS	181	138.372	15.310	64.785	1.00	20.38
1236 1237	N CA	SER SER	171 171	124.095 125.340	10.156 10.899	56.066 55.918	$\frac{1.00}{1.00}$	16.15 12.93		1309 1310	CB CG	HIS HIS	181 181	138.359 137.686	13.955 13.989	65.498 66.837	$\frac{1.00}{1.00}$	22.19 20.19
1238	СВ	SER	171	125.476	11.429	54.488	1.00	12.97		1311	CD2	HIS	181	138.077	14.524	68.018	1.00	22.10
1239	OG	SER	171	124.397	12.281	54.152	1.00	12.50		1312	ND1	HIS	181	136.437	13.448	67.055	1.00	24.79
1240 1241	C	SER SER	171 171	125.479 126.567	12.047 12.297	56.912 57.420	$\frac{1.00}{1.00}$	13.92 15.50	55	1313 1314	CE1 NE2	HIS HIS	181 181	136.086 137.064	13.649 14.300	68.313 68.919	1.00 1.00	28.39 34.90
1242	N	THR	172	124.372	12.726	57.205	1.00	16.38	33	1315	С	HIS	181	139.073	15.184	63.443	1.00	16.56
1243	CA	THR	172	124.383	13.854	58.137	1.00	16.69		1316	O	HIS	181	140.138	14.575	63.351	1.00	20.78
1244 1245	CB OG1	THR THR	172 172	123.000 122.758	14.564 15.259	58.196 56.966	$\frac{1.00}{1.00}$	14.70 12.00		1317 1318	N CA	LEU LEU	182 182	138.496 139.095	15.775 15.698	62.407 61.082	1.00 1.00	19.38 19.81
1246	CG2	THR	172	122.946	15.559	59.348	1.00	6.76		1319	CB	LEU	182	138.023	15.838	59.999	1.00	12.64
1247	С	THR	172	124.813	13.486	59.556	1.00	19.18	60	1320	CG CD1	LEU	182	136.883	14.822	60.017	1.00	9.36
1248 1249	O N	THR ILE	172 173	125.759 124.129	14.067 12.516	60.086 60.160	1.00 $1.00$	22.93 22.35		1321 1322	CD1 CD2	LEU LEU	182 182	135.883 137.414	15.191 13.405	58.946 59.808	1.00 1.00	6.26 5.44
1250	CA	ILE	173	124.439	12.112	61.529	1.00	26.29		1323	С	LEU	182	140.164	16.760	60.884	1.00	23.01
1251	CB	ILE	173	123.428	11.061	62.070	1.00	29.19		1324	O	LEU	182	140.177	17.787	61.567	1.00	24.75
1252 1253	CG2 CG1	ILE ILE	173 173	123.553 123.657	9.747 10.644	61.305 63.572	1.00 $1.00$	26.89 34.59		1325 1326	N CA	LYS LYS	183 183	141.071 142.139	16.492 17.426	59.953 59.646	1.00 1.00	23.75 25.95
1254	CD1	ILE	173	122.655	9.928	64.240	1.00	35.92	65	1327	CB	LYS	183	143.300	16.712	58.948	1.00	25.08
1255	С	ILE	173	125.868	11.602	61.702	1.00	27.07		1328	CG	LYS	183	142.946	16.106	57.600	1.00	32.88

TABLE 10-continued

S	tructura			of Tobacco				hase	· 5	s	tructura			of Tobacco l Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
1329	CD	LYS	183	144.157	15.465	56.949	1.00	39.75		1402	CD2	HIS	192	130.320	25.503	52.428	1.00	14.30
1330	CE	LYS	183	143.804	14.905	55.581	1.00	45.64	10	1403	ND1	HIS	192	129.981	26.814	54.141	1.00	17.04
1331 1332	NZ C	LYS LYS	183 183	144.980 141.590	14.277 18.522	54.913 58.747	$\frac{1.00}{1.00}$	50.66 26.52	10	1404 1405	CE1 NE2	HIS HIS	192 192	129.488 129.681	27.425 26.651	53.078 52.025	1.00 1.00	22.40 9.94
1333	O	LYS	183	140.579	18.333	58.068	1.00	31.05		1406	C	HIS	192	129.061	23.321	54.939	1.00	15.72
1334 1335	N CA	SER	184 184	142.247 141.842	19.675 20.806	58.769 57.949	$\frac{1.00}{1.00}$	25.24 18.75		1407 1408	O N	HIS	192 193	128.002	23.896 22.085	54.696	1.00 1.00	19.37 22.40
1336	CB	SER SER	184	142.202	22.111	58.656	1.00	15.62		1408	CA	ALA ALA	193	129.331 128.367	21.288	54.524 53.766	1.00	17.47
1337	OG	SER	184	141.536	22.192	59.906	1.00	17.72	15	1410	CB	ALA	193	128.993	19.976	53.333	1.00	13.16
1338 1339	C O	SER SER	184 184	142.553 143.666	20.707 20.186	56.605 56.528	1.00 $1.00$	13.85 23.56		1411 1412	C O	ALA ALA	193 193	127.104 125.991	21.027 21.093	54.584 54.063	1.00 1.00	18.46 20.11
1340	N	PRO	185	141.930	21.221	55.526	1.00	14.69		1413	N	LEU	193	127.285	20.747	55.870	1.00	13.26
1341	CD	PRO	185	142.636	21.342	54.235	1.00	6.08		1414	CA	LEU	194	126.165	20.488	56.763	1.00	18.67
1342 1343	CA CB	PRO	185 185	140.622	21.886	55.462 54.213	1.00 $1.00$	13.32		1415	CB CG	LEU LEU	194 194	126.669 127.424	19.948	58.103 58.050	1.00 1.00	22.98
1343	СG	PRO PRO	185	140.758 141.553	22.747 21.860	53.309	1.00	7.14 4.25	20	1416 1417	CD1	LEU	194	127.424	18.615 18.249	59.439	1.00	17.83 22.28
1345	С	PRO	185	139.378	20.990	55.368	1.00	19.26		1418	CD2	LEU	194	126.526	17.524	57.494	1.00	9.41
1346 1347	O N	PRO LEU	185 186	138.268 139.547	21.502 19.671	55.198 55.478	1.00 $1.00$	22.11 15.43		1419 1420	C	LEU LEU	194 194	125.325 124.169	21.745 21.662	56.977 57.390	1.00 1.00	20.18 28.31
1347	CA	LEU	186	138.410	18.757	55.385	1.00	7.82		1421	N	GLU	195	125.913	22.906	56.701	1.00	23.69
1349	CB	LEU	186	138.859	17.304	55.533	1.00	10.45	25	1422	CA	GLU	195	125.217	24.182	56.845	1.00	23.91
1350	CG CD1	LEU LEU	186 186	137.743 137.199	16.259	55.379 53.953	$\frac{1.00}{1.00}$	14.68 2.00	25	1423 1424	CB CG	GLU GLU	195 195	126.145 126.558	25.235 24.930	57.459 58.897	1.00 $1.00$	28.38 48.57
1351 1352	CD1	LEU	186	137.199	16.257 14.885	55.744	1.00	10.39		1425	CD	GLU	195	120.556	25.905	59.449	1.00	61.92
1353	С	LEU	186	137.339	19.061	56.424	1.00	14.43		1426	OE1	GLU	195	128.341	26.515	58.652	1.00	66.87
1354 1355	O N	LEU ARG	186 187	136.147 137.774	19.062 19.318	56.114 57.653	1.00 $1.00$	15.11		1427 1428	OE2 C	GLU GLU	195 195	127.658 124.693	26.052 24.670	60.690 55.497	$\frac{1.00}{1.00}$	55.96
1356	CA	ARG	187	136.868	19.516	58.755	1.00	11.19 10.77	30	1429	Ö	GLU	195	123.721	25.422	55.436	1.00	14.54 17.77
1357	CB	ARG	187	137.675	19.929	60.019	1.00	9.64		1430	N	GLN	196	125.327	24.207	54.422	1.00	10.51
1358 1359	CG CD	ARG ARG	187 187	136.839 137.724	20.202 20.530	61.251 62.429	$\frac{1.00}{1.00}$	12.29 17.99		1431 1432	CA CB	GLN GLN	196 196	124.951 125.488	24.584 25.984	53.064 52.740	1.00 1.00	9.57 9.74
1360	NE	ARG	187	136.944	20.796	63.633	1.00	40.99		1433	CG	GLN	196	125.212	26.461	51.321	1.00	13.51
1361	CZ	ARG	187	137.468	20.982	64.841	1.00	53.63		1434	CD	GLN	196	123.737	26.672	51.051	1.00	18.27
1362 1363	NH1 NH2	ARG ARG	187 187	138.785 136.674	20.931 21.217	65.014 65.879	1.00 $1.00$	54.42 49.07	35	1435 1436	OE1 NE2	GLN GLN	196 196	123.111 123.174	27.556 25.862	51.633 50.162	1.00 1.00	31.94 21.35
1364	C	ARG	187	135.949	20.804	58.424	1.00	19.68		1437	C	GLN	196	125.174	23.583	52.039	1.00	8.87
1365	О	ARG	187	134.754	20.771	58.731	1.00	20.19		1438	O	GLN	196	126.695	23.481	51.830	1.00	16.57
1366 1367	N CA	GLU GLU	188 188	136.512 135.758	21.831 23.026	57.789 57.405	1.00 $1.00$	19.04 14.75		1439 1440	N CA	CYS CYS	197 197	124.577 124.963	22.837 21.865	51.415 550.398	1.00 1.00	11.72 13.36
1368	CB	GLU	188	136.708	24.179	57.052	1.00	18.38	40	1441	CB	CYS	197	123.821	20.882	50.114	1.00	20.25
1369	OG	GLU	188	137.416	24.825	58.248	1.00	25.44	40	1442	SG	CYS	197	122.310	21.605	49.432	1.00	16.36
1370 1371	CD OE1	GLU GLU	188 188	138.326 138.143	23.865 23.708	59.000 60.228	1.00 $1.00$	37.02 36.96		1443 1444	C	CYS CYS	197 197	125.351 124.948	22.614 23.758	49.126 48.924	1.00 $1.00$	11.32 16.69
1372	OE2	GLU	188	139.224	23.271	58.362	1.00	33.70		1445	N	LEU	198	126.134	21.965	48.274	1.00	14.40
1373	C	GLU	188	134.819	22.758	56.236	1.00	15.06		1446	CA	LEU	198	126.594	22.580	47.038	1.00	14.31
1374 1375	O N	GLU GLN	188 189	133.720 135.263	23.317 21.920	56.176 55.301	$\frac{1.00}{1.00}$	16.38 12.50	45	1447 1448	GB CG	LEU LEU	198 198	127.688 128.283	21.717 22.186	46.394 45.060	$\frac{1.00}{1.00}$	14.46 13.58
1376	CA	GLN	189	134.458	21.567	54.134	1.00	10.80		1449	CD1	LEU	198	128.949	23.541	45.225	1.00	9.55
1377	CB	GLN	189	135.269	20.711	53.153	1.00	10.38		1450	CD2	LEU	198	129.279	21.162	44.547	1.00	12.33
1378 1379	CG CD	GLN GLN	189 189	134.529 135.415	20.384 19.722	51.856 50.811	$\frac{1.00}{1.00}$	9.35 16.04		1451 1452	C	LEU LEU	198 198	125.478 125.389	22.848 23.945	46.034 45.481	$\frac{1.00}{1.00}$	18.46 26.64
1380	OE1	GLN	189	135.319	20.022	49.617	1.00	8.08		1453	N	HIS	199	124.614	21.857	45.829	1.00	20.84
1381	NE2	GLN	189	136.277	18.812	51.254	1.00	11.78	50	1454	CA	HIS	199	123.519	21.965	44.869	1.00	13.67
1382 1383	C O	GLN GLN	189 189	133.204 132.117	20.814 21.057	54.574 54.059	$\frac{1.00}{1.00}$	12.51 19.40		1455 1456	CB CG	HIS HIS	199 199	122.756 121.733	20.640 20.603	44.781 43.688	$\frac{1.00}{1.00}$	11.33 6.17
1384	N	VAL	190	133.363	19.920	55.546	1.00	11.87		1457	CD2	HIS	199	120.389	20.445	43.729	1.00	12.38
1385	CA	VAL	190	132.250	19.139	56.070	1.00	14.05		1458	ND1	HIS	199	122.061	20.738	42.356	1.00	10.08
1386 1387	CB CG1	VAL VAL	190 190	132.750 131.574	17.975 17.265	56.967 57.637	$\frac{1.00}{1.00}$	18.34 19.94	55	1459 1460	CE1 NE2	HIS HIS	199 199	120.964 119.935	20.663 20.486	41.624 42.432	1.00 1.00	10.18 2.01
1388	CG2	VAL	190	133.556	16.986	56.135	1.00	2.59	33	1461	С	HIS	199	122.540	23.111	45.108	1.00	14.52
1389	С	VAL	190	131.300	20.031	56.865	1.00	12.69		1462	O	HIS	199	122.174	23.813	44.166	1.00	13.39
1390 1391	O N	VAL THR	190 191	130.091 131.858	20.012 20.822	56.642 57.777	$\frac{1.00}{1.00}$	16.38 19.11		1463 1464	N CA	LYS LYS	200 200	122.120 121.161	23.300 24.353	46.357 46.698	1.00 1.00	15.92 14.65
1392	CA	THR	191	131.065	21.727	58.606	1.00	20.76		1465	CS	LYS	200	120.205	23.859	47.789	1.00	15.18
1393	CB OG1	THR	191	131.964	22.557	59.551	1.00	23.59	60	1466	CG	LYS	200	119.425	22.609	47.416	1.00	12.45
1394 1395	OG1 CG2	THR THR	191 191	132.681 131.130	21.675 23.511	60.424 60.391	1.00 $1.00$	29.20 29.68	-	1467 1468	CD CE	LYS LYS	200 200	118.523 117.827	22.158 20.849	48.554 48.213	1.00 1.00	3.23 14.66
1396	C	THR	191	130.241	22.664	57.731	1.00	19.82		1469	NZ	LYS	200	116.966	20.369	49.332	1.00	22.03
1397	O	THR	191	129.073	22.927	58.023	1.00	24.05		1470	С	LYS	200	121.786	25.688	47.120	1.00	22.11
1398 1399	N CA	HIS HIS	192 192	130.843 130.160	23.136 24.032	56.641 55.719	1.00 $1.00$	11.94 13.13		1471 1472	O N	LYS GLY	200 201	121.101 123.078	26.541 25.860	47.693 46.844	1.00 1.00	20.41 21.90
1400	CB	HIS	192	131.148	24.658	54.741	1.00	14.49	65	1473	CA	GLY	201	123.764	27.097	47.191	1.00	14.85
1401	CG	HIS	192	130.512	25.600	53.764	1.00	13.85		1474	С	GLY	201	124.048	27.969	45.978	1.00	16.72

TABLE 10-continued

S	tructura			of Tobacco				hase	. 5	s	tructura			of Tobacco l Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
1475	О	GLY	201	123.992	27.489	44.842	1.00	12.09		1548	CB	PHE	210	135.384	26.105	46.901	1.00	5.35
1476	N CA	VAL VAL	202 202	124.329 124.627	29.253 30.183	46.209 45.114	$\frac{1.00}{1.00}$	11.53	10	1549	CG CD1	PHE PHE	210 210	136.131	24.854 23.773	46.525 47.392	$\frac{1.00}{1.00}$	2.00 7.13
1477 1478	CB	VAL	202	124.627	31.661	45.555	1.00	11.31 9.93	10	1550 1551	CD1	PHE	210	136.182 136.794	24.763	47.392	1.00	13.50
1479	CG1	VAL	202	124.960	32.617	44.491	1.00	2.00		1552	CE1	PHE	210	136.883	22.617	47.052	1.00	13.12
1480	CG2	VAL VAL	202 202	122.964 126.054	31.937 29.940	45.803 44.612	$\frac{1.00}{1.00}$	6.34 10.17		1553 1554	CE2 CZ	PHE PHE	210 210	137.498 137.542	23.613 22.539	44.956 45.830	1.00 1.00	10.64 12.55
1481 1482	C O	VAL	202	126.034	29.883	45.405	1.00	8.95		1555	C	PHE	210	137.342	27.297	47.844	1.00	15.50
1483	N	PRO	203	126.222	29.774	43.286	1.00	2.10	15	1556	O	PHE	210	138.339	26.801	48.436	1.00	21.01
1484 1485	CD CA	PRO PRO	203 203	125.136 127.509	29.796 29.524	42.290 42.628	1.00 $1.00$	5.19 8.34		1557 1558	N CA	ILE ILE	211 211	137.500 138.798	28.290 28.834	46.970 46.601	1.00 1.00	11.89 14.18
1486	CS	PRO	203	127.168	29.704	41.154	1.00	5.82		1559	CB	ILE	211	138.663	30.000	45.604	1.00	13.95
1487	CG	PRO	203	125.785	29.152	41.087	1.00	2.00		1560	CG2	ILE	211	140.040	30.517	45.218	1.00	23.94
1488 1489	C O	PRO	203 203	128.699	30.381	43.069 43.516	1.00 $1.00$	18.54		1561	CG1	ILE	211 211	137.925	29.547	44.346 43.335	1.00 1.00	14.06
1490	N	PRO ARG	203	129.709 128.591	29.836 31.704	42.951	1.00	26.42 $11.17$	20	1562 1563	CD1 C	ILE ILE	211	137.734 139.622	30.656 29.318	47.790	1.00	19.84 11.48
1491	CA	ARG	204	129.687	32.582	43.357	1.00	5.94		1564	O	ILE	211	140.730	28.838	48.010	1.00	22.99
1492 1493	CB CG	ARG	204 204	129.366 129.405	34.047 34.440	43.061 41.587	1.00 $1.00$	4.29 10.69		1565 1566	N CA	SER SER	212 212	139.069 139.799	30.238 30.797	48.574 49.708	1.00 1.00	11.99 19.48
1493	CD	ARG ARG	204	130.821	34.543	41.033	1.00	8.35		1567	CB	SER	212	139.799	32.205	50.044	1.00	19.48
1495	NE	ARG	204	131.410	33.242	40.725	1.00	21.05	2.5	1568	OG	SER	212	137.939	32.174	50.500	1.00	32.56
1496	CZ	ARG	204	132.555	33.071	40.068	1.00	20.78	25	1569	С	SER	212	139.902	29.954	50.979	1.00	15.60
1497 1498	NH1 NH2	ARG ARG	204 204	133.250 132.996	34.121 31.844	39.644 39.818	$\frac{1.00}{1.00}$	14.75 16.69		1570 1571	O N	SER SER	212 213	140.992 138.785	29.800 29.398	51.530 51.437	$\frac{1.00}{1.00}$	26.35 18.79
1499	С	ARG	204	130.016	32.420	44.836	1.00	6.43		1572	CA	SER	213	138.780	28.607	52.665	1.00	15.21
1500	0	ARG	204	131.185	32.304	45.207	1.00	15.01		1573	CB	SER	213	137.426	28.737	53.372	1.00	13.39
1501 1502	N CA	VAL VAL	205 205	128.983 129.159	32.380 32.238	45.672 47.116	$\frac{1.00}{1.00}$	2.00 3.42	30	1574 1575	OG C	SER SER	213 213	137.168 139.141	30.074 27.126	53.766 52.543	$\frac{1.00}{1.00}$	19.66 21.19
1502	CB	VAL	205	127.809	32.238	47.855	1.00	2.00	30	1576	Õ	SER	213	139.540	26.503	53.534	1.00	22.69
1504	CG1	VAL	205	128.027	32.047	49.342	1.00	13.49		1577	N	ILE	214	139.021	26.558	51.345	1.00	16.22
1505 1506	CG2 C	VAL VAL	205 205	127.064 129.904	33.530 30.963	47.594 47.488	$\frac{1.00}{1.00}$	2.00 7.19		1578 1579	CA CB	ILE ILE	214 214	139.308 138.047	25.138 24.354	51.177 50.712	1.00 1.00	12.65 17.01
1507	Ö	VAL	205	130.785	30.982	48.342	1.00	17.39		1580	CG2	ILE	214	138.343	22.853	50.628	1.00	14.54
1508	N	GLU	206	129.543	29.854	46.851	1.00	11.33	35	1581	CG1	ILE	214	136.879	24.602	51.673	1.00	7.09
1509	CA	GLU	206	130.188	28.579	47.136	1.00	11.44		1582	CD1	ILE	214	137.175	24.247	53.124	1.00	2.16
1510 1511	CB CG	GLU GLU	206 206	129.348 128.033	27.417 27.237	46.606 47.340	$\frac{1.00}{1.00}$	10.75 4.42		1583 1584	C O	ILE ILE	214 214	140.477 141.486	24.759 24.247	50.276 50.759	1.00 1.00	14.51 20.94
1512	CD	GLU	206	128.208	27.226	48.845	1.00	7.68		1585	N	TYR	215	140.342	25.006	48.975	1.00	10.71
1513	OE1	GLU	206	128.858	26.298	49.366	1.00	15.79		1586	CA	TYR	215	141.378	24.634	48.016	1.00	16.76
1514 1515	OE2 C	GLU GLU	206 206	127.700 131.598	28.153 28.528	49.509 46.568	1.00 $1.00$	12.42 13.88	40	1587 1588	CB CG	TYR TYR	215 215	140.914 141.523	24.914 23.975	46.587 45.569	1.00 1.00	10.15 16.49
1516	O	GLU	206	132.484	27.897	47.144	1.00	16.22		1589	CD1	TYR	215	141.526	22.595	45.777	1.00	14.36
1517	N	THR	207	131.792	29.199	45.438	1.00	9.73		1590	CE1	TYR	215	142.079	21.722	44.837	1.00	16.84
1518 1519	CA CB	THR THR	207 207	133.090 132.970	29.268 29.928	44.785 43.400	1.00 $1.00$	13.84 15.72		1591 1592	CD2 CE2	TYR TYR	215 215	142.090 142.645	24.463 23.601	44.393 43.447	1.00 1.00	18.93 14.61
1520	OG1	THR	207	132.272	29.045	42.513	1.00	13.12		1593	CZ	TYR	215	142.636	22.232	43.676	1.00	18.72
1521	CG2	THR	207	134.338	30.250	42.827	1.00	11.92	45	1594	OH	TYR	215	143.191	21.375	42.749	1.00	21.97
1522 1523	C O	THR THR	207 207	134.059 135.177	30.066 29.623	45.658 45.917	1.00 $1.00$	19.41 27.17		1595 1596	C	TYR TYR	215 215	142.753 143.772	25.251 24.567	48.256 48.154	1.00 1.00	22.56 21.88
1524	Ň	ARG	208	133.608	31.226	46.133	1.00	22.15		1597	N	ASP	216	142.780	26.538	48.582	1.00	25.86
1525	CA	ARG	208	134.417	32.091	46.988	1.00	17.49		1598	CA	ASP	216	144.032	27.239	48.841	1.00	28.90
1526 1527	CB CG	ARG ARG	208 208	133.595 134.349	33.309 34.352	47.429 48.264	$\frac{1.00}{1.00}$	23.71 24.20	50	1599 1600	CB CG	ASP ASP	216 216	143.745 145.000	28.708 29.514	49.155 49.373	$\frac{1.00}{1.00}$	35.55 32.91
1528	CD	ARG	208	135.532	34.935	47.501	1.00	35.41		1601	OD1	ASP	216	145.170	30.035	50.494	1.00	33.80
1529	NE	ARG	208	136.060	36.169	48.090	1.00	40.81		1602	OD2	ASP	216	145.811	29.630	48.427	1.00	38.29
1530 1531	CZ NH1	ARG ARG	208 208	136.736 136.978	36.242 35.150	49.235 49.947	$\frac{1.00}{1.00}$	40.71 42.82		1603 1604	C O	ASP ASP	216 216	144.782 146.013	26.590 26.552	50.002 50.017	$\frac{1.00}{1.00}$	28.68 35.11
1532	NH2	ARG	208	137.194	37.412	49.658	1.00	37.98		1605	N	LYS	217	144.026	26.052	50.954	1.00	24.04
1533	C	ARG	208	134.906	31.313	48.208	1.00	19.76	55	1606	CA	LYS	217	144.604	25.406	52.129	1.00	25.31
1534 1535	O N	ARG PHE	208 209	136.075 134.010	31.395 30.534	48.576 48.809	$\frac{1.00}{1.00}$	27.03 16.90		1607 1608	CB CG	LYS LYS	217 217	143.768 143.687	25.741 27.234	53.368 53.646	$\frac{1.00}{1.00}$	17.45 28.42
1536	CA	PHE	209	134.350	29.734	49.979	1.00	12.93		1609	CD	LYS	217	142.811	27.551	54.844	1.00	36.77
1537	CB	PHE	209	133.090	29.165	50.632	1.00	3.91		1610	CE	LYS	217	142.729	29.057	55.071	1.00	36.92
1538 1539	CG CD1	PHE PHE	209 209	133.377 133.605	28.292 28.852	51.818 53.070	$\frac{1.00}{1.00}$	8.27 5.35		1611 1612	NZ C	LYS LYS	217 217	141.758 144.754	29.426 23.892	56.143 51.973	1.00 1.00	37.10 25.97
1540	CD2	PHE	209	133.472	26.912	51.676	1.00	11.24	60	1613	Ö	LYS	217	145.170	23.201	52.905	1.00	23.71
1541	CE1	PHE	209	133.928	28.052	54.162	1.00	10.99		1614	N	GLU	218	144.429	23.385	50.788	1.00	30.40
1542 1543	CE2 CZ	PHE PHE	209 209	133.794 134.023	26.105 26.677	52.760 54.007	1.00 $1.00$	7.23 2.00		1615 1616	CA CB	GLU GLU	218 218	144.528 143.655	21.958 21.603	50.504 49.297	1.00 1.00	34.37 41.93
1544	C	PHE	209	135.305	28.581	49.664	1.00	16.94		1617	CG	GLU	218	143.462	20.114	49.297	1.00	45.33
1545	O	PHE	209	136.176	28.248	50.473	1.00	13.43	65	1618	CD	GLU	218	142.740	19.451	50.226	1.00	52.39
1546 1547	N CA	PHE PHE	210 210	135.112 135.960	27.942 26.823	48.514 48.126	1.00 $1.00$	11.91 12.01	65	1619 1620	OE1 OE2	GLU GLU	218 218	141.553 143.364	19.780 18.612	50.449 50.916	1.00 1.00	49.41 46.07
157/	CA	LIL	210	100.700	20.020	70.120	1.00	12.01		1020	OL2	OLU	210	170.007	10.012	50.510	1.00	70.07

TABLE 10-continued

s	tructura			of Tobacco				hase	5	s	tructura			of Tobacco				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
1621	С	GLU	218	145.982	21.585	50.228	1.00	34.68		1694	NH2	ARG	227	149.963	35.561	38.831	1.00	19.58
1622 1623	O N	GLU GLN	218 219	146.624 146.493	22.166 20.611	49.356 50.974	$\frac{1.00}{1.00}$	31.92 37.87	10	1695 1696	C O	ARG ARG	227 227	144.911 144.475	30.745 31.894	39.367 39.402	1.00 1.00	17.48 22.08
1624	CA	GLN	219	140.493	20.011	50.827	1.00	41.61	10	1697	N	PHE	228	144.474	29.822	38.516	1.00	22.08 17.49
1625	CB	GLN	219	148.180	19.105	51.896	1.00	52.08		1698	CA	PHE	228	143.439	30.073	37.516	1.00	20.78
1626 1627	CG CD	GLN GLN	219 219	149.617 149.709	18.615 17.102	51.900 51.943	$\frac{1.00}{1.00}$	67.09 78.27		1699 1700	CB CG	PHE PHE	228 228	143.184 142.261	28.770 28.908	36.741 35.556	1.00 1.00	17.58 10.74
1628	OE1	GLN	219	149.709	16.470	52.917	1.00	81.71		1701	CD1	PHE	228	142.201	30.128	35.214	1.00	18.32
1629	NE2	GLN	219	150.233	16.510	50.870	1.00	80.10	15	1702	CD2	PHE	228	141.958	27.791	34.785	1.00	13.57
1630 1631	C O	GLN GLN	219 219	148.173 149.260	19.583 19.789	49.438 48.893	1.00 $1.00$	38.79 34.89		1703 1704	CE1 CE2	PHE PHE	228 228	140.819 141.095	30.230 27.883	34.122 33.692	1.00 1.00	18.76 18.93
1632	N	SER	220	147.205	18.867	48.874	1.00	39.35		1704	CZ	PHE	228	140.525	29.106	33.361	1.00	12.22
1633	CA	SER	220	147.359	18.252	47.556	1.00	36.65		1706	C	PHE	228	142.158	30.552	38.205	1.00	20.33
1634 1635	CB OG	SER SER	220 220	146.658 145.291	16.891 17.018	47.537 47.899	1.00 $1.00$	48.69 58.57		1707 1708	O N	PHE ALA	228 229	141.585 141.746	31.580 29.817	37.834 39.233	1.00 1.00	17.12 15.85
1636	C	SER	220	146.824	19.117	46.418	1.00	30.45	20	1709	CA	ALA	229	140.541	30.125	39.989	1.00	14.57
1637	O	SER	220	146.651	18.639	45.297	1.00	31.98		1710	CB	ALA	229	140.320	29.069	41.059	1.00	11.49
1638 1639	N CA	LYS LYS	221 221	146.581 146.052	20.392 21.327	46.704 45.716	1.00 $1.00$	25.39 18.20		1711 1712	C	ALA ALA	229 229	140.572 139.606	31.513 32.272	40.619 40.505	1.00 1.00	22.21 26.82
1640	CB	LYS	221	145.949	22.731	46.316	1.00	17.94		1712	N	LYS	230	141.683	31.841	41.278	1.00	20.82 17.14
1641	CG	LYS	221	147.292	23.352	46.659	1.00	24.63	25	1714	CA	LYS	230	141.836	33.136	41.933	1.00	13.72
1642 1643	CD CE	LYS LYS	221 221	147.136 148.444	24.772 25.312	47.155 47.716	$\frac{1.00}{1.00}$	32.69 42.07	25	1715	CB CG	LYS LYS	230 230	143.118 143.067	33.168 32.332	42.766 44.030	1.00 1.00	17.71 14.07
1644	NZ	LYS	221	149.547	25.257	46.720	1.00	42.25		1716 1717	CD	LYS	230	144.343	32.505	44.835	1.00	23.37
1645	С	LYS	221	146.879	21.412	44.444	1.00	15.22		1718	CE	LYS	230	144.253	31.802	46.177	1.00	31.01
1646	O	LYS	221	148.097	21.243	44.467	1.00	21.32		1719	NZ	LYS	230	145.477	32.021	46.994	1.00	32.57
1647 1648	N CA	ASN ASN	222 222	146.196 146.853	21.660 21.818	43.333 42.048	$\frac{1.00}{1.00}$	11.20 9.54	30	1720 1721	C	LYS LYS	230 230	141.816 141.111	23.310 35.292	40.956 41.176	1.00 $1.00$	16.07 15.73
1649	СВ	ASN	222	145.993	21.250	40.919	1.00	2.46	20	1722	N	LEY	231	142.585	34.202	39.876	1.00	17.93
1650	CG	ASN	222	146.599	21.488	39.550	1.00	12.07		1723	CA	LEU	231	142.646	35.260	38.872	1.00	20.80
1651 1652	OD1 ND2	ASN ASN	222 222	146.698 147.003	22.626 20.414	39.097 38.881	$\frac{1.00}{1.00}$	10.03 11.12		1724 1725	CB CG	LEU LEU	231 231	143.653 145.141	34.911 34.870	37.775 38.116	1.00 1.00	18.38 16.32
1653	C	ASN	222	147.032	23.322	41.885	1.00	17.59		1726	CD1	LEU	231	145.920	34.518	36.863	1.00	14.82
1654	0	ASN	222	146.060	24.061	41.717	1.00	21.24	35	1727	CD2	LEU	231	145.593	36.212	38.658	1.00	15.83
1655 1656	N CA	ASN ASN	223 223	148.281 148.619	23.765 25.175	41.958 41.850	1.00 $1.00$	18.52 9.91		1728 1729	C O	LEU LEU	231 231	141.287 140.828	35.506 36.647	38.233 38.151	1.00 1.00	24.81 28.14
1657	CB	ASN	223	150.127	25.349	41.972	1.00	11.52		1730	N	ASP	232	140.648	34.427	37.790	1.00	28.52
1658	CG	ASN	223	150.664	24.821	43.282	1.00	23.77		1731	CA	ASP	232	139.344	34.503	37.139	1.00	23.60
1659 1660	OD1 ND2	ASN ASN	223 223	150.579 151.208	25.491 23.605	44.311 43.258	1.00 $1.00$	21.40 21.52		1732 1733	CB CG	ASP ASP	232 232	138.878 137.737	33.104 33.137	36.736 35.742	1.00 1.00	20.22 29.93
1661	C C	ASN	223	148.104	25.870	40.594	1.00	15.35	40	1734	OD1	ASP	232	138.019	33.223	34.527	1.00	24.38
1662	O	ASN	223	147.668	27.019	40.662	1.00	21.88		1735	OD2	ASP	232	136.564	33.079	36.174	1.00	26.36
1663 1664	N CA	VAL VAL	224 224	148.157 147.677	25.184 25.755	39.455 38.195	1.00 $1.00$	10.58 18.60		1736 1737	C	ASP ASP	232 232	138.300 137.622	35.170 36.111	38.032 37.612	1.00 1.00	20.67 15.86
1665	CB	VAL	224	147.957	24.811	37.001	1.00	24.63		1738	N	PHE	233	137.022	34.707	39.274	1.00	12.27
1666	CG1	VAL	224	147.405	25.406	35.709	1.00	25.94	15	1739	CA	PHE	233	137.244	35.260	40.219	1.00	12.59
1667 1668	CG2 C	VAL VAL	224 224	149.449 146.177	24.558 26.049	36.873 38.259	$\frac{1.00}{1.00}$	17.27 23.24	45	1740 1741	CB CG	PHE PHE	233 233	137.355 136.243	34.549 34.887	41.569 42.524	1.00 1.00	14.20 30.75
1669	Ö	VAL	224	145.716	27.071	37.740	1.00	25.12		1742	CD1	PHE	233	135.130	34.058	42.634	1.00	33.32
1670	N	LEU	225	145.423	25.146	38.886	1.00	23.80		1743	CD2	PHE	233	136.298	36.042	43.302	1.00	29.41
1671 1672	CA CB	LEU LEU	225 225	143.980 143.314	25.313 23.994	39.032 39.434	1.00 $1.00$	18.51 18.23		1744 1745	CE1 CE2	PHE PHE	233 233	134.088 135.263	34.372 36.365	43.502 44.172	$\frac{1.00}{1.00}$	34.63 31.81
1673	СБ	LEU	225	143.337	23.994	38.424	1.00	20.30	50	1745	CE2	PHE	233	133.203	35.528	44.172	1.00	32.38
1674	CD1	LEU	225	142.613	21.645	39.010	1.00	18.77		1747	С	PHE	233	137.452	36.760	40.407	1.00	18.84
1675 1676	CD2 C	LEU LEU	225 225	142.691 143.652	23.270 26.392	37.115 40.061	$\frac{1.00}{1.00}$	10.44 18.24		1748 1749	O N	PHE ASN	233 234	136.495 138.710	37.534 37.160	40.394 40.572	1.00 1.00	24.79 25.72
1677	Ö	LEU	225	143.032	26.392	39.872	1.00	22.06		1750	CA	ASN	234	138.710	38.563	40.372	1.00	20.56
1678	N	LEU	226	144.431	26.448	41.141	1.00	18.51		1751	CB	ASN	234	140.509	38.694	41.239	1.00	25.13
1679 1680	CA CB	LEU LEU	226 226	144.230 145.128	27.445 27.149	42.197 43.401	$\frac{1.00}{1.00}$	17.64 15.88	55	1752 1753	CG OD1	ASN ASN	234 234	140.702 139.738	38.249 37.996	42.683 43.406	1.00 1.00	30.65 21.26
1681	CG	LEU	226	145.013	28.096	44.605	1.00	19.63		1754	ND2	ASN	234	141.957	38.162	43.109	1.00	35.99
1682	CD1	LEU	226	143.633	27.996	45.235	1.00	8.24		1755	C	ASN	234	138.818	39.427	39.536	1.00	18.33
1683	CD2	LEU	226	146.086	27.764	45.627	1.00	2.89		1756	O N	ASN	234	138.457	40.599	39.662	1.00	15.79 17.45
1684 1685	C	LEU LEU	226 226	144.507 143.753	28.855 29.785	41.681 41.966	$\frac{1.00}{1.00}$	15.54 28.19		1757 1758	N CA	LEU LEU	235 235	139.019 138.814	38.848 39.567	38.353 37.097	1.00 1.00	17.45 16.13
1686	N	ARG	227	145.595	29.002	40.927	1.00	22.25	60	1759	CB	LEU	235	139.402	38.789	35.920	1.00	14.09
1687	CA	ARG	227	145.995	30.282	40.338	1.00	21.24		1760	CG CD1	LEU	235	139.233	39.426	34.534	1.00	25.16
1688 1689	CB CG	ARG ARG	227 227	147.320 147.831	30.108 31.335	39.587 38.844	1.00 $1.00$	25.23 26.56		1761 1762	CD1 CD2	LEU LEU	235 235	139.947 139.762	40.774 38.487	34.472 33.458	1.00 1.00	16.26 11.51
1690	CD	ARG	227	148.575	32.292	39.760	1.00	33.23		1763	С	LEU	235	137.329	39.812	36.866	1.00	23.53
1691	NE CZ	ARG	227	149.114	33.433	39.021	1.00	27.48	65	1764	O	LEU	235	136.929	40.918	36.502	1.00	30.18
1692 1693	CZ NH1	ARG ARG	227 227	149.516 149.447	34.568 34.722	39.585 40.902	1.00 $1.00$	29.44 31.95	05	1765 1766	N CA	LEU LEU	236 236	136.517 135.071	38.773 38.894	37.065 36.900	1.00 1.00	26.97 21.97
1070	. 1221				22		2.00	- 1.70		2,00	~ 1			220.071	231077	20.000	2.00	-4.27

TABLE 10-continued

S	tructura			of Tobacco				hase	5	s	tructura			of Tobacco				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor	_	Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
1767	CB	LEU	236	134.375	37.538	37.041	1.00	22.99		1840	CD2	LEU	244	127.780	46.582	43.908	1.00	9.05
1768	CG CD1	LEU	236 236	134.550	36.506	35.931	1.00	24.74	10	1841	С	LEU	244 244	127.213	49.191	40.335	1.00	22.78
1769 1770	CD1 CD2	LEU LEU	236	133.601 134.259	35.347 37.133	36.187 34.579	$\frac{1.00}{1.00}$	22.83 26.05	10	1842 1843	O N	LEU ALA	244	126.328 128.325	49.868 49.725	40.863 39.838	1.00 1.00	27.46 31.72
1771	C	LEU	236	134.511	39.858	37.935	1.00	15.40		1844	CA	ALA	245	128.560	51.167	39.856	1.00	30.71
1772 1773	O N	LEU GLN	236 237	133.581 135.080	40.602 39.837	37.646 39.139	$\frac{1.00}{1.00}$	21.06 13.42		1845 1846	CB C	ALA	245 245	129.998 127.589	51.476 51.860	39.466 38.893	1.00 1.00	26.12 29.98
1774	CA	GLN	237	134.645	40.721	40.217	1.00	14.55		1847	Ö	ALA ALA	256	127.005	52.887	39.226	1.00	32.44
1775	CB	GLN	237	135.477	40.481	41.475	1.00	14.51	15	1848	N	GLN	246	127.410	51.256	37.718	1.00	29.03
1776 1777	CG CD	GLN GLN	237 237	135.051 135.967	41.318 41.131	42.671 43.862	1.00 $1.00$	12.72 $12.21$		1849 1850	CA CB	GLN GLN	246 246	126.528 126.689	51.754 50.868	36.668 35.430	1.00 1.00	31.14 31.92
1778	OE1	GLN	237	137.121	41.131	43.847	1.00	18.76		1851	CG	GLN	246	125.845	51.244	34.232	1.00	41.36
1779	NE2	GLN	237	135.460	40.483	44.900	1.00	4.88		1852	CD	GLN	246	125.970	50.235	33.109	1.00	48.09
1780 1781	C	GLN	237 237	134.760	42.180	39.788 40.192	1.00 $1.00$	22.23 28.74		1853	OE1 NE2	GLN	246 246	127.023	49.627 50.043	32.919 32.361	1.00 1.00	49.29
1781	N	GLN MET	238	133.950 135.770	43.011 42.481	38.970	1.00	29.92	20	1854 1855	C	GLN GLN	246	124.887 125.074	51.762	37.119	1.00	55.11 34.84
1783	CA	MET	238	135.985	43.833	38.458	1.00	23.66		1856	O	GLN	246	124.297	52.637	36.732	1.00	42.31
1784 1785	CB CG	MET MET	238 238	137.275 138.552	43.906 43.791	37.638 38.454	1.00 $1.00$	25.72 26.38		1857 1858	N CA	VAL VAL	247 247	124.719 123.360	50.762 50.631	37.921 38.441	1.00 1.00	39.27 38.03
1786	SD	MET	238	140.030	43.684	37.408	1.00	30.42		1859	CB	VAL	247	123.360	49.138	38.742	1.00	38.04
1787	CE	MET	238	141.233	43.058	38.580	1.00	23.83	25	1860	CG1	VAL	247	122.330	48.954	40.059	1.00	39.31
1788 1789	С	MET MET	238 238	134.801 134.344	44.227 45.367	38.584 37.628	$\frac{1.00}{1.00}$	20.92 20.70	25	1861 1862	CG2	VAL VAL	247 247	122.270 123.144	48.526 51.507	37.603 39.667	$\frac{1.00}{1.00}$	37.60 37.49
1790	O N	LEU	239	134.344	43.274	36.792	1.00	23.17		1863	C O	VAL	247	123.144	51.862	39.998	1.00	33.11
1791	CA	LEU	239	133.159	43.509	35.920	1.00	21.15		1864	N	SER	248	124.231	51.871	40.340	1.00	39.80
1792 1793	CB CG	LEU LEU	239 239	132.938	42.323	34.978	1.00 $1.00$	10.92 20.20		1865	CA	SER SER	248 248	124.173	52.736	41.515 42.352	$\frac{1.00}{1.00}$	43.54 41.35
1793	CD1	LEU	239	131.684 131.748	42.381 43.579	34.100 33.166	1.00	9.66	30	1866 1867	CB OG	SER	248	125.456 125.482	52.604 51.368	43.057	1.00	23.81
1795	CD2	LEU	239	131.541	41.089	33.309	1.00	10.55		1868	С	SER	248	123.963	54.195	41.093	1.00	41.56
1796 1797	C O	LEU LEU	239 239	131.908 131.129	43.732 44.645	36.764 36.501	$\frac{1.00}{1.00}$	14.68 23.63		1869 1870	O N	SER ARG	248 249	123.288 124.591	54.976 54.559	41.783 39.974	$\frac{1.00}{1.00}$	39.28 40.01
1798	N	HIS	240	131.735	42.904	37.788	1.00	15.07		1871	CA	ARG	249	124.391	55.901	39.421	1.00	47.97
1799	CA	HIS	240	130.587	43.015	38.680	1.00	19.33		1872	CB	ARG	249	125.475	56.127	38.290	1.00	51.84
1800 1801	CB CG	HIS HIS	240 240	130.619 130.661	41.913 40.525	39.746 39.185	1.00 $1.00$	16.76 $12.10$	35	1873 1874	CG CD	ARG ARG	249 249	126.912 127.836	56.292 56.501	38.747 37.563	1.00 1.00	62.62 69.45
1802	CD2	HIS	240	130.001	40.039	37.973	1.00	12.10		1875	NE	ARG	249	129.065	55.731	37.713	1.00	78.07
1803	ND1	HIS	240	131.144	39.449	39.897	1.00	8.57		1876	CZ	ARG	249	129.491	54.824	36.840	1.00	81.11
1804 1805	CE1 NE2	HIS HIS	240 240	131.077 130.567	38.362 38.692	39.150 37.979	1.00 $1.00$	16.05 13.30		1877 1878	NH1 NH2	ARG ARG	249 249	128.795 130.602	54.570 54.145	35.737 37.087	1.00 1.00	75.69 86.20
1805	C	HIS	240	130.610	44.383	39.344	1.00	21.46	40	1879	C	ARG	249	123.051	56.058	38.890	1.00	43.68
1807	0	HIS	240	129.572	45.034	39.481	1.00	28.61	40	1880	0	ARG	249	122.402	57.072	39.127	1.00	45.81
1808 1809	N CA	LYS LYS	241 241	131.809 132.008	44.819 46.118	39.728 40.364	1.00 $1.00$	24.98 19.85		1881 1882	N CA	TRP TRP	250 250	122.588 121.247	55.030 55.000	38.183 37.613	1.00 1.00	38.57 34.46
1810	CB	LYS	241	133.469	46.281	40.782	1.00	18.41		1883	CB	TRP	250	121.060	53.682	36.851	1.00	37.42
1811	CG	LYS	241	133.855	45.556	42.057	1.00	18.41		1884	CG	TRP	250	119.635	53.325	36.531	1.00	37.50
1812 1813	CD CE	LYS LYS	241 241	135.348 135.667	45.688 45.861	42.297 43.769	$\frac{1.00}{1.00}$	29.39 43.69	45	1885 1886	CD2 CE2	TRP TRP	250 250	118.745 117.516	52.545 52.464	37.341 36.652	$\frac{1.00}{1.00}$	32.98 39.28
1814	NZ	LYS	241	137.131	45.987	43.991	1.00	46.04		1887	CE3	TRP	250	118.867	51.909	38.585	1.00	32.90
1815	C	LYS	241	131.604	47.252	39.419	1.00	23.69		1888	CD1	TRP	250	118.931	53.672	35.413	1.00	28.29
1816 1817	O N	LYS GLN	241 242	130.983 131.954	48.229 47.107	39.845 38.140	$\frac{1.00}{1.00}$	14.81 20.70		1889 1890	NE1 CZ2	TRP TRP	250 250	117.658 116.411	53.159 51.771	35.479 37.167	$\frac{1.00}{1.00}$	36.07 40.28
1818	CA	GLN	242	131.615	48.099	37.120	1.00	28.94		1891	CZ3	TRP	250	117.770	51.221	39.098	1.00	33.00
1819	CB	GLN	242	132.262	47.748	35.775	1.00	29.06	50	1892	CH2	TRP	250	116.557	51.159	38.388	1.00	34.10
1820 1821	CG CD	GLN GLN	242 242	133.775 134.359	47.862 47.517	35.748 34.392	$\frac{1.00}{1.00}$	39.14 42.97		1893 1894	C O	TRP TRP	250 250	120.215 119.207	55.131 55.820	38.731 38.583	$\frac{1.00}{1.00}$	31.76 38.62
1822	OE1	GLN	242	134.324	46.363	33.962	1.00	48.36		1895	N	TRP	251	120.499	54.493	39.861	1.00	29.34
1823	NE2	GLN	242	134.904	48.519	33.710	1.00	39.66		1896	CA	TRP	251	119.611	54.513	41.017	1.00	28.23
1824 1825	C O	GLN GLN	242 242	130.103 129.514	48.163 49.246	36.943 36.938	1.00 $1.00$	33.40 40.80	55	1897 1898	CB CG	TRP TRP	251 251	120.041 119.164	53.431 53.309	42.003 43.196	1.00 1.00	24.43 27.54
1826	N	GLU	243	129.487	46.992	36.807	1.00	33.80	33	1899	CD2	TRP	251	117.813	52.824	43.224	1.00	28.47
1827	CA	GLU	243	128.044	46.884	36.631	1.00	22.54		1900	CE2	TRP	251	117.394	52.848	44.576	1.00	30.61
1828 1829	CB CG	GLU GLU	243 243	127.647 128.204	45.420 44.778	36.466 35.210	$\frac{1.00}{1.00}$	15.53 15.70		1901 1902	CE3 CD1	TRP TRP	251 251	116.921 119.493	52.373 53.603	42.246 44.486	1.00 1.00	22.56 29.85
1830	CD	GLU	243	127.938	43.290	35.137	1.00	18.78		1903	NE1	TRP	251	118.439	53.327	45.321	1.00	28.21
1831	OE1	GLU	243	127.639	42.675	36.178	1.00	18.01	60	1904	CZ2	TRP	251	116.117	52.434	44.974	1.00	27.80
1832 1833	OE2 C	GLU GLU	243 243	128.040 127.290	42.727 47.495	34.032 37.806	1.00 $1.00$	15.89 20.49	-	1905 1906	CZ3 CH2	TRP TRP	251 251	115.646 115.261	51.959 51.995	42.641 43.993	1.00 1.00	16.40 23.32
1834	Ö	GLU	243	126.351	48.266	37.611	1.00	18.81		1907	C	TRP	251	119.575	55.877	41.703	1.00	37.98
1835	N	LEU	244	127.715	47.159	39.022	1.00	12.97		1908	O	TRP	251	118.564	56.243	42.309	1.00	45.16
1836 1837	CA CB	LEU LEU	244 244	127.079 127.676	47.675 46.999	40.231 41.467	1.00 $1.00$	15.01 12.34		1909 1910	N CA	LYS LYS	252 252	120.685 120.758	56.610 57.938	41.636 42.238	1.00 1.00	42.00 42.12
1838	CG	LEU	244	127.144	47.436	42.832	1.00	12.62	65	1911	CB	LYS	252	122.198	58.328	42.551	1.00	47.95
1839	CD1	LEU	244	125.628	47.332	42.881	1.00	24.09		1912	CG	LYS	252	122.598	58.028	43.975	1.00	54.42

TABLE 10-continued

s	tructura			of Tobacco				hase	5	S	tructura			of Tobacco l Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor	_	Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
1913	CD	LYS	252	123.720	58.943	44.418	1.00	63.61		1986	CG	PRO	261	109.865	59.476	51.972	1.00	61.83
1914 1915	CE NZ	LYS LYS	252 252	123.889 124.827	58.896 59.944	45.923 46.407	$\frac{1.00}{1.00}$	76.47 82.00	10	1987 1988	C O	PRO PRO	261 261	110.685	56.401 55.811	52.265 53.143	$\frac{1.00}{1.00}$	56.67 54.94
1915	C	LYS	252	124.827	58.992	41.348	1.00	43.32	10	1989	N	TYR	262	111.317 109.475	56.019	51.869	1.00	55.28
1917	O	LYS	252	119.528	59.955	41.845	1.00	40.24		1990	CA	TYR	262	108.813	54.855	52.455	1.00	52.74
1918 1919	N CA	ASP ASP	253 253	120.220 119.621	58.802 59.715	40.033 39.061	$\frac{1.00}{1.00}$	41.81 42.20		1991 1992	CB CG	TYR TYR	262 262	107.309 106.954	54.898 54.869	52.167 50.695	1.00 1.00	46.52 38.58
1919	CB	ASP	253	119.021	59.715	37.632	1.00	46.23		1992	CD1	TYR	262	106.624	53.671	50.062	1.00	30.09
1921	CG	ASP	253	121.475	59.448	37.340	1.00	56.62	15	1994	CE1	TYR	262	106.295	53.636	48.711	1.00	29.27
1922 1923	OD1 OD2	ASP ASP	253 253	122.222 121.896	59.973 59.029	38.197 36.240	1.00 $1.00$	59.66 59.43		1995 1996	CD2 CE2	TYR TYR	262 262	106.944 106.614	56.039 56.015	49.936 48.581	1.00 1.00	34.60 37.08
1923	C C	ASP	253	118.100	59.701	39.208	1.00	46.95		1990	CZ	TYR	262	106.014	54.809	47.974	1.00	32.64
1925	O	ASP	253	117.404	60.508	38.597	1.00	47.82		1998	OH	TYR	262	105.961	54.776	46.635	1.00	17.43
1926 1927	N CA	LEU LEU	254 254	117.600 116.172	58.743 58.596	39.987 40.253	1.00 $1.00$	50.43 52.57		1999 2000	C O	TYR TYR	262 262	109.398 109.356	53.534 52.522	51.956 52.658	1.00 1.00	52.07 51.95
1928	CB	LEU	254	115.777	57.116	40.236	1.00	51.09	20	2001	N	ALA	263	109.957	53.565	50.748	1.00	48.22
1929	CG	LEU	254	116.036	56.357	38.930	1.00	52.73		2002	CA	ALA	263	110.545	52.386	50.120	1.00	49.01
1930 1931	CD1 CD2	LEU LEU	254 254	115.673 115.244	54.894 56.974	39.102 37.788	1.00 $1.00$	50.16 48.48		2003 2004	CB C	ALA ALA	263 263	110.701 111.879	52.627 51.943	48.623 50.723	1.00 1.00	44.24 52.04
1932	C C	LEU	254	115.867	59.205	41.619	1.00	50.66		2005	Ö	ALA	263	112.458	52.631	51.567	1.00	52.30
1933	O	LEU	254	114.780	59.735	41.848	1.00	48.89	25	2006	N	ARG	264	112.333	50.766	50.297	1.00	53.43
1934 1935	N CA	ASP ASP	255 255	116.838 116.750	59.099 59.641	42.522 43.875	$\frac{1.00}{1.00}$	54.73 59.42	25	2007 2008	CA CB	ARG ARG	264 264	113.592 113.499	50.196 48.670	50.752 50.889	$\frac{1.00}{1.00}$	48.17 40.34
1936	CB	ASP	255	116.930	61.167	43.829	1.00	63.16		2009	CG	ARG	264	112.624	48.166	52.030	1.00	43.53
1937	CG	ASP	255	117.232	61.774	45.193	1.00	70.47		2010	CD	ARG	264	112.450	46.639	51.996	1.00	35.92
1938 1939	OD1 OD2	ASP ASP	255 255	117.674 117.030	61.045 62.997	46.110 45.344	1.00 $1.00$	70.14 79.11		2011 2012	NE CZ	ARG ARG	264 264	111.772 110.964	46.200 45.144	50.774 50.679	$\frac{1.00}{1.00}$	44.75 49.82
1940	C C	ASP	255	115.476	59.260	44.640	1.00	56.10	30	2012	NH1	ARG	264	110.714	44.385	51.738	1.00	51.07
1941	0	ASP	255	114.834	60.106	45.263	1.00	54.97		2014	NH2	ARG	264	110.385	44.857	49.518	1.00	37.04
1942 1943	N CA	PHE PHE	256 256	115.127 113.946	57.977 57.486	44.602 45.308	$\frac{1.00}{1.00}$	55.78 55.28		2015 2016	C O	ARG ARG	264 264	114.676 114.453	50.512 50.444	49.742 48.527	$\frac{1.00}{1.00}$	49.03 47.55
1944	CB	PHE	256	113.556	56.093	44.808	1.00	51.79		2017	N	ASP	265	115.848	50.870	50.252	1.00	50.80
1945	CG	PHE	256	113.024	56.079	43.407	1.00	52.55		2018	CA	ASP	265	116.987	51.164	49.392	1.00	51.29
1946 1947	CD1 CD2	PHE PHE	256 256	113.356 112.186	55.051 57.091	42.537 42.955	$\frac{1.00}{1.00}$	55.14 58.21	35	2019 2020	CB CG	ASP ASP	265 265	117.728 118.690	52.418 52.955	49.877 48.843	1.00 1.00	52.76 54.37
1948	CE1	PHE	256	112.862	55.028	41.236	1.00	57.90		2021	OD1	ASP	265	118.782	52.363	47.747	1.00	50.00
1949	CE2	PHE	256	111.687	57.077	41.656	1.00	60.53		2022	OD2	ASP	265	119.358	53.972	49.120	1.00	60.30
1950 1951	CZ C	PHE PHE	256 256	112.026 114.199	56.042 57.438	40.796 46.812	1.00 $1.00$	57.76 60.70		2023 2024	C O	ASP ASP	265 265	117.903 118.824	49.933 49.846	49.419 50.238	1.00 1.00	45.08 39.53
1952	O	PHE	256	113.292	57.162	47.596	1.00	62.84	40	2025	N	ARG	266	117.614	48.973	48.541	1.00	41.67
1953	N	VAL	257	115.442	57.704	47.203	1.00	64.41	40	2026	CA	ARG	266	118.377	47.732 46.574	48.462	1.00	37.99
1954 1955	CA CB	VAL VAL	257 257	115.834 117.373	57.697 57.799	48.606 48.757	1.00 $1.00$	64.01 62.36		2027 2028	CB CG	ARG ARG	266 266	117.528 116.957	46.771	48.983 50.372	1.00 1.00	38.78 29.99
1956	CG1	VAL	257	117.789	57.482	50.187	1.00	60.29		2029	CD	ARG	266	118.028	46.593	51.418	1.00	37.48
1957 1958	CG2 C	VAL VAL	257 257	118.068 115.179	56.864 58.870	47.774 49.333	1.00 $1.00$	58.27 65.72		2030 2031	NE CZ	ARG ARG	266 266	117.503 117.416	46.781 47.958	52.764 53.376	1.00 1.00	38.27 45.66
1959	Ö	VAL	257	113.179	58.771	50.517	1.00	64.55	45	2031	NH1	ARG	266	117.410	49.066	52.763	1.00	38.28
1960	N	THR	258	114.977	59.971	48.609	1.00	66.40		2033	NH2	ARG	266	116.920	48.027	54.603	1.00	42.90
1961 1962	CA CB	THR THR	258 258	114.364 115.200	61.171 62.437	49.175 48.870	$\frac{1.00}{1.00}$	65.40 65.98		2034 2035	C O	ARG ARG	266 266	118.826 118.671	47.429 46.306	47.034 46.542	$\frac{1.00}{1.00}$	33.86 40.81
1963	OG1	THR	258	115.282	62.633	47.453	1.00	65.25		2036	N	VAL	267	119.392	48.431	46.371	1.00	25.69
1964	CG2	THR	258	116.608	62.302	49.442	1.00	64.28		2037	CA	VAL	267	119.845	48.257	45.000	1.00	20.97
1965 1966	C O	THR THR	258 258	112.919 112.066	61.399 61.769	48.716 49.524	$\frac{1.00}{1.00}$	62.28 62.30	50	2038 2039	CB CG1	VAL VAL	267 267	120.143 121.384	49.611 50.264	44.326 44.933	$\frac{1.00}{1.00}$	21.69 20.36
1967	N	THR	259	112.649	61.182	47.428	1.00	58.00		2040	CG2	VAL	267	120.292	49.420	42.828	1.00	8.30
1968	CA	THR	259	111.303	61.372	46.879	1.00	53.45		2041	С	VAL	267	121.058	47.333	44.913	1.00	27.40
1969 1970	CB OG1	THR THR	259 259	111.300 111.730	61.402 60.136	45.332 44.818	$\frac{1.00}{1.00}$	48.24 47.10		2042 2043	O N	VAL VAL	267 268	121.231 121.889	46.616 47.347	43.926 45.952	$\frac{1.00}{1.00}$	36.12 30.76
1971	CG2	THR	259	112.221	62.496	44.817	1.00	44.78	55	2044	CA	VAL	268	123.080	46.503	46.008	1.00	32.68
1972	С	THR	259	110.320	60.303	47.358	1.00	54.85		2045	CB	VAL	268	123.998	46.904	47.190	1.00	35.53
1973 1974	O N	THR LEU	259 260	109.147 110.802	60.593 59.069	47.593 47.492	$\frac{1.00}{1.00}$	58.40 54.85		2046 2047	CG1 CG2	VAL VAL	268 268	125.220 124.420	46.001 48.355	47.245 47.058	1.00 1.00	32.24 33.69
1975	CA	LEU	260	109.977	57.952	47.958	1.00	57.07		2048	C	VAL	268	122.623	45.058	46.196	1.00	29.06
1976	CB	LEU	260	109.811	56.907 57.346	46.845	1.00	54.53		2049	O	VAL	268	123.119	44.144	45.533	1.00	25.58
1977 1978	CG CD1	LEU LEU	250 250	109.191 109.216	57.346 56.192	45.511 44.523	1.00 $1.00$	54.02 47.30	60	2050 2051	N CA	GLU GLU	269 269	121.662 121.087	44.878 43.573	47.100 47.406	1.00 1.00	24.94 22.59
1979	CD2	LEU	260	107.765	57.838	45.715	1.00	53.62		2052	CB	GLU	269	120.083	43.692	48.558	1.00	16.86
1980	C	LEU	260	110.655	57.326 56.100	49.183	1.00	59.06		2053	CG	GLU	269	120.705	43.939	49.942	1.00	22.84
1981 1982	O N	LEU PRO	260 261	111.135 110.672	56.190 58.057	49.132 50.314	$\frac{1.00}{1.00}$	61.18 59.80		2054 2055	CD OE1	GLU GLU	269 269	121.136 121.417	45.385 45.713	50.200 51.374	1.00 1.00	25.18 24.26
1983	CD	PRO	261	110.004	59.362	50.474	1.00	57.03	/-	2056	OE2	GLU	269	121.194	46.198	49.255	1.00	23.69
1984	CA	PRO	261	111.281	57.634	51.582	1.00	60.02 50.64	65	2057	С	GLU	269	120.404	43.001	46.167	1.00	21.52
1985	CB	PRO	261	111.144	58.883	52.452	1.00	59.64		2058	О	GLU	269	120.423	41.789	45.941	1.00	22.99

TABLE 10-continued

s	tructura			of Tobacco				hase	. 5	s	tructura			of Tobacco l Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
2059	N	CYS	270	119.814	43.885	45.365	1.00	16.93		2132	CD2	TYR	278	121.581	34.952	33.698	1.00	10.48
2060	CA CB	CYS CYS	270 270	119.144 118.305	43.487 44.637	44.133 43.577	$\frac{1.00}{1.00}$	15.61 14.26	10	2133	CE2 CZ	TYR	278 278	120.532 120.371	35.762 37.012	33.290 33.860	$\frac{1.00}{1.00}$	15.32 20.32
2061 2062	SG	CYS	270	116.923	45.097	44.637	1.00	24.62	10	2134 2135	OH	TYR TYR	278	119.341	37.820	33.432	1.00	20.32 16.49
2063	C	CYS	270	120.191	43.065	43.117	1.00	19.83		2136	C	TYR	278	125.236	35.423	36.790	1.00	22.42
2064 2065	O N	CYS TYR	270 271	119.922 121.382	42.239 43.648	42.245 43.220	$\frac{1.00}{1.00}$	25.94 25.82		2137 2138	O N	TYR PHE	278 279	125.024 126.454	36.609 34.883	37.052 36.756	1.00 1.00	24.68 20.65
2066	CA	TYR	271	122.464	43.290	42.315	1.00	22.31		2139	CA	PHE	279	127.665	35.662	36.998	1.00	24.23
2067	CB	TYR	271	123.616	44.296	42.366	1.00	15.29	15	2140	CB	PHE	279	128.474	35.036	38.140	1.00	19.97
2068 2069	CG CD1	TYR TYR	271 271	124.715 124.592	43.914 44.174	41.408 40.043	1.00 $1.00$	11.26 9.23		2141 2142	CG CD1	PHE PHE	279 279	129.063 130.278	33.694 33.600	37.800 37.124	1.00 1.00	27.50 27.56
2070	CE1	TYR	271	125.534	43.705	39.135	1.00	12.75		2142	CD1	PHE	279	128.387	32.523	38.120	1.00	26.31
2071	CD2	TYR	271	125.821	43.184	41.844	1.00	6.39		2144	CE1	PHE	279	130.804	32.363	36.770	1.00	27.48
2072 2073	CE2	TYR	271 271	126.767	42.709	40.946 39.594	1.00 $1.00$	6.94		2145	CE2	PHE	279 279	128.906 130.116	31.283	37.770 37.093	1.00 1.00	27.51
2073	CZ OH	TYR TYR	271	126.615 127.525	42.970 42.467	38.702	1.00	11.07 9.47	20	2146 2147	CZ C	PHE PHE	279	128.564	31.202 35.797	35.773	1.00	26.58 22.60
2075	C	TYR	271	122.973	41.904	42.692	1.00	23.07		2148	O	PHE	279	129.420	36.681	35.727	1.00	29.14
2076 2077	O N	TYR PHE	271 272	123.318 123.037	41.104 41.639	41.820 43.997	1.00 $1.00$	22.92 17.89		2149 2150	N CA	GLU GLU	280 280	128.404 129.217	34.893 34.909	34.811 33.599	1.00 1.00	26.53 25.23
2077	CA	PHE	272	123.484	40.344	44.500	1.00	17.87		2151	CB	GLU	280	128.759	33.832	32.608	1.00	31.74
2079	CB	PHE	272	123.481	40.329	46.033	1.00	20.14	25	2152	CG	GLU	280	129.004	32.392	33.056	1.00	26.55
2080	CG CD1	PHE PHE	272 272	123.722 124.998	38.967 38.409	46.625	$\frac{1.00}{1.00}$	18.24 20.70	25	2153 2154	CD OE1	GLU GLU	280 280	127.873 127.909	31.806 30.581	33.899 34.149	$\frac{1.00}{1.00}$	40.41 41.01
2081 2082	CD1	PHE	272	124.998	38.232	46.631 47.160	1.00	19.50		2155	OE1	GLU	280	126.949	32.549	34.307	1.00	31.27
2083	CE1	PHE	272	125.219	37.139	47.159	1.00	15.25		2156	C	GLU	280	129.195	36.276	32.928	1.00	28.72
2084 2085	CE2 CZ	PHE PHE	272 272	122.881 124.159	36.961 36.414	47.690 47.689	1.00 $1.00$	17.71 $12.72$		2157 2158	O N	GLU PRO	280 281	128.169 130.346	36.958 36.702	32.918 32.382	$\frac{1.00}{1.00}$	21.31 31.20
2085	C	PHE	272	124.139	39.273	43.972	1.00	20.61	30	2159	CD	PRO	281	131.607	35.942	32.368	1.00	29.80
2087	O	PHE	272	122.974	38.200	43.550	1.00	26.50		2160	CA	PRO	281	130.511	37.988	31.697	1.00	31.18
2088 2089	N CA	TRP TRP	273 273	121.248 120.203	39.594 38.695	43.982 43.506	$\frac{1.00}{1.00}$	24.31 20.50		2161 2162	CB CG	PRO PRO	281 281	131.976 132.274	37.949 36.479	31.246 31.137	$\frac{1.00}{1.00}$	36.16 35.87
2099	CB	TRP	273	118.831	39.335	43.724	1.00	22.07		2163	C	PRO	281	129.561	38.213	30.522	1.00	29.70
2091	CG	TRP	273	117.820	38.395	44.280	1.00	22.11		2164	O	PRO	281	129.196	39.352	30.226	1.00	29.95
2092 2093	CD2 CE2	TRP TRP	273 273	117.499 116.513	38.205 37.193	45.661 45.726	1.00 $1.00$	17.16 18.77	35	2165 2166	N CA	GLN GLN	282 282	129.161 128.252	37.126 37.194	29.866 28.722	1.00 1.00	27.63 28.39
2093	CE3	TRP	273	117.949	38.789	46.851	1.00	16.10		2167	CB	GLN	282	128.232	35.832	28.028	1.00	34.10
2095	CD1	TRP	273	117.036	37.522	43.580	1.00	22.90		2168	CG	GLN	282	127.630	34.717	28.912	1.00	45.80
2096 2097	NE1 CZ2	TRP TRP	273 273	116.250 115.969	36.794 36.750	44.442 46.938	1.00 $1.00$	15.33 8.52		2169 2170	CD OE1	GLN GLN	282 282	127.714 128.543	33.351 32.523	28.264 28.647	1.00 1.00	47.83 54.06
2098	CZ3	TRP	273	117.408	38.351	48.057	1.00	19.38	40	2170	NE2	GLN	282	126.858	33.101	27.285	1.00	41.38
2099	CH2	TRP	273	116.428	37.339	48.088	1.00	25.09	40	2172	С	GLN	282	126.851	37.640	29.133	1.00	25.75
2100 2101	C O	TRP TRP	273 273	120.401 120.291	38.389 37.239	42.024 41.596	1.00 $1.00$	19.25 26.32		2173 2174	O N	GLN TYR	282 283	126.061 126.553	38.071 37.521	28.294 30.425	1.00 1.00	34.41 26.54
2102	N	ALA	274	120.705	39.424	41.247	1.00	15.42		2175	CA	TYR	283	125.254	37.910	30.972	1.00	26.30
2103	CA	ALA	274	120.925	39.267	39.815	1.00	17.00		2176	CB	TYR	283	124.765	36.853	31.966	1.00	19.61
2104 2105	CB C	ALA ALA	274 274	120.927 122.240	40.622 38.538	39.138 39.553	$\frac{1.00}{1.00}$	8.15 21.87	45	2177 2178	CG CD1	TYR TYR	283 283	124.537 125.030	35.506 34.339	31.323 31.901	$\frac{1.00}{1.00}$	13.47 13.33
2106	Ö	ALA	274	122.394	37.858	38.535	1.00	24.79		2179	CE1	TYR	283	124.852	33.099	31.286	1.00	15.49
2107	N	LEU	275	123.188	38.694	40.474	1.00	18.76		2180	CD2	TYR	283	123.853	35.402	30.112	1.00	18.67
2108 2109	CA CB	LEU LEU	275 275	124.487 125.505	38.045 38.712	40.354 41.281	$\frac{1.00}{1.00}$	20.76 15.67		2181 2182	CE2 CZ	TYR TYR	283 283	123.669 124.172	34.173 33.026	29.490 30.079	$\frac{1.00}{1.00}$	23.42 14.24
2110	CG	LEU	275	126.937	38.176	41.221	1.00	6.06		2183	OH	TYR	283	124.002	31.817	29.448	1.00	22.41
2111	CD1	LEU	275	127.475	38.257	39.798	1.00	5.14	50	2184	С	TYR	283	125.304	39.287	31.632	1.00	29.24
2112 2113	CD2 C	LEU LEU	275 275	127.812 124.351	38.960 36.560	42.179 40.684	$\frac{1.00}{1.00}$	9.22 19.71		2185 2186	O N	TYR SER	283 284	124.504 126.244	39.599 40.108	32.517 31.170	$\frac{1.00}{1.00}$	27.80 29.09
2114	Ö	LEU	275	125.130	35.731	40.206	1.00	17.59		2187	CA	SER	284	126.438	41.461	31.673	1.00	24.35
2115	N	GLY	276	123.356	36.239	41.507	1.00	22.38		2188	CB	SER	284	127.644	42.103	30.981	1.00	29.60
2116 2117	CA C	GLY GLY	276 276	123.098 122.429	34.860 34.101	41.880 40.747	$\frac{1.00}{1.00}$	15.02 8.27	55	2189 2190	OG C	SER SER	284 284	127.873 125.192	43.418 42.315	31.456 31.451	1.00 1.00	29.67 23.95
2118	Ö	GLY	276	122.574	32.885	40.641	1.00	21.39	33	2191	Ö	SER	284	124.647	42.882	32.396	1.00	16.24
2119	N	VAL	277	121.693	34.825	39.904	1.00	9.55		2192	N	GLN	285	124.743	42.393	30.199	1.00	29.77
2120 2121	CA CB	VAL VAL	277 277	120.992 119.950	34.245 35.238	38.758 38.201	$\frac{1.00}{1.00}$	6.19 4.45		2193 2194	CA CB	GLN GLN	285 285	123.556 123.313	43.173 43.138	29.852 28.339	1.00 1.00	36.37 36.36
2122	CG1	VAL	277	119.236	34.660	36.994	1.00	2.00		2195	CG	GLN	285	122.119	43.974	27.883	1.00	42.33
2123	CG2	VAL	277	118.946	35.576	39.284	1.00	2.00	60	2196	CD OF 1	GLN	285	121.887	42.913	26.382	1.00	47.38
2124 2125	C O	VAL VAL	277 277	122.003 121.872	33.848 32.807	37.686 37.042	1.00 $1.00$	9.32 17.27	-	2197 2198	OE1 NE2	GLN GLN	285 285	122.208 121.321	42.919 44.981	25.727 25.832	1.00 1.00	45.82 47.01
2126	N	TYR	278	122.992	34.711	37.481	1.00	13.83		2199	C	GLN	285	122.328	42.638	30.588	1.00	40.35
2127	CA	TYR	278	124.082	34.466	36.543	1.00	17.57		2200	O	GLN	285	121.503	43.413	31.076	1.00	46.67
2128 2129	CB CG	TYR TYR	278 278	123.644 122.485	34.476 35.368	35.067 34.675	1.00 $1.00$	14.31 21.92		2201 2202	N CA	ALA ALA	286 286	122.242 121.136	41.312 40.637	30.686 31.356	1.00 1.00	38.34 27.14
2130	CD1	TYR	278	122.304	36.630	35.242	1.00	28.06	65	2202	CB	ALA	286	121.150	39.136	31.170	1.00	34.43
2131	CE1	TYR	278	121.249	37.454	34.839	1.00	17.69		2204	С	ALA	286	121.067	40.983	32.837	1.00	24.71

TABLE 10-continued

s	tructura			of Tobacco				hase	5	s	tructura			of Tobacco l Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor	_	Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
2205	О	ALA	286	119.996	41.319	33.346	1.00	32.18		2278	CA	MET	296	111.843	48.816	40.685	1.00	33.20
2206	N CA	ARG	287 287	122.203 122.261	40.891 41.212	33.526 34.951	$\frac{1.00}{1.00}$	17.67 17.77	10	2279	CB	MET MET	296 296	111.659 110.820	48.829 49.978	39.165 38.637	$\frac{1.00}{1.00}$	33.02 34.84
2207 2208	CB	ARG ARG	287	122.201	41.212	35.504	1.00	14.04	10	2280 2281	CG SD	MET	295	111.653	51.571	38.807	1.00	40.22
2209	CG	ARG	287	124.013	39.659	36.008	1.00	21.57		2282	CE	MET	296	110.937	52.162	40.336	1.00	31.58
2210 2211	CD NE	ARG ARG	287 287	125.294 126.451	39.673 40.078	36.828 36.033	$\frac{1.00}{1.00}$	20.38 14.48		2283 2284	C O	MET MET	296 296	110.617 109.831	48.181 48.871	41.343 41.997	1.00 1.00	36.25 35.27
2211	CZ	ARG	287	120.431	39.293	35.162	1.00	22.09		2285	N	ILE	297	110.462	46.867	41.172	1.00	36.41
2213	NH1	ARG	287	126.670	38.049	34.959	1.00	17.96	15	2286	CA	ILE	297	109.327	46.145	41.743	1.00	31.22
2214 2215	NH2 C	ARG ARG	287 287	128.132 121.802	39.749 42.642	34.497 35.207	1.00 $1.00$	25.61 24.30		2287 2288	CB CG2	ILE ILE	297 297	109.240 110.401	44.681 43.850	41.222 41.748	1.00 1.00	30.61 32.13
2216	Ö	ARG	287	120.942	42.877	36.046	1.00	22.53		2289	CG2	ILE	297	107.915	44.039	41.647	1.00	25.21
2217	N	VAL	288	122.358	43.583	34.449	1.00	32.79		2290	CD1	ILE	297	106.681	44.725	41.088	1.00	9.77
2218 2219	CA	VAL	288 288	122.031	45.001	34.586 33.543	1.00 $1.00$	35.93		2291	C	ILE	297 297	109.362	46.151	43.266 43.914	1.00 1.00	28.01
2220	CB CG1	VAL VAL	288	122.800 122.484	45.853 47.329	33.718	1.00	42.80 45.42	20	2292 2293	N	ILE SER	298	108.333 110.544	45.964 46.369	43.834	1.00	33.37 26.49
2221	CG2	VAL	288	124.294	45.622	33.682	1.00	39.26		2294	CA	SER	298	110.682	46.410	45.284	1.00	31.31
2222 2223	C O	VAL VAL	288 288	120.525 119.927	45.246 45.888	34.470 35.339	1.00 $1.00$	30.79 27.62		2295 2296	CB OG	SER SER	298 298	112.152 112.281	46.511 46.533	45.692 47.106	1.00 1.00	36.65 34.52
2224	N	MET	289	119.927	44.698	33.422	1.00	25.59		2290	C	SER	298	109.921	47.616	45.810	1.00	31.13
2225	CA	MET	289	118.473	44.837	33.202	1.00	20.40	25	2298	O	SER	298	109.331	47.567	46.888	1.00	32.44
2226 2227	CB CG	MET MET	289 289	118.055 118.675	44.123 44.684	31.908 30.646	$\frac{1.00}{1.00}$	11.02 17.69	25	2299 2300	N CA	ILE ILE	299 299	109.932 109.239	48.693 49.923	45.029 45.390	$\frac{1.00}{1.00}$	31.80 37.55
2228	SD	MET	289	118.236	43.769	29.151	1.00	29.61		2301	CB	ILE	299	109.239	51.086	44.462	1.00	44.76
2229	CE	MET	289	117.076	44.873	28.424	1.00	23.00		2302	CG2	ILE	299	108.809	52.326	44.753	1.00	45.64
2230 2231	C O	MET MET	289 289	117.692 116.762	44.246 44.861	34.383 34.901	1.00 $1.00$	20.89 25.06		2303 2304	CG1 CD1	ILE ILE	299 299	111.135 111.656	51.390 52.438	44.645 43.707	$\frac{1.00}{1.00}$	46.63
2232	N	LEU	290	118.104	43.063	34.825	1.00	20.72	30	2304	CDI	ILE	299	107.735	49.710	45.315	1.00	51.58 31.48
2233	CA	LEU	290	117.448	42.379	35.935	1.00	15.74		2306	O	ILE	299	107.008	50.072	46.238	1.00	32.42
2234 2235	CB CG	LEU LEU	290 290	118.020 117.497	40.969 40.044	36.078 37.174	$\frac{1.00}{1.00}$	14.98 16.12		2307 2308	N CA	VAL VAL	300 300	107.277 105.855	49.101 48.824	44.224 44.037	$\frac{1.00}{1.00}$	27.09 23.95
2236	CD1	LEU	290	117.497	39.981	37.174	1.00	16.12		2309	CB	VAL	300	105.598	48.080	42.709	1.00	22.51
2237	CD2	LEU	290	118.098	38.659	36.964	1.00	19.76		2310	CG1	VAL	300	104.108	47.876	42.494	1.00	20.37
2238 2239	C O	LEU LEU	290 290	117.530 116.561	43.139 43.172	37.261 38.019	$\frac{1.00}{1.00}$	21.50 23.53	35	2311 2312	CG2 C	VAL VAL	300 300	106.190 105.349	48.867 47.990	41.552 45.211	1.00 1.00	20.21 23.59
2240	N	VAL	291	118.675	43.761	37.534	1.00	24.67		2312	Ö	VAL	300	103.349	48.204	45.714	1.00	30.31
2241	CA	VAL	291	118.858	44.518	38.773	1.00	28.92		2314	N	ASP	301	106.186	47.072	45.674	1.00	24.66
2242 2243	CB CG1	VAL VAL	291 291	120.280 120.439	45.113 45.822	38.893 40.234	1.00 $1.00$	30.77 27.10		2315 2316	CA CB	ASP ASP	301 301	105.837 106.879	46.226 45.121	46.802 46.975	1.00 1.00	31.65 25.48
2244	CG2	VAL	291	121.324	44.033	38.742	1.00	35.91		2317	CG	ASP	301	106.523	44.163	48.087	1.00	24.95
2245	С	VAL	291	117.872	45.679	38.826	1.00	33.69	40	2318	OD1	ASP	301	105.672	43.277	47.869	1.00	37.24
2246 2247	O N	VAL LYS	291 292	117.266 117.722	45.950 46.360	39.867 37.693	1.00 $1.00$	38.64 32.65		2319 2320	OD2 C	ASP ASP	301 301	107.075 105.762	44.309 47.078	49.193 48.065	1.00 1.00	32.62 33.30
2248	CA	LYS	292	116.819	47.500	37.589	1.00	29.16		2321	Ö	ASP	301	103.762	46.930	48.874	1.00	36.41
2249	CB	LYS	292	116.961	48.155	36.213	1.00	28.67		2322	N	ASP	302	106.737	47.971	48.218	1.00	41.65
2250 2251	CG CD	LYS LYS	292 292	118.314 118.440	48.814 49.353	35.986 34.575	$\frac{1.00}{1.00}$	28.14 36.09	45	2323 2324	CA CB	ASP ASP	302 302	106.805 108.124	48.866 49.650	49.369 49.353	$\frac{1.00}{1.00}$	43.94 49.38
2252	CE	LYS	292	119.765	50.059	34.370	1.00	37.49		2325	CG	ASP	302	109.322	48.798	49.744	1.00	55.74
2253	NZ	LYS	292	119.962	50.417	32.940	1.00	42.24		2326	OD1	ASP	302	109.246	48.101	50.780	1.00	59.49
2254 2255	0	LYS LYS	292 292	115.369 114.633	47.102 47.829	37.849 38.514	$\frac{1.00}{1.00}$	25.46 23.99		2327 2328	OD2 C	ASP ASP	302 302	110.344 105.619	48.833 49.831	49.024 49.416	$\frac{1.00}{1.00}$	52.10 43.11
2256	N	THR	293	114.033	34.922	37.365	1.00	28.08		2329	Ö	ASP	302	105.019	50.257	50.493	1.00	44.18
2257	CA	THR	293	113.627	45.401	37.536	1.00	20.11	50	2330	N	THR	303	105.081	50.159	48.243	1.00	38.88
2258 2259	CB OG1	THR THR	293 293	113.385 113.325	44.183 44.619	36.617 35.252	$\frac{1.00}{1.00}$	19.93 19.22		2331 2332	CA CB	THR THR	303 303	103.945 103.745	51.069 51.514	48.123 46.660	$\frac{1.00}{1.00}$	33.20 41.14
2260	CG2	THR	293	112.095	43.472	36.972	1.00	14.44		2333	OG1	THR	303	104.965	52.075	46.158	1.00	39.42
2261	С	THR	293	113.326	45.026	38.987	1.00	22.63		2334	CG2	THR	303	102.643	52.555	46.564	1.00	44.39
2262 2263	O N	THR ILE	293 294	112.286 114.239	45.405 44.295	39.524 39.621	$\frac{1.00}{1.00}$	30.10 23.15		2335 2336	C O	THR THR	303 303	102.652 101.930	50.426 51.016	48.624 49.425	$\frac{1.00}{1.00}$	36.28 44.91
2264	CA	ILE	294	114.058	43.884	41.015	1.00	22.01	55	2337	N	PHE	304	102.367	49.218	48.143	1.00	33.21
2265	CB	ILE	294	115.232	43.007	41.522	1.00	19.34		2338	CA	PHE	304	101.167	48.478	48.532	1.00	26.28
2266 2267	CG2 CG1	ILE ILE	294 294	114.962 115.430	42.546 41.799	42.958 40.604	$\frac{1.00}{1.00}$	19.45 12.44		2339 2340	CB CG	PHE PHE	304 304	101.005 100.431	47.234 47.509	47.653 46.293	1.00 1.00	23.04 14.95
2268	CD1	ILE	294	116.564	40.876	41.017	1.00	23.70		2341	CD1	PHE	304	101.250	47.901	45.239	1.00	16.64
2269	C	ILE	294	113.959	45.113	41.910	1.00	21.14	60	2342	CD2	PHE	304	99.068	47.352	46.059	1.00	16.64
2270 2271	O N	ILE SER	294 295	113.097 114.841	45.193 46.075	42.789 41.664	1.00 $1.00$	23.19 27.23		2343 2344	CE1 CE2	PHE PHE	304 304	100.720 98.527	48.132 47.580	43.968 44.793	1.00 1.00	20.58 13.24
2271	CA	SER	295 295	114.879	47.310	42.435	1.00	36.44		2345	CZ	PHE	304	99.355	47.971	43.746	1.00	15.24
2273	CB	SER	295	116.063	48.167	41.979	1.00	38.02		2346	С	PHE	304	101.183	48.032	49.993	1.00	34.53
2274 2275	OG C	SER SER	295 295	116.508 113.566	49.015 48.077	43.021 42.265	1.00 $1.00$	50.33 33.20		2347 2348	O N	PHE ASP	304 305	100.135 102.379	47.926 47.778	50.632 50.515	1.00 1.00	38.46 40.19
2276	Ö	SER	295	112.984	48.562	43.239	1.00	27.63	65	2349	CA	ASP	305	102.544	47.778	51.881	1.00	46.42
2277	N	MET	296	113.083	48.124	41.026	1.00	32.18		2350	CB	ASP	305	103.774	46.399	51.968	1.00	58.96

TABLE 10-continued

s _	tructura					ristoloche nate Boun		hase	5	s _	tructura			of Tobacco yl Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	occ	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
2351	CG	ASP	305	103.431	44.933	51.845	1.00	70.39		2424	С	GLU	314	93.417	51.357	41.880	1.00	38.64
2352	OD1	ASP	305	102.647	44.572	50.940	1.00	75.74		2425	O	GLU	314	93.315	50.828	40.772	1.00	41.77
2353	OD2	ASP	305	103.957	44.140	52.656	1.00	75.60	10	2426	N	ALA	315	93.742	52.634	42.047	1.00	37.56
2354	C	ASP	305	102.605	48.324	53.001	1.00	47.79		2427	CA	ALA	315	94.012	53.513	40.917	1.00	37.09
2355	O	ASP	305	101.936	48.161	54.022	1.00	46.25		2428	CB	ALA	315	94.024	54.961	41.375	1.00	41.02
2356 2357	N CA	ALA ALA	306 306	103.425 103.574	49.357 50.362	52.831 53.875	$\frac{1.00}{1.00}$	52.07 54.53		2429 2430	C O	ALA ALA	315 315	95.354 95.522	53.146 53.222	40.292 39.074	1.00 1.00	37.51 40.72
2358	CB	ALA	306	104.958	50.240	54.518	1.00	57.28		2431	N	TYR	316	96.301	52.741	41.138	1.00	37.47
2359	C	ALA	306	103.312	51.807	53.462	1.00	53.39	15	2432	CA	TYR	316	97.641	52.351	40.698	1.00	36.60
2360	O	ALA	306	103.971	52.718	53.965	1.00	58.62	13	2433	CB	TYR	316	98.567	52.189	41.908	1.00	42.76
2361	N	TYR	307	102.345	52.030	52.576	1.00	51.30		2434	CG	TYR	316	100.045	52.214	41.576	1.00	47.70
2362	CA	TYR	307	102.035	53.395	52.165	1.00	53.81		2435	CD1	TYR	316	100.701	53.421	41.323	1.00	50.35
2363 2364	CB CG	TYR TYR	307 307	103.107	53.918	51.195 51.392	$\frac{1.00}{1.00}$	48.52 55.69		2436	CE1 CD2	TYR TYR	316 316	102.064	53.456 51.038	41.029 41.526	1.00 1.00	47.09 50.58
2365	CD1	TYR	307	103.396 103.963	55.388 55.847	52.581	1.00	59.86		2437 2438	CE2	TYR	316	100.792 102.158	51.058	41.232	1.00	51.96
2366	CE1	TYR	307	104.206	57.200	52.789	1.00	62.54	20	2439	CZ	TYR	316	102.785	52.276	40.986	1.00	46.31
2367	CD2	TYR	307	103.077	56.324	50.411	1.00	56.20		2440	ОН	TYR	316	104.130	52.308	40.697	1.00	45.69
2368	CE2	TYR	307	103.315	57.683	50.608	1.00	59.89		2441	С	TYR	316	97.582	51.047	39.909	1.00	38.30
2369	CZ	TYR	307	103.878	58.112	51.801	1.00	62.92		2442	O	TYR	316	98.142	50.949	38.812	1.00	29.96
2370	OH	TYR	307	104.112	59.451	52.009	1.00	66.19		2443	N	THR	317	96.890	50.058	40.473	1.00	34.49
2371	С	TYR	307	100.647	53.640 54.078	51.597	1.00	59.33	25	2444	CA	THR	317	96.731	48.752	39.839	1.00	33.58
2372 2373	O N	TYR GLY	307 308	99.752 100.473	53.364	52.320 50.307	$\frac{1.00}{1.00}$	66.09 59.69	20	2445 2446	CB OG1	THR THR	317 317	95.811 96.347	47.831 47.676	40.671 41.990	1.00 1.00	29.35 30.19
2374	CA	GLY	308	99.199	53.580	49.636	1.00	58.39		2447	CG2	THR	317	95.691	46.460	40.020	1.00	28.64
2375	C	GLY	308	97.924	53.122	50.326	1.00	60.86		2448	С	THR	317	96.125	48.922	38.448	1.00	38.84
2376	O	GLY	308	97.925	52.163	51.101	1.00	61.04		2449	O	THR	317	96.624	48.363	37.470	1.00	37.70
2377	N	THR	309	96.833	53.833	50.044	1.00	60.72		2450	N	ASP	318	95.070	49.731	38.369	1.00	39.63
2378	CA	THR	309	95.522	53.524	50.609	1.00	57.07	30	2451	CA	ASP	318	94.385	49.987	37.110	1.00	43.25
2379	CB	THR	309	94.751	54.807	509.89	1.00	55.19		2452	CB	ASP	318	93.115	50.806	37.351 36.094	1.00	53.81
2380 2381	OG1 CG2	THR THR	309 309	94.651 95.461	55.667 55.538	49.847 52.117	$\frac{1.00}{1.00}$	47.62 46.57		2453 2454	CG OD1	ASP ASP	318 318	92.282 91.830	50.972 49.947	35.538	1.00 1.00	64.26 68.93
2382	C	THR	309	94.693	52.722	49.609	1.00	59.08		2455	OD2	ASP	318	92.088	52.126	35.656	1.00	69.91
2383	Ö	THR	309	94.996	52.709	48.415	1.00	56.21		2456	C	ASP	318	95.292	50.706	36.118	1.00	38.89
2384	N	VAL	310	93.631	52.089	50.107	1.00	60.80	35	2457	О	ASP	318	95.280	50.406	34.922	1.00	35.65
2385	CA	VAL	310	92.737	51.264	49.295	1.00	62.84		2458	N	ALA	319	96.081	51.651	36.622	1.00	39.85
2386	CB	VAL	310	91.430	50.932	50.059	1.00	64.23		2459	CA	ALA	319	97.001	52.409	35.783	1.00	39.48
2387 2388	CG1 CG2	VAL VAL	310 310	90.667 91.737	49.821 50.534	49.351 51.498	$\frac{1.00}{1.00}$	66.80 61.87		2460 2461	CB C	ALA ALA	319 319	97.716 98.007	53.462 51.469	36.610 35.123	1.00 1.00	45.21 36.47
2389	C	VAL	310	92.390	51.903	47.947	1.00	63.74		2462	Ö	ALA	319	98.261	51.564	33.920	1.00	28.06
2390	ō	VAL	310	92.469	51.244	46.904	1.00	60.65	40	2463	N	ILE	320	98.547	50.541	35.912	1.00	36.88
2391	N	LYS	311	92.038	53.189	47.972	1.00	65.40	40	2464	CA	ILE	320	99.514	49.560	35.422	1.00	35.98
2392	CA	LYS	311	91.687	53.926	46.755	1.00	65.81		2465	CB	ILE	320	99.994	48.620	36.561	1.00	43.40
2393	CB	LYS	311	91.121	55.301	47.105	1.00	70.45		2466	CG2	ILE	320	100.784	47.443	35.991	1.00	44.99
2394 2395	CG CD	LYS LYS	311 311	89.696 89.175	55.305 56.734	47.621 47.692	1.00 $1.00$	74.65 77.16		2467 2468	CG1 CD1	ILE ILE	320 320	100.834 102.173	49.497 49.891	37.582 37.054	1.00 1.00	45.58 39.26
2396	CE	LYS	311	87.719	56.787	48.120	1.00	75.45		2469	CD1	ILE	320	98.911	48.712	34.307	1.00	31.85
2397	NZ	LYS	311	87.239	58.194	48.137	1.00	78.70	45	2470	ŏ	ILE	320	99.544	48.506	33.271	1.00	32.45
2398	С	LYS	311	92.889	54.116	45.842	1.00	63.97		2471	N	GLN	321	97.680	48.245	34.518	1.00	27.13
2399	O	LYS	311	92.840	53.788	44.658	1.00	64.41		2472	CA	GLN	321	96.980	47.414	33.538	1.00	29.80
2400	N	GLU	312	93.961	54.670	46.403	1.00	60.54		2473	CB	GLN	321	95.592	47.021	34.053	1.00	37.41
2401	CA	GLU	312	95.195	54.931	45.665	1.00	57.43		2474	CG	GLN	321	95.581	46.336	35.422	1.00	37.33
2402 2403	CB CG	GLU GLU	312 312	96.263 95.900	55.516 56.859	46.596 47.194	$\frac{1.00}{1.00}$	59.85 66.27	50	2475 2476	CD OE1	GLN GLN	321 321	96.510 96.690	45.136 44.398	35.508 34.536	$\frac{1.00}{1.00}$	42.33 43.16
2404	CD	GLU	312	97.024	57.438	48.033	1.00	69.45	50	2477	NE2	GLN	321	97.108	44.938	36.679	1.00	35.06
2405	OE1	GLU	312	97.396	56.817	49.051	1.00	71.29		2478	C	GLN	321	96.856	48.101	32.180	1.00	30.46
2406	OE2	GLU	312	97.544	58.514	47.666	1.00	70.23		2479	O	GLN	321	97.066	47.474	31.139	1.00	25.13
2407	С	GLU	312	95.750	53.679	44.989	1.00	53.28		2480	N	ARG	322	96.519	49.390	32.199	1.00	36.61
2408	0	GLU	312	96.133	53.715	43.815	1.00	44.50		2481	CA	ARG	322	96.384	50.171	30.971	1.00	40.97
2409	N CA	LEU	313	95.787 96.290	52.577	45.736	1.00	45.10	55	2482	CB	ARG ARG	322 322	95.779 94.261	51.549 51.612	31.264	1.00	48.13 58.29
2410 2411	CA CB	LEU LEU	313 313	96.290	51.310 50.267	45.222 46.343	$\frac{1.00}{1.00}$	40.42 35.64		2483 2484	CG CD	ARG	322	94.261	50.836	31.176 32.290	1.00 $1.00$	58.29 66.64
2412	CG	LEU	313	97.263	50.614	47.534	1.00	32.14		2485	NE	ARG	322	92.125	50.822	32.134	1.00	76.38
2413	CD1	LEU	313	97.226	49.501	48.569	1.00	28.15		2486	CZ	ARG	322	91.326	51.868	32.344	1.00	74.78
2414	CD2	LEU	313	98.687	50.861	47.061	1.00	26.11		2487	NH1	ARG	322	91.827	53.035	32.727	1.00	73.98
2415	C	LEU	313	95.430	50.800	44.071	1.00	41.37	60	2488	NH2	ARG	322	90.019	51.751	32.157	1.00	73.84
2416	O	LEU	313	95.950	50.275	43.085	1.00	39.82	50	2489	С	ARG	322	97.737	50.347	30.290	1.00	39.65
2417	N	GLU	314	94.116	50.981	44.193	1.00	41.16		2490	O	ARG	322	97.848	50.233	29.067	1.00	46.16
2418 2419	CA CB	GLU GLU	314 314	93.180 91.737	50.553 50.728	43.156 43.636	1.00 $1.00$	41.35 45.36		2491 2492	N CA	TRP TRP	323 323	98.757 100.118	50.635 50.828	31.094 30.607	1.00 1.00	37.39 35.58
2420	CG	GLU	314	90.674	50.726	42.612	1.00	51.16		2493	CB	TRP	323	100.118	49.535	29.990	1.00	36.83
2421	CD	GLU	314	90.717	48.848	42.251	1.00	56.61		2494	CG	TRP	323	102.169	49.447	29.979	1.00	41.47
2422	OE1	GLU	314	90.469	48.007	43.144	1.00	55.46	65	2495	CD2	TRP	323	103.017	49.048	31.063	1.00	38.37
2423	OE2	GLU	314	90.988	48.530	41.071	1.00	49.16		2496	CE2	TRP	323	104.351	49.089	30.594	1.00	38.29

TABLE 10-continued

s	tructura			of Tobacco				hase	5	s	tructura			of Tobacco				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	occ	B- factor	-	Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
2497	CE3	TRP	323	102.779	48.656	32.390	1.00	36.27		2570	CD2	LEU	331	97.789	55.739	39.806	1.00	72.35
2498	CD1	TRP	323 323	103.003	49.714	28.922	1.00	40.25	10	2571	С	LEU	331 331	98.081	60.056	41.057	1.00	59.96
2499 2500	NE1 CZ2	TRP TRP	323	104.312 105.439	49.500 48.753	29.286 31.402	$\frac{1.00}{1.00}$	39.37 30.80	10	2572 2573	O N	LEU PRO	332	98.432 98.145	61.069 59.962	40.456 42.397	1.00 1.00	60.57 56.66
2501	CZ3	TRP	323	103.863	48.322	33.192	1.00	34.06		2574	CD	PRO	332	97.661	58.870	43.257	1.00	52.18
2502	CH2	TRP	323	105.178	48.374	32.691	1.00	37.92		2575	CA	PRO	332	98.666	61.070	43.204	1.00	60.33
2503 2504	C O	TRP TRP	323 323	100.182 100.522	51.976 51.788	29.606 28.437	$\frac{1.00}{1.00}$	36.88 30.80		2576 2577	CB CG	PRO PRO	332 332	98.458 98.462	60.578 59.090	44.640 44.507	1.00 1.00	54.14 53.96
2505	N	ASP	324	99.781	53.157	30.066	1.00	48.71	15	2578	C	PRO	332	100.144	61.294	42.871	1.00	65.03
2506	CA	ASP	324	99.797	54.366	29.249	1.00	54.17	15	2579	O	PRO	332	100.817	60.386	42.381	1.00	68.68
2507 2508	CB CG	ASP ASP	324 324	98.462 98.585	54.586 55.547	28.537 27.366	1.00 $1.00$	56.16 58.43		2580 2581	N CA	ASP ASP	333 333	100.637 102.021	62.500 62.876	43.136 42.839	1.00 1.00	70.86 72.50
2509	OD1	ASP	324	98.956	56.724	27.575	1.00	59.06		2582	CB	ASP	333	102.021	64.220	43.489	1.00	76.30
2510	OD2	ASP	324	98.328	55.120	26.222	1.00	57.24		2583	CG	ASP	333	101.737	65.396	42.760	1.00	73.02
2511	С	ASP	324	100.076	55.543	30.165	1.00	57.09	20	2584	OD1	ASP	333	101.290	66.343	43.438	1.00	75.36
2512 2513	O N	ASP ILE	324 325	99.468 100.977	55.671 56.414	31.230 29.729	1.00 $1.00$	54.89 60.16		2585 2586	OD2 C	ASP ASP	333 333	101.700 103.146	65.378 61.873	41.510 43.105	1.00 1.00	70.83 69.45
2514	CA	ILE	325	101.377	57.584	30.495	1.00	64.12		2587	Ö	ASP	333	104.019	61.694	42.454	1.00	64.53
2515	CB	ILE	325	102.559	58.286	29.788	1.00	67.78		2588	N	TYR	334	103.139	61.226	44.269	1.00	65.65
2516 2517	CG2 CG1	ILE ILE	325 325	102.072 103.325	59.013 59.185	28.542 30.768	$\frac{1.00}{1.00}$	68.84 71.40		2589 2590	CA CB	TYR TYR	334 334	104.195 104.180	60.267 59.900	44.590 46.080	1.00 1.00	64.14 67.04
2518	CD1	ILE	325	104.716	59.579	30.273	1.00	78.93	25	2591	CG	TYR	334	103.162	58.858	46.484	1.00	71.84
2519	C	ILE	325	100.219	58.560	30.760	1.00	62.87		2592	CD1	TYR	334	101.827	59.199	46.688	1.00	74.21
2520 2521	O N	ILE ASN	325 326	100.248 99.181	59.325 58.489	31.723 29.931	$\frac{1.00}{1.00}$	55.53 63.16		2593 2594	CE1 CD2	TYE TYR	334 334	100.895 103.542	58.243 57.531	47.086 46.685	1.00 1.00	74.47 72.86
2522	CA	ASN	326	98.008	59.347	30.075	1.00	60.43		2595	CE2	TYR	334	102.620	56.570	47.081	1.00	72.07
2523	CB	ASN	326	97.060	59.157	28.891	1.00	58.85		2596	CZ	TYR	334	101.299	56.932	47.281	1.00	72.74
2524	CG	ASN	326	97.208	60.240	27.858	1.00	58.26	30	2597	OH	TYR	334	100.386	55.982	47.675	1.00	69.90
2525 2526	OD1 ND2	ASN ASN	326 326	97.005 97.564	61.420 59.854	28.150 26.640	1.00 $1.00$	61.96 59.50		2598 2599	C O	TYR TYR	334 334	104.143 105.181	59.015 58.466	43.714 43.341	$\frac{1.00}{1.00}$	59.49 58.89
2527	C	ASN	326	97.247	59.093	31.370	1.00	61.39		2600	N	MET	335	102.933	58.575	43.379	1.00	51.53
2528	0	ASN	326	96.561	59.982	31.875	1.00	62.82		2601	CA	MET	335	102.762	57.401	42.533	1.00	48.12
2529 2530	N CA	GLU GLU	327 327	97.378 96.691	57.881 57.498	31.904 33.136	$\frac{1.00}{1.00}$	59.66 62.98	2.5	2602 2603	CB CG	MET MET	335 335	101.340 100.979	56.854 56.325	42.637 44.006	$\frac{1.00}{1.00}$	45.40 34.82
2531	CB	GLU	327	96.563	55.973	33.216	1.00	64.39	35	2604	SD	MET	335	99.387	55.502	44.005	1.00	37.30
2532	CG	GLU	327	96.087	55.299	31.933	1.00	69.30		2605	CE	MET	335	99.776	53.994	44.867	1.00	41.41
2533 2534	CD OE1	GLU GLU	327 327	94.708 93.784	55.750 55.783	31.491 32.335	1.00 $1.00$	70.48 72.78		2606 2607	C	MET MET	335 335	103.082 103.354	57.727 56.826	41.081 40.287	1.00 1.00	48.13 55.57
2535	OE2	GLU	327	94.548	56.067	30.291	1.00	63.31		2608	N	LYS	336	103.032	59.013	40.738	1.00	48.65
2536	C	GLU	327	97.414	58.011	34.380	1.00	62.97	40	2609	CA	LYS	336	103.332	59.465	39.380	1.00	50.84
2537 2538	O N	GLU ILE	327 328	96.972 98.510	57.771 58.734	35.505 34.169	1.00 $1.00$	62.71 64.24		2610 2611	CB CG	LYS LYS	336 336	103.004 101.524	60.953 61.301	39.213 39.255	1.00 1.00	55.94 63.40
2539	CA	ILE	328	99.316	59.270	35.264	1.00	67.30		2612	CD	LYS	336	101.324	62.758	38.857	1.00	63.31
2540	CB	ILE	328	100.636	59.886	34.729	1.00	69.70		2613	CE	LYS	336	99.820	63.092	38.764	1.00	60.22
2541 2542	CG2 CG1	ILE ILE	328 328	100.372	61.245 60.013	34.069	1.00 $1.00$	69.85		2614	NZ C	LYS	336	99.580 104.810	64.473 59.237	38.271 39.080	1.00 1.00	62.57
2543	CD1	ILE	328	101.657 103.047	60.424	35.863 35.409	1.00	72.49 71.67	45	2615 2616	Ö	LYS LYS	336 336	104.810	58.938	37.943	1.00	51.33 47.74
2544	C	ILE	328	98.577	60.298	36.122	1.00	66.37		2617	N	ILE	337	105.638	59.382	40.114	1.00	45.10
2545	O	ILE	328	98.763	60.349	37.340	1.00	61.63		2618	CA	ILE	337	107.079	59.195	39.996	1.00	46.36
2546 2547	N CA	ASP ASP	329 329	97.711	61.082 62.128	35.485	1.00 $1.00$	70.35		2619 2620	CG2	ILE	337	107.805 109.309	59.607 59.651	41.297	1.00 $1.00$	48.91 50.47
2548	CB	ASP	329	96.212	62.987	35.134	1.00	73.34		2621	CG1	ILE	337	107.330	60.986	41.759	1.00	50.88
2549	CG	ASP	329	97.154	63.620	34.123	1.00	75.99	50	2622	CD1	ILE	337	107.888	61.407	43.105	1.00	47.89
2550 2551	OD1 OD2	ASP ASP	329 329	97.861 97.198	64.584 63.140	34.486 32.970	$\frac{1.00}{1.00}$	75.75 74.82		2623 2624	C	ILE ILE	337 337	107.380 108.140	57.725 57.402	39.712 38.795	$\frac{1.00}{1.00}$	47.32 52.27
2552	C	ASP	329	95.978	61.611	37.219	1.00	73.26		2625	N	SER	338	106.755	56.844	40.491	1.00	42.57
2553	0	ASP	329	95.637	62.332	38.159	1.00	73.84		2626	CA	SER	338	106.928	55.401	40.351	1.00	32.89
2554 2555	N CA	ARG ARG	330 330	95.539 94.616	60.366 59.756	37.065 38.019	$\frac{1.00}{1.00}$	70.87 70.53		2627 2628	CB OG	SER SER	338 338	106.120 106.339	54.663 55.198	41.424 42.718	1.00 1.00	29.02 33.47
2556	CB	ARG	330	93.932	58.535	37.393	1.00	71.49	55	2629	C	SER	338	106.465	54.933	38.975	1.00	31.60
2557	CG	ARG	330	93.145	58.845	36.129	1.00	78.19		2630	0	SER	338	107.214	54.287	38.243	1.00	27.59
2558 2559	CD NE	ARG ARG	330 330	92.435 91.756	57.612 57.889	35.591 34.326	$\frac{1.00}{1.00}$	85.11 94.74		2631 2632	N CA	TYR TYR	339 339	105.239 104.622	55.311 54.932	38.621 37.353	1.00 1.00	33.89 39.75
2560	CZ	ARG	330	90.865	57.082	33.754	1.00	98.54		2633	CB	TYR	339	104.022	55.508	37.265	1.00	42.11
2561	NH1	ARG	330	90.532	55.934	34.331	1.00	100.00	60	2634	CG	TYR	339	102.367	54.908	36.157	1.00	46.50
2562	NH2	ARG	330	90.309	57.424	32.599	1.00	94.40	00	2635	CD1	TYR	339	101.682	53.709	36.348	1.00	48.25
2563 2564	C O	ARG ARG	330 330	95.358 94.749	59.345 59.145	39.291 40.434	$\frac{1.00}{1.00}$	67.20 61.95		2636 2637	CE1 CD2	TYR TYR	339 339	100.924 102.270	53.144 55.530	35.327 34.915	1.00 1.00	52.79 42.98
2565	N	LEU	331	96.681	59.252	39.183	1.00	66.11		2638	CE2	TYR	339	101.515	54.976	33.890	1.00	51.56
2566	CA	LEU	331	97.539	58.857	40.295	1.00	65.38		2639	CZ	TYR	339	100.845	53.784	34.100	1.00	54.29
2567 2568	CB CG	LEU LEU	331 331	98.727 98.430	58.047 56.802	39.768 38.933	1.00 $1.00$	70.15 71.98	65	2640 2641	OH C	TYR TYR	339 339	100.100 105.414	53.236 55.309	33.080 36.101	1.00 1.00	56.73 43.13
2569	CD1	LEU	331	99.710	56.289	38.300	1.00	67.56		2642	Ö	TYR	339	105.531	54.502	35.174	1.00	41.07

TABLE 10-continued

S	tructura			of Tobacco				hase	5	S	tructura			of Tobacco l Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor	_	Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
2643	N	LYS	340	105.941	56.531	36.064	1.00	49.11		2716	C	ASP	348	115.308	50.325	27.394	1.00	44.57
2644	CA CB	LYS LYS	340 340	106.706 106.894	56.989 58.508	34.903 34.934	1.00	49.74	10	2717	O	ASP TYR	348 349	116.186 114.585	50.116 49.355	26.555 27.959	$\frac{1.00}{1.00}$	45.03 43.55
2645 2646	СБ	LYS	340	100.894	59.059	33.674	$\frac{1.00}{1.00}$	58.08 64.41	10	2718 2719	N CA	TYR	349	114.773	47.943	27.627	1.00	42.30
2647	CD	LYS	340	107.642	60.573	33.694	1.00	69.77		2720	CB	TYR	349	113.813	47.054	28.429	1.00	41.29
2648 2649	CE NZ	LYS LYS	340 340	108.246 108.256	61.097 62.584	32.403 32.365	$\frac{1.00}{1.00}$	73.44 82.14		2721 2722	CG CD1	TYR TYR	349 349	114.128 113.181	46.916 46.393	29.906 30.785	1.00 1.00	37.24 30.06
2650	C	LYS	340	108.230	56.297	34.800	1.00	44.71		2723	CE1	TYR	349	113.161	46.247	32.142	1.00	36.46
2651	O	LYS	340	108.506	55.938	33.703	1.00	34.93	15	2724	CD2	TYR	349	115.368	47.293	30.424	1.00	41.76
2652 2653	N CA	ALA ALA	341 341	108.712 110.013	56.114 55.456	35.948 36.003	$\frac{1.00}{1.00}$	36.45 36.97		2725 2726	CE2 CZ	TYR TYR	349 349	115.656 114.694	47.151 46.627	31.783 32.633	1.00 1.00	40.50 36.99
2654	CB	ALA	341	110.513	55.415	37.439	1.00	35.42		2727	OH	TYR	349	114.094	46.491	33.975	1.00	36.59
2655	C	ALA	341	109.897	54.041	35.444	1.00	35.71		2728	C	TYR	349	114.520	47.741	26.139	1.00	42.51
2656 2657	O	ALA	341 342	110.791	53.561	34.746 35.734	1.00 $1.00$	34.74		2729	O	TYR	349 350	115.308 113.411	47.094	25.446 25.656	1.00 1.00	42.03 44.24
2658	N CA	ILE ILE	342	108.766 108.487	53.399 52.041	35.283	1.00	29.99 21.08	20	2730 2731	N CA	GLU GLU	350	113.411	48.296 48.199	24.244	1.00	46.89
2659	CB	ILE	342	107.231	51.472	35.982	1.00	16.81		2732	CB	GLU	350	111.734	48.929	23.969	1.00	49.81
2660	CG2 CG1	ILE ILE	342 342	106.786 107.523	50.171 51.275	35.309 37.476	$\frac{1.00}{1.00}$	10.96 10.78		2733 2734	CG CD	GLU GLU	350 350	110.509 109.214	48.270 49.033	24.589 24.347	1.00 1.00	54.69 58.54
2661 2662	CD1	ILE	342	107.323	50.884	38.324	1.00	2.00		2735	OE1	GLU	350	109.214	48.491	24.695	1.00	62.24
2663	С	ILE	342	108.336	51.939	33.771	1.00	28.67	25	2736	OE2	GLU	350	109.253	50.168	23.822	1.00	64.71
2664 2665	O N	ILE LEU	342 343	108.949 107.530	51.071 52.821	33.150 33.180	$\frac{1.00}{1.00}$	32.50 33.81	25	2737 2738	C	GLU GLU	350 350	114.162	48.811 48.294	23.397 22.334	1.00 1.00	48.29 45.94
2666	CA	LEU	343	107.320	52.809	31.732	1.00	37.97		2739	N	LYS	351	114.491 114.763	49.884	23.909	1.00	53.89
2667	CB	LEU	343	106.208	53.774	31.317	1.00	41.50		2740	CA	LYS	351	115.841	50.591	23.222	1.00	58.14
2668 2669	CG CD1	LEU LEU	343 343	104.822 103.831	53.594 54.537	31.932 31.267	$\frac{1.00}{1.00}$	46.77		2741 2742	CB CG	LYS LYS	351 351	116.053 116.916	51.971 52.921	23.855 23.031	$\frac{1.00}{1.00}$	63.43 71.72
2670	CD1	LEU	343	103.831	52.165	31.759	1.00	48.11 43.32	30	2742	CD	LYS	351	116.247	53.286	23.031	1.00	71.72 77.97
2671	C	LEU	343	108.596	53.177	30.995	1.00	41.41		2744	CE	LYS	351	117.122	54.218	20.885	1.00	83.68
2672 2673	O N	LEU ASP	343 344	108.880 109.348	62.626 54.126	29.932 31.552	$\frac{1.00}{1.00}$	43.16 45.08		2745 2746	NZ C	LYS LYS	351 351	116.483 117.155	54.588 49.795	19.591 23.215	$\frac{1.00}{1.00}$	83.73 57.54
2674	CA	ASP	344	110.601	54.563	30.942	1.00	51.08		2747	Ö	LYS	351	117.133	49.784	22.209	1.00	56.27
2675	CB	ASP	344	111.144	55.820	31.628	1.00	57.96		2748	N	GLU	352	117.465	49.142	24.336	1.00	56.46
2676 2677	CG OD1	ASP ASP	344 344	110.754 110.680	57.098 57.090	30.903 29.654	$\frac{1.00}{1.00}$	64.16 69.36	35	2749 2750	CA CB	GLU GLU	352 352	118.684 118.847	48.334 47.801	24.458 25.390	1.00 1.00	52.60 50.25
2678	OD1	ASP	344	110.526	58.117	31.588	1.00	65.04		2751	CG	GLU	352	119.239	48.828	26.943	1.00	58.10
2679	C	ASP	344	111.643	53.461	30.980	1.00	50.69		2752	CD	GLU	352	119.464	48.194	28.311	1.00	59.00
2680 2681	O N	ASP LEU	344 345	112.415 111.661	53.301 52.709	30.034 32.078	1.00 $1.00$	53.80 47.98		2753 2754	OE1 OE2	GLU GLU	352 352	118.655 120.447	48.447 47.435	29.232 28.468	1.00 1.00	57.55 53.94
2682	CA	LEU	345	112.594	51.599	32.242	1.00	43.23		2755	C	GLU	352	118.645	47.140	23.508	1.00	49.57
2683	CB	LEU	345	112.384	50.925	33.599	1.00	45.16	40	2756	O	GLU	352	119.671	46.735	22.957	1.00	45.55
2684 2685	CG CD1	LEU LEU	345 345	113.317 114.752	49.773 50.267	33.977 34.070	1.00 $1.00$	44.62 45.63		2757 2758	N CA	LEU LEU	353 353	117.448 117.239	46.587 45.432	23.327 22.463	1.00 1.00	44.88 44.80
2686	CD2	LEU	345	112.875	49.186	35.307	1.00	42.33		2759	CB	LEU	353	116.116	44.561	23.034	1.00	35.61
2687	С	LEU	345	112.364	50.585	31.129	1.00	42.19		2760	CG	LEU	353	116.304	44.125	24.489	1.00	30.68
2688 2689	O N	LEU TYR	345 346	113.315 111.094	50.068 50.311	30.541 30.844	$\frac{1.00}{1.00}$	47.47 38.09	45	2761 2762	CD1 CD2	LEU LEU	353 353	115.030 117.468	43.507 43.156	25.030 24.597	$\frac{1.00}{1.00}$	31.93 32.93
2690	CA	TYR	346	111.094	49.372	29.793	1.00	36.61		2763	CD2	LEU	353	116.937	45.806	21.011	1.00	48.25
2691	CB	TYR	346	109.298	48.878	29.983	1.00	32.28		2764	O	LEU	353	116.878	44.933	20.140	1.00	48.95
2692 2693	CG CD1	TYR TYR	346 346	109.211 108.903	47.802 48.110	31.038 32.361	$\frac{1.00}{1.00}$	30.73 22.11		2765 2766	N CA	SER SER	354 354	116.756 116.468	47.101 47.595	20.751 19.403	$\frac{1.00}{1.00}$	54.12 58.83
2694	CE1	TYR	346	108.895	47.122	33.346	1.00	24.69		2767	CB	SER	354	116.356	49.122	19.395	1.00	64.47
2695	CD2	TYR	346	109.503	46.477	30.722	1.00	35.15	50	2768	OG	SER	354	115.196	49.571	20.072	1.00	73.04
2696 2697	CE2 CZ	TYR TYR	346 346	109.499 109.198	45.484 45.809	31.694 33.000	$\frac{1.00}{1.00}$	26.34 27.54		2769 2770	С О	SER SER	354 354	117.534 117.226	47.171 46.900	18.400 17.237	$\frac{1.00}{1.00}$	58.85 60.25
2698	OH	TYR	346	109.136	44.812	33.948	1.00	22.51		2771	N	SER	355	118.784	47.119	18.857	1.00	59.55
2699	C	TYR	346	110.954	49.953	28.403	1.00	39.45		2772	CA	SER	355	119.918	46.731	18.022	1.00	60.93
2700 2701	O N	TYR LYS	346 347	111.086 110.995	49.213 51.281	27.429 28.320	$\frac{1.00}{1.00}$	37.79 44.42		2773 2774	CB OG	SER SER	355 355	121.219 122.333	46.840 46.392	18.823 18.071	1.00 1.00	58.82 61.05
2701	CA	LYS	347	111.256	51.261	27.056	1.00	45.72	55	2775	C	SER	355	119.772	45.316	17.455	1.00	67.50
2703	CB	LYS	347	110.797	53.418	27.105	1.00	49.09		2776	O	SER	355	119.753	45.125	16.239	1.00	72.60
2704 2705	CG CD	LYS LYS	347 347	109.313 108.959	53.604 53.084	26.824 25.433	$\frac{1.00}{1.00}$	54.73 58.88		2777 2778	N CA	ALA ALA	356 356	119.640 119.501	44.338 42.943	18.345 17.946	1.00 1.00	68.57 67.55
2705	CE	LYS	347	108.939	53.195	25.149	1.00	58.88 58.70		2779	CB	ALA	356	119.501	42.943	17.946	1.00	63.74
2707	NZ	LYS	347	107.129	52.632	23.816	1.00	45.34	60	2780	C	ALA	356	118.163	42.642	17.278	1.00	69.18
2708 2709	C	LYS	347 347	112.756	51.787	26.810	1.00	44.56	00	2781	O N	ALA	356 357	118.071	41.754	16.434	1.00	70.23
2709	O N	LYS ASP	347 348	113.201 113.524	51.803 51.865	25.666 27.901	$\frac{1.00}{1.00}$	44.20 45.25		2782 2783	N CA	GLY GLY	357 357	117.131 115.811	43.385 43.152	17.661 17.102	1.00 1.00	69.13 62.21
2711	CA	ASP	348	114.977	51.748	27.829	1.00	43.43		2784	С	GLY	357	115.027	42.258	18.039	1.00	59.93
2712	CB	ASP	348	115.630	52.041	29.188	1.00	41.08		2785	O	GLY	357	114.203	41.443	17.604	1.00	60.71
2713 2714	CG OD1	ASP ASP	348 348	115.545 115.741	53.509 54.388	29.584 28.716	1.00 $1.00$	45.56 50.29	65	2786 2787	N CA	ARG ARG	358 358	115.322 114.669	42.391 41.616	19.332 20.385	1.00 1.00	54.34 52.71
2715	OD2	ASP	348		53.787	30.775	1.00	46.84		2788	CB	ARG	358	115.713	40.882	21.231	1.00	45.26

TABLE 10-continued

s	tructura			of Tobacco				hase	5	s	tructura			of Tobacco				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor	_	Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
2789	CG	ARG	358	116.561	39.896	20.442	1.00	47.78		2862	OE1	GLU	367	99.829	41.633	29.321	1.00	48.68
2790 2791	CD NE	ARG ARG	358 358	117.644 117.083	39.275 38.456	21.309 22.383	$\frac{1.00}{1.00}$	47.12 44.42	10	2863 2864	OE2 C	GLU GLU	367 367	101.083 102.539	42.670 40.448	27.841 32.585	$\frac{1.00}{1.00}$	43.16 20.02
2791	CZ	ARG	358	117.206	38.724	23.681	1.00	39.46	10	2865	O	GLU	367	102.559	40.448	33.314	1.00	20.02
2793	NH1	ARG	358	117.871	39.797	24.083	1.00	37.90		2866	N	ARG	368	103.766	40.057	32.923	1.00	21.53
2794 2795	NH2 C	ARG ARG	358 358	116.684 113.817	37.905 42.522	24.583 21.282	$\frac{1.00}{1.00}$	43.78 55.62		2867 2868	CA CB	ARG ARG	368 368	104.055 105.406	39.422 38.709	34.205 34.159	1.00 1.00	14.50 17.59
2796	Ö	ARG	358	113.676	42.268	22.479	1.00	60.74		2869	CG	ARG	368	105.427	37.442	33.319	1.00	13.40
2797	N	SER	359	113.286	43.596	20.699	1.00	54.17	15	2870	CD	ARG	368	104.602	36.338	33.960	1.00	17.58
2798 2799	CA CB	SER SER	359 359	112.440 112.373	44.548 45.887	21.419 20.671	1.00 $1.00$	49.75 46.10		2871 2872	NE CZ	ARG ARG	368 368	104.843 104.380	35.044 33.884	33.325 33.784	1.00 1.00	25.37 29.88
2800	OG	SER	350	113.659	46.424	20.441	1.00	39.24		2873	NH1	ARG	368	103.641	33.847	34.887	1.00	15.72
2801	C	SER	359	111.030	43.979	21.584	1.00	51.35		2874	NH2	ARG	368	104.669	32.747	33.146	1.00	25.60
2802 2803	O N	SER HIS	359 360	110.321 110.642	44.294 43.145	22.549 20.619	1.00 $1.00$	51.03 48.18		2875 2876	C O	ARG ARG	368 368	104.058 103.674	40.473 40.193	35.306 36.444	1.00 1.00	22.28 25.28
2804	CA	HIS	360	109.339	42.484	20.566	1.00	49.55	20	2877	N	MET	369	103.074	41.686	34.965	1.00	23.23
2805	CB	HIS	360	109.165	41.769	19.214	1.00	55.45		2878	CA	MET	369	104.513	42.774	35.933	1.00	22.69
2806 2807	CG CD2	HIS HIS	360 360	110.191 111.485	40.706 40.791	18.955 18.565	1.00 $1.00$	57.94 58.70		2879 2880	CB CG	MET MET	369 369	105.234 105.216	44.001 45.178	35.371 36.332	1.00 1.00	21.94 33.04
2808	ND1	HIS	360	109.933	39.363	19.134	1.00	62.62		2881	SD	MET	369	105.216	46.580	35.855	1.00	30.05
2809	CE1	HIS	360	111.028	38.668	18.875	1.00	63.83	25	2882	CE	MET	369	106.788	47.105	37.492	1.00	22.46
2810 2811	NE2 C	HIS HIS	360 360	111.985 109.105	39.511 41.483	18.527 21.705	$\frac{1.00}{1.00}$	63.77 51.67	25	2883 2884	C O	MET MET	369 369	103.088 102.794	43.138 43.316	36.439 37.513	1.00 1.00	21.30 25.30
2812	Ö	HIS	360	108.023	40.886	21.795	1.00	56.00		2885	N	LYS	370	102.207	43.230	35.332	1.00	22.42
2813	N	ILE	361	110.115	41.294	22.552	1.00	44.35		2886	CA	LYS	370	100.798	43.555	35.562	1.00	21.17
2814 2815	CA CB	ILE ILE	361 361	110.005 111.217	40.352 39.396	23.659 23.719	$\frac{1.00}{1.00}$	40.02 40.34		2887 2888	CB CG	LYS LYS	370 370	100.033 100.498	43.596 44.679	34.237 33.272	$\frac{1.00}{1.00}$	18.37 19.38
2816	CG2	ILE	351	111.350	38.631	22.412	1.00	38.81	30	2889	CD	LYS	370	99.724	44.628	31.959	1.00	22.90
2817	CG1	ILE	361	112.490	40.174	24.052	1.00	45.99		2890	CE	LYS	370	100.144	45.754	31.026	1.00	25.30
2818 2819	CD1 C	ILE ILE	361 361	113.742 109.837	39.324 41.012	24.096 25.022	$\frac{1.00}{1.00}$	38.31 38.58		2891 2892	NZ C	LYS LYS	370 370	99.370 100.184	45.760 42.503	29.753 36.480	$\frac{1.00}{1.00}$	26.14 22.60
2820	Ö	ILE	361	109.620	40.323	25.018	1.00	46.53		2893	Ö	LYS	370	99.433	42.830	37.404	1.00	22.82
2821	N	VAL	362	109.920	42.339	25.068	1.00	34.37		2894	N	GLU	371	100.540	41.243	36.233	1.00	20.87
2822 2823	CA CB	VAL VAL	362 362	109.784 110.133	43.073 44.567	26.323 26.131	$\frac{1.00}{1.00}$	36.05 44.38	35	2895 2896	CA CB	GLU GLU	371 371	100.060 100.633	40.117 38.805	37.027 36.473	1.00 1.00	17.91 17.04
2824	CG1	VAL	362	110.157	45.290	27.474	1.00	28.09		2897	CG	GLU	371	100.291	37.558	37.281	1.00	14.89
2825	CG2	VAL	362	111.470	44.705	25.420	1.00	50.90		2898	CD	GLU	371	100.951	36.307	36.737	1.00	33.96
2826 2827	C O	VAL VAL	362 362	108.372 108.187	42.959 43.012	26.899 28.113	1.00 $1.00$	35.07 27.77		2899 2900	OE1 OE2	GLU GLU	371 371	100.246 102.162	35.281 36.342	36.624 36.433	1.00 1.00	49.48 39.35
2828	N	CYS	363	107.383	42.770	26.025	1.00	37.13	40	2901	С	GLU	371	100.459	40.291	38.491	1.00	14.70
2829	CA CB	CYS	363	105.980	42.653 42.483	26.437 25.215	1.00 $1.00$	34.16 29.73	70	2902	O	GLU	371 372	99.629 101.727	40.146 40.624	39.389 38.718	1.00	15.00
2830 2831	SG	CYS CYS	363 363	105.066 105.447	42.483	24.179	1.00	41.96		2903 2904	N CA	VAL VAL	372	101.727	40.824	40.055	1.00 1.00	16.52 17.45
2832	C	CYS	363	105.730	41.520	27.434	1.00	32.61		2905	CB	VAL	372	103.747	41.232	40.043	1.00	17.58
2833 2834	O N	CYS HIS	363 364	104.887 106.481	41.646 40.429	28.325 27.292	1.00 $1.00$	25.38		2906 2907	CG1 CG2	VAL VAL	372 372	104.258 104.575	41.453 40.156	41.450 39.365	1.00 1.00	3.56 18.80
2835	CA	HIS	364	106.356	39.267	28.168	1.00	28.31 20.38	45	2907	C	VAL	372	104.373	41.919	40.782	1.00	21.27
2836	CB	HIS	364	107.304	38.159	27.713	1.00	19.91		2909	O	VAL	372	101.101	41.757	41.952	1.00	23.46
2837	CD2	HIS	364	107.064	37.696	26.309	1.00	25.64		2910	N CA	VAL	373	101.155	43.000	50.063	1.00	26.12
2838	ND1	HIS	364 364	107.777 105.976	37.887	25.173 25.954	1.00 $1.00$	29.90 34.65		2911 2912	CA CB	VAL VAL	373 373	100.407 100.425	44.123 45.356	40.629 39.694	1.00 $1.00$	29.37 33.84
2840	CE1	HIS	364	106.028	36.667	24.659	1.00	33.79		2913	CG1	VAL	373	99.736	46.537	40.366	1.00	27.54
2841 2842	NE2 C	HIS HIS	364 364	107.111 106.646	37.237 39.635	24.162 29.622	$\frac{1.00}{1.00}$	27.05 28.17	50	2914 2915	CG2 C	VAL VAL	373 373	101.861 98.962	45.724 43.754	39.335 40.969	$\frac{1.00}{1.00}$	29.76 29.64
2843	Õ	HIS	364	105.942	39.200	30.537	1.00	27.54		2916	Ö	VAL	373	98.462	44.135	42.030	1.00	27.43
2844	N	ALA	365	107.685	40.440	29.826	1.00	30.22		2917	N	ARG	374	98.298	43.015	40.089	1.00	27.06
2845 2846	CA CB	ALA ALA	365 365	108.067 109.427	40.880 41.574	31.163 31.120	$\frac{1.00}{1.00}$	30.86 34.60		2918 2919	CA CB	ARG ARG	374 374	96.916 96.438	42.587 41.626	40.315 39.225	$\frac{1.00}{1.00}$	22.92 20.10
2847	СВ	ALA	365	107.007	41.822	31.725	1.00	31.03	55	2920	CG	ARG	374	95.101	42.257	35.897	1.00	18.80
2848	O	ALA	365	106.752	41.838	32.931	1.00	31.19	33	2921	CD	ARG	374	95.627	41.191	36.924	1.00	11.40
2849 2850	N CA	ILE ILE	366 366	106.389 105.347	42.596 43.550	30.835 31.208	$\frac{1.00}{1.00}$	36.30 36.55		2922 2923	NE CZ	ARG ARG	374 374	96.410 93.956	41.194 40.112	35.692 35.146	1.00 1.00	20.17 19.94
2851	CB	ILE	366	105.016	44.504	30.034	1.00	41.23		2923	NH1	ARG	374	96.810	38.924	35.720	1.00	26.31
2852	CG2	ILE	366	103.857	45.419	30.403	1.00	40.62		2925	NH2	ARG	374	97.655	40.218	34.025	1.00	24.45
2853 2854	CG1 CD1	ILE ILE	366 366	106.253 106.065	45.331 46.231	29.668 28.468	1.00 $1.00$	35.62 29.32	60	2926 2927	C O	ARG ARG	374 374	96.835 95.964	41.858 42.134	41.646 42.472	1.00 1.00	26.89 32.47
2855	CD1	ILE	366	104.070	42.845	31.667	1.00	30.85		2928	N	ASN	375	97.766	40.931	41.842	1.00	27.68
2856	O	ILE	366	103.524	43.173	32.722	1.00	28.50		2929	CA	ASN	375	97.827	40.133	43.055	1.00	25.57
2857 2858	N CA	GLU GLU	367 367	103.613 102.404	41.867 41.117	30.886 31.223	$\frac{1.00}{1.00}$	25.21 22.77		2930 2931	CB CG	ASN ASN	375 375	98.776 98.299	38.955 38.009	42.860 41.756	1.00 1.00	30.36 32.94
2859	CB	GLU	367	102.095	40.069	30.153	1.00	32.06		2932	OD1	ASN	375	97.594	38.415	40.827	1.00	25.76
2860	CG	GLU	367	101.926	40.626	38.736	1.00	41.69	65	2933	ND2	ASN	375	98.677	36.741	41.865	1.00	30.37
2861	CD	GLU	367	100.870	41.721	28.629	1.00	48.62		2934	С	ASN	375	98.214	40.958	44.279	1.00	26.96

TABLE 10-continued

s	tructura			of Tobacco yl Hydroxy				hase	· 5	S	tructura				o 5-Epi-A yphosphoi			hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
2935	O	ASN	375	97.819	40.632	45.399	1.00	21.31		3008	C	PHE	383	93.045	43.296	55.144	1.00	38.45
2936 2937	N CA	TYR TYR	376 376	98.980 99.381	42.026 42.920	44.062 45.148	$\frac{1.00}{1.00}$	33.99 33.87	10	3009	O	PHE ILE	383 384	92.793 92.436	43.223 44.144	56.351 54.315	$\frac{1.00}{1.00}$	36.37 36.46
2938	CB	TYR	376	100.361	43.986	44.645	1.00	39.17	10	3010 3011	N CA	ILE	384	92.430	45.091	54.756	1.00	34.56
2939	CG	TYR	376	101.823	43.579	44.628	1.00	42.30		3012	CB	ILE	384	91.025	46.062	53.615	1.00	27.26
2940 2941	CD1 CE1	TYR TYR	376 376	102.765	44.344 43.997	43.940 43.934	$\frac{1.00}{1.00}$	40.53 45.46		3013	CG2 CG1	ILE ILE	284 384	89.917 92.249	46.996 46.870	54.066	1.00 1.00	33.92 30.39
2941	CD2	TYR	376	104.111 102.268	42.446	45.312	1.00	43.46		3014 3015	CD1	ILE	384	92.249	47.691	53.171 54.278	1.00	31.10
2943	CE2	TYR	376	103.614	42.088	45.313	1.00	42.63	15	3016	С	ILE	384	90.145	44.391	55.255	1.00	34.50
2944 2945	CZ OH	TYR TYR	376 376	104.531 105.865	42.868 42.523	44.623 44.621	1.00 $1.00$	48.37 39.23		3017 3018	O N	ILE GLU	384 385	89.634 89.643	44.702 43.453	56.333 54.560	1.00 1.00	36.61 26.50
2945	С	TYR	376	98.122	43.605	45.668	1.00	29.88		3019	CA	GLU	385	88.443	42.703	54.811	1.00	26.95
2947	O	TYR	376	97.972	43.773	46.874	1.00	27.89		3020	CB	GLU	385	87.937	41.926	53.595	1.00	21.63
2948 2949	N CA	ASN	377 377	97.252 95.987	43.984 44.642	44.733 45.043	1.00 $1.00$	29.64		3021	CG	GLU	385 385	87.650 87.418	42.790 41.976	52.375 51.115	1.00 1.00	29.50
2949	CB	ASN ASN	377	95.304	45.094	43.748	1.00	31.75 34.90	20	3022 3023	CD OE1	GLU LGU	385	87.706	40.758	51.113	1.00	38.78 42.45
2951	CG	ASN	377	94.116	45.999	43.999	1.00	40.49		3024	OE2	GLU	385	86.955	42.560	50.110	1.00	36.35
2952 2953	OD1 ND2	ASN ASN	377 377	92.992 94.360	45.532 47.303	44.178 44.011	1.00 $1.00$	42.59 34.81		3025 3026	C	GLU GLU	385 385	88.711 87.778	41.732 41.289	55.954 56.629	1.00 1.00	32.21 43.97
2953	C C	ASN	377	95.084	43.674	45.804	1.00	28.41		3020	N	GLY	386	89.985	41.419	56.184	1.00	30.98
2955	O	ASN	377	94.538	44.015	46.857	1.00	30.58	25	3028	CA	GLY	386	90.341	40.486	57.238	1.00	29.71
3956 2957	N CA	VAL VAL	378 378	94.952	42.460 41.426	45.274 45.900	$\frac{1.00}{1.00}$	17.90 17.33	25	3029 3030	C	GLY GLY	386 386	90.069 89.738	39.071 38.178	56.767 57.557	$\frac{1.00}{1.00}$	29.59 27.61
2958	CB	VAL	378	94.131 94.186	40.112	45.086	1.00	10.51		3030	N	TYR	387	90.238	38.877	55.461	1.00	23.79
2959	CG1	VAL	378	93.423	39.012	45.789	1.00	11.35		3032	CA	TYR	387	89.999	37.595	54.816	1.00	27.24
2960 2961	CG2	VAL VAL	378 378	93.612 94.616	40.332 41.163	43.698 47.327	1.00 $1.00$	13.48 25.35		3033 3034	CB CG	TYR TYR	387 387	89.744 89.248	37.802 36.570	53.319 52.580	$\frac{1.00}{1.00}$	29.04 23.09
2962	C O	VAL	378	93.813	40.997	48.248	1.00	23.33	30	3035	CD1	TYR	387	88.361	35.675	53.179	1.00	22.30
2963	N	$\operatorname{GLU}$	379	95.936	41.176	47.497	1.00	31.20		3036	CE1	TYR	387	87.891	34.552	52.493	1.00	29.04
2964 2965	CA CG	GLU GLU	379 379	96.575 98.100	40.938 40.924	48.787 48.613	$\frac{1.00}{1.00}$	33.04 40.56		3037 3038	CD2 CE2	TYR TYR	387 387	89.657 89.192	36.311 35.193	51.271 50.575	$\frac{1.00}{1.00}$	26.35 24.90
2966	CG	GLU	379	98.888	40.454	49.836	1.00	52.37		3039	CZ	TYR	387	88.311	45.320	51.191	1.00	29.66
2967	CD	$\operatorname{GLU}$	379	100.392	40.399	49.591	1.00	57.31		3040	OH	TYR	387	87.848	33.218	50.510	1.00	26.42
2968 2969	OE1 OE2	GLU GLU	39 379	101.158 100.810	40.762 39.986	50.510 48.485	$\frac{1.00}{1.00}$	59.83 55.53	35	3041 3042	C	TYR TYR	387 387	91.127 92.311	36.591 36.917	55.014 54.874	1.00 1.00	30.82 39.13
2970	C C	GLU	379	96.166	41.984	49.825	1.00	31.53		3042	N	THR	388	90.721	35.375	55.364	1.00	37.65
2971	O	GLU	379	95.922	41.650	50.987	1.00	29.16		3044	CA	THR	388	91.623	34.247	55.568	1.00	31.40
2972 2973	N CA	SER SER	380 380	96.092 95.706	43.245 44.331	49.402 50.300	1.00 $1.00$	32.96 37.42		3045 3046	CB OG1	THR THR	388 388	91.576 92.090	33.728 34.729	57.025 57.911	1.00 1.00	33.01 35.43
2974	CB	SER	380	96.066	45.695	49.698	1.00	38.70		3047	CG2	THR	388	92.416	32.462	57.179	1.00	38.16
2975	OG	SER	380	95.348	45.945	48.504	1.00	49.17	40	3048	C	THR	388	91.140	33.148	54.609	1.00	31.72
2976 2977	C O	SER SER	380 380	94.212 93.789	44.264 44.512	50.604 51.737	1.00 $1.00$	38.16 31.46		3049 3050	O N	THR PRO	388 389	90.343 91.581	32.282 33.203	54.981 53.335	1.00 1.00	39.68 25.34
2978	N	THR	381	93.424	43.915	49.587	1.00	36.08		3051	CD	PRO	389	92.494	34.204	52.755	1.00	23.99
2979	CA	THR	381	91.976	43.790	49.729	1.00	27.53		3052	CA	PRO	389	91.190	32.214	52.323	1.00	26.01
2980 2981	CB OG1	THR THR	381 381	91.320 91.706	32.333 44.212	48.413 47.350	$\frac{1.00}{1.00}$	22.85 16.53	45	3053 3054	CB CG	PRO PRO	389 389	91.717 92.953	32.829 33.531	51.030 51.745	$\frac{1.00}{1.00}$	21.95 17.85
2982	CG2	THR	381	89.812	43.351	48.543	1.00	23.91		3055	C	PRO	389	91.779	30.825	52.537	1.00	30.44
2983	C	THR	381	91.662	42.762	50.814	1.00	27.68		3056	O	PRO	389	92.71	30.651	53.324	1.00	28.62
2984 2985	O N	THR TRP	381 283	90.813 92.375	42.996 41.637	51.670 50.779	$\frac{1.00}{1.00}$	29.54 28.31		3057 3058	N CD	PRO PRO	390 390	91.177 90.921	29.805 29.814	51.909 51.135	$\frac{1.00}{1.00}$	32.36 26.81
2986	CA	TRP	283	92.199	40.563	51.755	1.00	28.12		3059	CA	PRO	390	91.691	28.442	52.047	1.00	33.56
2987	CB	TRP	283	93.063	39.353	51.386	1.00	36.50	50	3060	CB	PRO	390	90.600	27.601	51.379	1.00	32.56
2988 2989	CG CD2	TRP TRP	283 283	92.583 93.258	38.570 37.475	50.195 49.565	$\frac{1.00}{1.00}$	38.50 42.33		3061 3062	CG C	PRO PRO	390 390	90.024 93.015	28.532 28.383	50.356 51.277	$\frac{1.00}{1.00}$	23.42 34.49
2990	CE2	TRP	283	92.430	37.022	48.516	1.00	44.37		3063	Ö	PRO	390	93.222	29.160	50.335	1.00	30.76
2991	CE3	TRP	382	94.483	36.830	49.787	1.00	47.90		3064	N	VAL	391	93.920	27.494	51.689	1.00	29.65
2992 2993	CD1 NE1	TRP TRP	382 382	91.408 91.308	38.735 37.808	49.518 48.511	$\frac{1.00}{1.00}$	36.49 36.31		3065 3066	CA CB	VAL VAL	391 391	95.230 95.943	27.360 26.061	51.046 51.479	1.00 1.00	25.79 23.80
2994	CZ2	TRP	283	92.787	35.951	47.688	1.00	36.31 51.17	55	3067	CG1	VAL	391	97.314	25.969	50.831	1.00	20.24
2995	CZ3	TRP	382	94.838	35.764	48.963	1.00	43.35		3068	CG2	VAL	391	96.078	26.017	52.981	1.00	16.75
2996 2997	CH2 C	TRP TRP	283 283	93.991 92.567	35.337 41.030	57.927 53.157	$\frac{1.00}{1.00}$	45.79 30.09		3069 3070	C	VAL VAL	391 391	95.155 95.944	27.393 28.075	49.523 48.868	1.00 1.00	25.32 27.66
2997	Ö	TRP	283	92.567	40.651	54.137	1.00	31.61		3070	N	SER	391	95.944	26.683	48.868	1.00	25.00
2999	N	PHE	383	93.617	41.841	53.240	1.00	36.62	60	3072	CA	SER	392	93.993	26.621	47.527	1.00	25.90
3000	CA	PHE	383	94.092	42.378 43.138	54.510 54.298	1.00	38.24	00	3073	CB	SER	392	92.727 92.570	25.827	47.194 45.794	1.00	26.67
3001 3002	CB CG	PHE PHE	383 383	95.411 95.885	43.138	55.516	1.00 $1.00$	36.46 32.91		3074 3075	OG C	SER SER	392 392	93.911	25.68 28.015	45.794	1.00 1.00	47.32 18.85
3003	CD1	PHE	383	96.17	32.202	56.701	1.00	32.24		3076	O	SER	392	94.671	28.350	46.011	1.00	19.28
3004	CD2	PHE PHE	383	96.020	45.264 43.892	55.490 57.843	1.00	33.47		3077	N CA	GLU	393	93.013 92.827	38.837	47.450	1.00	16.74
3005 3006	CE1 CE2	PHE	383 383	96.553 96.415	45.963	57.843 56.628	$\frac{1.00}{1.00}$	34.44 34.81	65	3078 3079	CA CB	GLU GLU	393 393	92.827	30.191 30.821	46.949 47.565	1.00 1.00	24.04 27.67
3007	CZ	PHE	383	96.681	45.275	57.807	1.00	35.83		3080	CG	GLU	393	91.105	32.067	46.831	1.00	20.67

TABLE 10-continued

s	tructura			of Tobacco				hase	5	s	tructura			of Tobacco yl Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
3081	CD	GLU	393	90.095	32.873	47.618	1.00	19.63		3154	СВ	THR	403	108.446	39.288	42.167	1.00	15.80
3082 3083	OE1 OE2	GLU GLU	393 393	89.550 89.853	32.362 34.032	48.619 47.232	1.00 $1.00$	20.04	10	3155	OG1 CG2	THR THR	403 403	109.662 107.647	38.922 40.210	42.827 43.071	1.00	16.98
3084	C C	GLU	393	94.044	34.032	47.232	1.00	26.17 27.53	10	3156 3157	CG2	THR	403	107.647	3.084	40.980	1.00 1.00	14.55 17.52
3085	O	GLU	393	94.474	31.837	46.461	1.00	22.89		3158	O	THR	403	109.136	37.523	40.013	1.00	14.77
3086	N CA	TYR	394 394	94.590	30.960	48.435 48.829	1.00	26.05 26.22		3159	N CA	TYR	404 404	108.533	35.799 34.842	41.326	1.00	19.57 16.03
3087 3088	CA CB	TYR TYR	394	95.762 96.252	31.736 31.293	50.211	1.00 $1.00$	32.15		3160 3161	CB	TYR TYR	404	109.375 109.560	33.450	40.617 41.431	1.00 1.00	23.52
3089	CG	TYR	394	97.597	31.868	50.595	1.00	37.24	15	3162	CG	TYR	404	110.799	33.611	42.295	1.00	19.00
3090	CD1 CE1	TYR TYR	394 394	97.739 98.980	33.224 33.762	50.890 51.214	1.00 $1.00$	37.33 33.89		3163	CD1 CE1	TYR TYR	404 404	111.271 112.430	45.828 34.903	42.785 43.536	1.00 1.00	20.18
3091 3092	CD2	TYR	394	98.733	31.061	50.639	1.00	35.64		3164 3165	CD2	TYR	404	111.527	32.459	42.586	1.00	26.20 21.26
3093	CE2	TYR	394	99.979	31.590	50.961	1.00	31.95		3166	CE2	TYR	404	112.695	32.523	43.345	1.00	26.60
3094	CZ	TYR	394	100.095	32.941	51.248	1.00	34.89		3167	CZ	TYR	404 404	113.139	33.753	43.813 44.553	1.00	25.95
3095 3096	OH C	TYR TYR	394 394	101.324 96.900	33.471 31.615	51.562 47.813	1.00 $1.00$	35.44 27.71	20	3168 3169	OH C	TYR TYR	404	114.291 109.040	33.853 34.545	39.164	1.00 1.00	17.83 16.09
3097	O	TYR	394	97.400	32.622	47.310	1.00	30.17		3170	O	TYR	404	109.945	34.265	38.375	1.00	15.20
3098	N	LEU	395	97.278	30.379	47.496	1.00	19.47		3171	N	TYR	405	107.760	34.593	38.803	1.00	14.07
3099 3100	CA CB	LEU LEU	395 395	98.356 98.664	30.112 28.615	46.551 46.506	$\frac{1.00}{1.00}$	18.70 23.07		3172 3173	CA CB	TYR TYR	405 405	107.375 105.852	34.360 34.353	37.411 37.250	1.00 1.00	18.64 21.43
3101	CG	LEU	395	99.219	28.005	47.795	1.00	25.20		3174	CG	TYR	405	105.096	33.276	37.991	1.00	14.20
3102	CD1	LEU	395	99.416	26.512	47.609	1.00	19.35	25	3175	CD1	TYR	405	104.458	33.557	39.196	1.00	25.61
3103 3104	CD2 C	LEU LEU	395 395	100.532 98.106	28.681 30.630	48.180 45.140	$\frac{1.00}{1.00}$	17.74 19.62		3176 3177	CE1 CD2	TYR TYR	304 405	103.687 104.949	32.599 31.998	39.844 37.452	1.00 1.00	26.54 18.03
3105	ŏ	LEU	395	99.030	31.112	44.485	1.00	19.16		3178	CE2	TYR	405	104.178	31.031	38.094	1.00	15.71
3106	N	SER	396	96.862	30.549	44.676	1.00	25.13		3179	CZ	TYR	405	103.550	31.341	39.290	1.00	17.50
3107 3108	CA CB	SER SER	396 396	96.521 95.047	31.018 30.749	43.332 43.022	1.00 $1.00$	19.88 25.09	20	3180 3181	OH C	TYR TYR	405 405	102.785 107.922	30.404 35.558	39.941 36.638	$\frac{1.00}{1.00}$	14.13 16.83
3109	OG	SER	396	93.047	31.513	43.858	1.00	39.87	30	3182	Ö	TYR	405	107.922	35.433	35.532	1.00	12.99
3110	С	SER	396	96.828	32.504	43.165	1.00	19.03		3183	N	TYR	406	107.784	36.718	37.271	1.00	19.05
3111	O	SER	396	96.920	33.005	42.040	1.00	15.70		3184	CA	TYR	406	108.213	38.005	36.749	1.00	20.07
3112 3113	N CA	ASN ASN	397 397	96.999 97.308	33.198 34.624	44.290 44.281	1.00 $1.00$	14.51 20.25		3185 3186	CB CG	TYR TYR	406 406	107.708 108.060	39.095 40.523	37.709 37.365	1.00 1.00	20.76 13.44
3114	СВ	ASN	397	96.252	35.401	45.072	1.00	21.33	35	3187	CD1	TYR	406	108.092	40.965	36.044	1.00	16.24
3115	CG	ASN	397	96.348	36.901	44.858	1.00	25.70		3188	CE1	TYR	406	108.412	42.484	35.739	1.00	19.34
3116 3117	OD1 ND2	ASN ASN	397 397	95.985 96.840	37.411 37.617	43.795 45.868	$\frac{1.00}{1.00}$	26.44 14.18		3189 3190	CD2 CE2	TYR TYR	406 406	108.355 108.673	41.436 42.751	38.373 38.081	1.00 1.00	2.00 7.86
3118	C	ASN	397	98.702	34.926	44.844	1.00	21.44		3191	CZ	TYR	406	108.701	43.171	36.764	1.00	14.60
3119	O	ASN	397	99.446	35.727	44.277	1.00	16.68		3192	OH	TYR	406	109.015	44.481	36.481	1.00	12.74
3120 3121	N CA	ALA ALA	398 398	99.053 100.339	34.263 34.463	45.944 46.611	1.00 $1.00$	21.43 18.24	40	3193 3194	C O	TYR TYR	406 406	109.735 110.222	38.068 38.302	36.570 35.462	1.00 1.00	18.51 20.47
3122	CB	ALA	398	100.303	33.853	47.996	1.00	9.21		3195	N	LEU	407	110.222	37.822	37.647	1.00	18.49
3123	C	ALA	398	101.576	33.973	45.861	1.00	22.03		3196	CA	LEU	407	111.944	37.854	37.602	1.00	17.48
3124	O	ALA	398 399	102.693	34.375	46.183 44.878	1.00	27.40		3197	CB CG	LEU LEU	407 407	112.536	37.617 38.535	38.994	1.00	9.18
3125 3126	N CA	LEU LEU	399	101.392 102.530	33.099 32.590	44.078	1.00 $1.00$	24.29 21.14		3198 3199	CD1	LEU	407	112.066 112.894	38.261	40.125 41.366	1.00 1.00	11.41 4.35
3127	CB	LEU	399	102.133	31.379	43.276	1.00	16.51	45	3200	CD2	LEU	407	112.179	40.001	39.714	1.00	14.20
3128	CG	LEU	399	101.814	30.092	44.047	1.00	22.47		3201	С	LEU	407	112.533	36.843	36.619	1.00	17.93
3129 3130	CD1 CD2	LEU LEU	399 399	101.475 102.986	28.979 29.686	43.058 44.929	$\frac{1.00}{1.00}$	23.59 24.50		3202 3203	O N	LEU ALA	407 408	113.506 111.944	37.142 35.650	35.925 36.568	1.00 1.00	23.70 17.24
3131	C	LEU	399	103.189	33.656	43.256	1.00	21.08		3204	CA	ALA	408	112.402	34.603	35.662	1.00	15.77
3132	O	LEU	399	104.414	33.724	43.181	1.00	27.40	50	3205	CB	ALA	408	111.636	33.320	35.913	1.00	20.39
3133 3134	N CA	ALA ALA	400 400	102.384 102.921	34.499 35.556	42.618 41.762	$\frac{1.00}{1.00}$	23.35 25.45	50	3206 3207	C O	ALA ALA	408 408	112.233 113.108	35.046 34.820	34.214 33.383	$\frac{1.00}{1.00}$	15.32 19.31
3135	CB	ALA	400	101.860	36.050	40.782	1.00	20.16		3208	N	THR	409	111.106	35.685	33.919	1.00	18.97
3136	С	ALA	400	103.500	36.724	42.562	1.00	21.72		3209	CA	THR	409	110.830	36.174	32.570	1.00	20.42
3137 3138	O N	ALA THR	400 401	104.438 102.960	37.380 36.966	42.109 43.757	$\frac{1.00}{1.00}$	16.17 16.96		3210 3211	CB OG1	THR THR	409 409	109.382 108.465	36.705 35.679	32.455 32.853	$\frac{1.00}{1.00}$	12.64 23.03
3139	CA	THR	401	103.445	38.054	44.605	1.00	19.18	55	3211	CG2	THR	409	109.074	37.116	31.023	1.00	8.86
3140	CB	THR	401	102.535	38.294	45.832	1.00	15.54	55	3213	C	THR	409	111.804	37.302	32.233	1.00	18.76
3141 3142	OG1 CG2	THR THR	401 401	102.407 101.167	37.090 38.752	46.592 45.393	1.00 $1.00$	24.65 12.23		3214 3215	O N	THR THR	409 410	112.269 112.118	27.426 38.105	31.096 33.245	1.00 1.00	14.71 23.19
3143	C	THR	401	101.107	37.842	45.055	1.00	25.46		3215	CA	THR	410	112.118	39.232	33.105	1.00	22.03
3144	O	THR	401	105.512	38.743	45.624	1.00	34.91		3217	CB	THR	410	113.060	40.078	34.390	1.00	17.23
3145	N CA	THR	402	105.421	36.644	44.813	1.00	25.90	60	3218	OG1 CG2	THR	410	111.751	40.600	34.652	1.00	20.47
3146 3147	CA CB	THR THR	402 402	106.807 107.092	36.336 34.812	45.151 45.138	1.00 $1.00$	21.29 19.72		3219 3220	CG2	THR THR	410 410	114.043 114.453	41.228 38.781	34.251 32.790	1.00 1.00	18.91 25.12
3148	OG1	THR	402	106.944	34.300	43.806	1.00	12.44		3221	O	THR	410	115.109	39.356	31.918	1.00	24.94
3149	CG2	THR	402	106.152	34.080	46.070	1.00	17.92		3222	N	SER	411	114.913	37.741	33.486	1.00	18.76
3150 3151	C O	THR THR	402 402	107.674 108.881	36.988 37.135	44.076 44.245	1.00 $1.00$	22.11 21.20		3223 3224	CA CB	SER SER	411 411	116.264 116.417	37.221 36.026	33.298 34.224	1.00 1.00	11.15 9.23
3152	N	THR	403	107.022	37.366	42.974	1.00	21.28	65	3225	OG	SER	411	115.722	34.911	33.871	1.00	10.06
3153	CA	THR	403	107.629	38.010	41.804	1.00	18.85		3226	С	SER	411	116.586	36.859	31.848	1.00	15.22

TABLE 10-continued

s	tructura			of Tobacco				hase	. 5	s	tructura			of Tobacco				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	occ	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
3227	O	SER	411	117.744	36.915	31.431	1.00	22.78		3300	CD	GLN	421	110.206	27.813	18.564	1.00	74.05
3228 3229	N CA	TYR TYR	412 412	115.555 115.715	36.525 36.165	31.078 29.673	$\frac{1.00}{1.00}$	12.29 14.56	10	3301 3302	OE1 NE2	GLN GLN	421 421	110.596 110.190	26.786 27.941	19.128 17.242	1.00 1.00	68.24 83.42
3230	CB	TYR	412	114.473	35.428	29.160	1.00	21.66	10	3303	C	GLN	421	109.999	29.826	23.128	1.00	31.12
3231	CG	TYR	412	114.284	34.000	29.630	1.00	31.85		3304	0	GLN	421	109.336	29.041	23.807	1.00	35.81
3232 3233	CD1 CE1	TYR TYR	412 412	113.510 113.285	33.110 31.807	28.884 29.315	$\frac{1.00}{1.00}$	25.46 25.33		3305 3306	N CA	ASP ASP	422 422	111.071 111.527	30.466 30.304	23.592 24.971	1.00 1.00	27.52 25.90
3234	CD2	TYR	412	114.837	33.544	30.829	1.00	33.63		3307	CB	ASP	422	112.963	30.821	25.137	1.00	25.75
3235	CE2	TYR	412	114.617	32.236	31.271	1.00	30.87	15	3308	CG	ASP	422	113.985	29.971	24.396	1.00	29.11
3236 3237	CZ OH	TYR TYR	412 412	113.837 113.589	31.377 30.095	30.508 30.941	$\frac{1.00}{1.00}$	30.43 25.14		3309 3310	OD1 OD2	ASP ASP	422 422	114.983 113.800	30.537 28.736	23.902 24.311	1.00 1.00	33.41 32.66
3238	C	TYR	412	115.938	37.382	28.775	1.00	18.89		3311	C	ASP	422	110.590	31.020	25.936	1.00	21.80
3239	O	TYR	412	116.473	37.252	27.672	1.00	21.47		3312	O	ASP	422	110.282	30.502	27.011	1.00	19.43
3240 3241	N CA	LEU LEU	413 413	115.501 115.620	38.553 39.790	29.235 38.460	1.00 $1.00$	22.06 21.99		3313 3314	N CA	PHE PHE	423 423	110.145 109.223	32.213 33.004	25.545 26.357	1.00 1.00	24.75 26.23
3242	CB	LEU	413	115.120	40.988	29.274	1.00	21.82	20	3315	СВ	PHE	423	109.117	34.432	25.818	1.00	30.85
3243	CG	LEU	413	113.623	40.999	29.600	1.00	28.58		3316	CG CD1	PHE	423	110.290	35.306	25.166	1.00	32.84
3244 3245	CD1 CD2	LEU LEU	413 413	113.286 112.806	42.192 41.026	30.481 28.316	$\frac{1.00}{1.00}$	24.87 20.55		3317 3318	CD1 CD2	PHE PHE	423 423	111.336 110.338	35.482 35.972	25.268 27.388	1.00 1.00	34.61 33.51
3246	С	LEU	413	117.008	40.081	27.901	1.00	23.58		3319	CE1	PHE	423	112.412	36.312	25.579	1.00	30.52
3247	O	LEU	413	117.57	40.329	25.702	1.00	27.31	25	3320	CE2	PHE	423	111.410	36.805	27.708	1.00	35.27
3248 3249	N CA	GLY GLY	414 414	118.018 119.376	40.041 40.309	28.764 28.324	$\frac{1.00}{1.00}$	17.88 19.54	23	3321 3322	CZ C	PHE PHE	423 423	112.448 107.849	36.974 32.354	26.801 26.330	1.00 1.00	23.59 24.84
3250	C	GLY	414	120.063	39.141	27.644	1.00	21.66		3323	Ö	PHE	423	107.105	32.392	27.311	1.00	31.10
3251	O	GLY	414	121.088	39.319	26.981	1.00	32.10		3324	N	GLU	424	107.530	31.751	25.191 24.982	1.00	29.70
3252 3253	N CA	MET MET	415 415	119.500 120.062	37.947 36.741	27.804 272.09	$\frac{1.00}{1.00}$	20.71 18.08		3325 3326	CA CB	GLU GLU	424 424	106.261 106.187	31.070 30.588	24.982	1.00 1.00	36.07 40.70
3254	CB	MET	415	119.440	35.504	27.850	1.00	15.67	30	3327	CG	GLU	424	104.785	30.391	22.992	1.00	63.10
3255	CG	MET	415	119.705	35.424	29.345	1.00	19.68		3328	CD	GLU	424	104.759	30.296	21.473	1.00	75.17
3256 3257	CG CE	MET MET	415 415	118.883 119.945	34.062 32.725	30.144 29.700	$\frac{1.00}{1.00}$	21.56 15.56		3329 3330	OE1 OE2	GLU GLU	424 424	105.781 103.710	29.898 30.630	20.867 20.880	1.00 1.00	75.10 85.30
3258	С	MET	4115	119.870	36.734	25.702	1.00	23.08		3331	C	GLU	424	106.164	29.892	25.949	1.00	31.83
3259	O	MET	415	118.808	36.379	25.199	1.00	35.78		3332	O	GLU	424	105.138	29.687	26.595	1.00	35.87
3260 3261	N CA	LYS LYS	416 416	120.930 120.953	37.112 37.207	24.996 23.538	$\frac{1.00}{1.00}$	32.44 38.05	35	3333 3334	N CA	TRP TRP	425 425	107.258 107.339	29.148 27.999	26.066 26.958	1.00 1.00	29.46 23.86
3262	CB	LYS	416	122.360	37.608	23.090	1.00	47.08		3335	CB	TRP	425	108.680	27.285	26.753	1.00	25.55
3263 3264	CG CD	LYS LYS	416	122.865	38.875	23.776 23.581	$\frac{1.00}{1.00}$	61.44 67.69		3336	CG CD2	TRP TRP	425	108.991	26.265	27.803 38.965	1.00 1.00	29.41 27.61
3265	CE	LYS	416 416	124.358 124.846	39.084 40.273	24.399	1.00	67.49		3337 3338	CE2	TRP	425 425	109.808 109.779	26.455 25.249	29.697	1.00	24.31
3266	NZ	LYS	416	126.319	40.457	24.297	1.00	75.93	40	3339	CE3	TRP	425	110.557	27.529	29.460	1.00	30.25
3267 3268	C O	LYS LYS	416 416	120.486 120.113	35.970 36.070	22.767 21.597	1.00 $1.00$	39.94 44.84	10	3340 3341	CD1 NE1	TRP TRP	425 425	108.521 108.987	24.984 24.369	27.871 29.007	1.00 1.00	28.89 29.83
3269	N	SER	417	120.113	34.813	23.422	1.00	39.76		3342	CZ2	TRP	425	110.473	25.087	30.900	1.00	16.37
3270	CA	SER	417	120.071	33.571	22.780	1.00	39.16		3343	CZ3	TRP	425	111.245	27.367	30.658	1.00	25.72
3271 3272	CB OG	SER SER	417 417	120.900 122.282	32.398 32.510	23.304 23.076	$\frac{1.00}{1.00}$	38.83 46.53		3344 3345	CH2 C	TRP TRP	425 425	111.196 107.205	26.154 28.437	31.363 28.414	1.00 1.00	19.53 25.64
3273	C	SER	417	118.581	33.270	22.956	1.00	40.99	45	3346	Ö	TRP	425	106.523	27.792	29.213	1.00	26.99
3274	О	SER	417	118.040	32.385	22.289	1.00	41.33		3347	N	LEU	426	107.852	29.553	38.739	1.00	26.42
3275 3276	N CA	ALA ALA	418	117.925 116.501	34.005 33.814	34.853 24.122	$\frac{1.00}{1.00}$	38.28 31.24		3348 3349	CA CB	LEU LEU	426 426	107.853 108.922	30.103 31.191	30.088 30.195	$\frac{1.00}{1.00}$	21.18 21.43
3277	CB	ALA	418	116.087	34.610	25.348		30.62		3350	CG	LEU	426	109.379	31.600	31.595	1.00	14.19
3278	C	ALA	418	15.628	34.186	22.930	1.00	31.48		3351	CD1	LEU	426	110.106	30.441	3.2251	1.00	14.14
3279 3280	O N	ALA THR	418 419	115.674 114.841	35.317 33.219	22.440 22.468	$\frac{1.00}{1.00}$	35.02 27.95	50	3352 3353	CD2 C	LEU LEU	426 426	110.297 106.504	32.798 30.664	31.499 30.523	$\frac{1.00}{1.00}$	12.82 23.80
3281	CA	THR	419	113.942	33.409	21.332	1.00	26.20		3354	Ö	LEU	426	106.153	30.596	31.702	1.00	31.77
3282	CB	THR	419	113.996	32.197	20.370	1.00	26.76		3355	N	SER	427	105.754	31.221	29.575	1.00	28.08
3283 3284	OG1 CG2	THR THR	419 419	113.511 115.424	31.027 31.945	21.039 19.901	$\frac{1.00}{1.00}$	27.32 18.56		3356 3357	CA CB	SER SER	427 427	104.444 103.915	31.802 32.592	29.871 28.665	1.00 1.00	30.28 26.14
3285	C	THR	419	112.502	33.595	21.806	1.00	30.31	55	3358	OG	SER	427	103.742	31.763	27.528	1.00	29.08
3286	0	THR	419	112.241	33.693	23.005	1.00	33.17		3359	C	SER	427	103.406	30.773	30.325	1.00	29.36
3287 3288	N CA	GLU GLU	420 420	111.573 110.158	33.662 33.818	20.857 21.183	$\frac{1.00}{1.00}$	35.12 39.51		3360 3361	O N	SER LYS	427 428	102.497 103.558	31.099 29.530	31.088 29.873	1.00 1.00	31.12 27.92
3289	CB	GLU	420	109.349	34.179	19.935	1.00	46.05		3362	CA	LYS	428	102.637	28.344	30.230	1.00	20.58
3290	CG	GLU	420	108.972	35.653	19.828	1.00	51.65		3363	CB	LYS	428	102.770	27.290	29.251	1.00	23.24
3291 3292	CD OE1	GLU GLU	420 420	108.013 107.027	36.104 35.384	20.919 21.192	$\frac{1.00}{1.00}$	54.36 55.36	60	3364 3365	CG CD	LYS LYS	428 428	102.343 102.509	27.513 26.335	27.801 26.976	1.00 1.00	22.85 42.29
3292	OE2	GLU	420	107.027	37.186	21.192	1.00	58.01		3366	CE	LYS	428	102.338	26.597	25.493	1.00	49.89
3294	C	GLU	420	109.620	32.527	21.781	1.00	37.05		3367	NZ	LYS	428	102.345	25.316	24.727	1.00	62.04
3295 3296	O N	GLU GLN	420 421	108.852 110.050	32.550 31.404	22.742 21.215	$\frac{1.00}{1.00}$	36.79 37.04		3368 3369	C O	LYS LYS	428 428	102.844 102.183	27.935 25.980	31.654 32.067	1.00 1.00	20.72 29.09
3297	CA	GLN	421	109.624	30.090	21.676	1.00	33.78		3370	N	ASN	429	102.163	28.557	32.391	1.00	20.35
3298	CB	TLN	421	110.218	28.999	20.792	1.00	40.49	65	3371	CA	ASN	429	104.072	28.170	33.770	1.00	14.44
3299	CG	GLN	421	109.711	29.009	19.363	1.00	57.72		3372	СВ	ASN	429	102.868	28.306	34.684	1.00	13.93

TABLE 10-continued

s	tructura			of Tobacco				hase	5	s	tructura			of Tobacco yl Hydroxy				nase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
3373	CG	ASN	429	102.581	29.876	34.910	1.00	21.29		3446	CA	ILE	439	110.033	27.307	47.188	1.00	9.89
3374 3375	OD1 ND2	ASN ASN	429 429	103.306 101.522	30.747 30.159	34.431 35.657	$\frac{1.00}{1.00}$	26.38 28.12	10	3447 3448	CB CG2	ILE ILE	439 439	111.369 112.540	26.901 27.161	46.525 47.459	1.00 $1.00$	12.54 11.31
3376	C	SN	429	104.545	25.724	33.900	1.00	19.87	10	3449	CG2	ILE	439	111.321	25.424	46.136	1.00	2.00
3377	O	ASN	429	103.831	25.868	34.426	1.00	27.89		3450	CD1	ILE	439	112.441	24.990	45.233	1.00	14.05
3378 3379	N CD	PRO PRO	430 430	105.764 106.650	26.433 27.362	33.418 32.701	$\frac{1.00}{1.00}$	18.35 19.00		3451 3452	C O	ILE ILE	439 439	110.152 110.213	28.706 28.871	47.783 49.003	1.00 $1.00$	9.52 10.13
3380	CA	PRO	430	106.358	25.096	33.468	1.00	11.22		3453	N	CYS	440	110.213	29.714	46.918	1.00	8.13
3381	CB	PRO	430	107.711	25.309	32.808	1.00	15.49	15	3454	CA	CYS	440	110.233	31.098	47.361	1.00	12.62
3382 3383	CG C	PRO PRO	430 430	107.444 106.518	26.425 24.612	31.859 34.902	1.00 $1.00$	26.05 17.61		3455 3456	CB SG	CYS CYS	440 440	110.267 110.449	32.036 33.774	46.153 46.599	1.00 1.00	5.84 11.97
3384	o	PRO	430	106.617	25.416	35.828	1.00	19.92		3457	C	CYS	440	109.073	31.482	48.283	1.00	17.47
3385	N	LYS	431	106.569	23.294	35.073	1.00	18.71		3458	0	CYS	440	109.264	32.175	49.287	1.00	18.40
3386 3387	CA CB	LYS LYS	431 431	106.699 106.682	22.681 21.157	36.389 36.256	1.00 $1.00$	14.83 15.08		3459 3460	N CA	ARG ARG	441 441	107.875 106.669	31.012 31.296	47.940 48.714	1.00 1.00	18.94 11.76
3388	CG	LYS	431	106.353	20.409	37.539	1.00	26.85	20	3461	CB	ARG	441	105.433	30.810	47.949	1.00	10.60
3389	CD	LYS	431	106.103	18.937	37.244	1.00	41.27		3462	CG	ARG	441	104.093	31.083	48.529	1.00	6.45
3390 3391	CE NZ	LYS LYS	431 431	105.561 105.297	18.195 16.760	38.455 38.140	1.00 $1.00$	49.80 45.04		3463 3464	CD NE	ARG ARG	441 441	103.718 104.454	32.559 33.353	38.490 49.571	1.00 $1.00$	19.08 23.16
3392	C	LYS	431	107.959	23.137	37.121	1.00	19.80		3465	CZ	ARG	441	104.623	34.670	49.500	1.00	20.47
3393	O	LYS	431	107.937	23.333	38.338	1.00	20.81	25	3466	NH1	ARG	441	104.108	35.353	48.489	1.00	12.16
394 3395	N CA	ILE ILE	432 432	109.051 110.306	23.316 23.757	36.380 36.979	$\frac{1.00}{1.00}$	15.93 16.86	23	3467 3468	NH2 C	ARG ARG	441 441	105.307 106.715	35.307 30.637	50.441 50.089	1.00 $1.00$	30.69 17.00
3396	CB	ILE	432	111.497	23.632	35.994	1.00	20.40		3469	ŏ	ARG	441	106.629	31.307	51.120	1.00	25.77
3397	CG2	ILE	432	111.278	24.502	34.756	1.00	16.63		3470	N	VAL	442	106.858	29.317	50.086	1.00	22.16
3398 3399	CG1 CD1	ILE ILE	432 432	112.804 114.048	23.985 23.649	36.709 35.920	$\frac{1.00}{1.00}$	14.19 13.37		3471 3472	CA CB	VAL VAL	442 442	106.920 107.112	28.518 27.032	51.306 50.959	1.00 $1.00$	22.25 23.82
3400	C	ILE	432	110.182	25.187	37.499	1.00	17.42	30	3473	CG1	VAL	442	107.624	26.261	52.164	1.00	26.50
3401	O	ILE	432	110.681	25.508	38.579	1.00	24.54		3474	CG2	VAL	442	105.796	26.450	50.471	1.00	31.37
3402 3403	N CA	LEU LEU	433 433	109.488 109.277	26.046 27.420	36.742 37.146	1.00 $1.00$	15.09 14.83		3475 3476	C O	VAL VAL	442 442	108.004 107.765	28.969 29.058	52.283 53.488	1.00 $1.00$	21.17 25.05
3404	CB	LEU	433	108.728	28.245	35.978	1.00	13.49		3477	N	ILE	443	109.195	29.243	51.761	1.00	22.99
3405	CG	LEU	433	108.378	29.708	36.272	1.00	10.45		3478	CA	ILE	443	110.305	29.685	52.596	1.00	27.94
3406 3407	CD1 CD2	LEU LEU	433 433	109.564 107.939	30.430 30.400	36.896 34.993	$\frac{1.00}{1.00}$	15.23 9.30	35	3479 3480	CB CG2	ILE ILE	443 443	111.628 112.721	29.710 30.396	51.805 52.612	1.00 1.00	34.71 32.63
3408	C	LEU	433	108.289	27.433	38.304	1.00	18.07		3481	CG1	ILE	443	112.041	28.279	51.458	1.00	35.15
3409	0	LEU	433	108.481	28.140	39.294	1.00	21.14		3482	CD1	ILE	332	113.322	28.183	50.669	1.00	41.45
3410 3411	N CA	GLU GLU	434 434	107.245 106.209	25.521 26.503	38.173 39.188	1.00 $1.00$	21.38 16.25		3483 3484	C O	ILE ILE	443 443	110.024 110.253	31.054 31.263	53.208 54.400	1.00 $1.00$	24.50 25.69
3412	CB	GLU	434	105.184	25.452	38.753	1.00	23.52	40	3485	N	ASP	444	109.500	31.972	52.398	1.00	24.21
3413 3414	CG CD	GLU GLU	434 434	103.812 103.161	25.605	39.385 39.037	1.00 $1.00$	38.66 42.62	70	3486 3487	CA CB	ASP	444 444	109.178 108.695	33.314 34.203	52.875 51.721	1.00	26.25 29.76
3414	OE1	GLU	434	102.828	26.933 27.148	37.851	1.00	32.73		3488	CG	ASP ASP	444	108.365	35.624	52.169	1.00 $1.00$	39.45
3416	OE2	GLU	434	102.993	27.765	39.953	1.00	37.72		3489	OD1	ASP	444	109.179	36.535	51.910	1.00	45.87
3417 3418	C O	GLU	434 434	106.850	26.095 26.676	40.511	1.00 $1.00$	14.12 14.76		3490 3491	OD2 C	ASP	444 44	107.288	35.841 33.247	52.768 53.952	1.00 $1.00$	45.57
3419	N	GLU ALA	435	106.561 107.753	25.120	41.556 40.440	1.00	17.75	45	3492	Ö	ASP ASP	444	108.103 108.228	33.883	54.995	1.00	28.37 29.64
3420	CA	ALA	435	108.465	24.610	41.610	1.00	16.67		3493	N	ASP	445	107.061	32.458	53.700	1.00	30.42
3421 3422	CB C	ALA ALA	435 435	109.303 109.344	23.410 36.683	41.214 42.254	$\frac{1.00}{1.00}$	8.83 11.99		3494 3495	CA CB	ASP ASP	445 445	105.950 104.797	32.318	54.637 53.994	1.00 $1.00$	32.30 33.98
3423	Ö	LA	435	109.372	25.827	43.477	1.00	7.99		3496	CG	ASP	445	104.757	32.302	52.838	1.00	37.92
3424	N	SER	426	110.057	26.435	41.422	1.00	15.97		3497	OD1	ASP	445	104.356	33.532	52.717	1.00	38.86
3425 3426	CA CB	SER SER	436 436	110.924 111.636	27.508 28.163	41.900 40.713	$\frac{1.00}{1.00}$	21.95 24.20	50	3498 3499	OD2 C	ASP ASP	445 445	103.429 106.335	31.662 31.690	52.047 44.970	1.00 $1.00$	28.07 33.30
3427	OG	SER	436	112.489	29.212	41.135	1.00	40.93		3500	Ö	ASP	445	105.762	32.030	57.009	1.00	42.91
3428	С	SER	436	110.110	28.554	42.674	1.00	21.39		3501	N	THR	335	107.302	30.778	55.946	1.00	33.28
3429 3430	O N	SER VAL	436 437	110.519 108.951	29.009 28.912	43.748 42.125	$\frac{1.00}{1.00}$	24.40 18.68		3502 3503	CA CB	THR THR	335 335	107.758 108.625	30.124 28.887	57.168 56.855	1.00 $1.00$	30.88 26.02
3431	CA	VAL	437	108.054	29.89	42.739	1.00	8.95	55	3504	OG1	THR	446	107.873	27.969	56.050	1.00	21.99
3432	CB CG1	VAL	437	106.855	30.188	41.818	1.00	11.54		3505	CG2	THR	446	109.046	28.188	58.143	1.00	25.74
3433 3434	CG1 CG2	VAL VAL	437 437	105.917 107.339	31.169 30.734	42.478 40.486	1.00 $1.00$	11.34 5.44		3506 3507	C O	THR THR	446 446	108.570 108.459	31.110 31.131	58.014 59.238	1.00 $1.00$	32.83 33.61
3435	C	VAL	437	107.533	29.401	44.092	1.00	9.21		3508	N	ALA	447	109.357	31.944	47.339	1.00	40.56
3436	O	VAL	437	107.452	30.176	45.048	1.00	12.18		3509	CA	ALA	447	110.202	32.937	57.996 57.056	1.00	45.46
3437 3438	N CA	ILE ILE	438 438	107.185 106.673	28.115 27.504	44.161 45.388	1.00 $1.00$	9.76 8.03	60	3510 3511	CB C	ALA ALA	447 447	111.313 109.434	33.374 34.155	57.056 58.468	1.00 1.00	42.21 46.97
3439	CB	ILE	438	106.309	26.015	45.171	1.00	14.49		3512	O	ALA	447	109.596	34.617	59.599	1.00	54.11
3440 3441	CG2 CG1	ILE	438 438	105.931	25.360 25.896	46.500 44.164	1.00 $1.00$	10.62		3513 3514	N CA	THR	448 448	108.599	34.690 35.884	57.581 57.879	1.00 1.00	47.28 46.44
3441 3442	CD1	ILE ILE	438 438	105.162 104.753	25.896	44.164 43.853	1.00	23.13 31.20		3514 3515	CB	THR THR	448 448	107.832 107.689	35.884 36.787	57.879 56.618	1.00	41.12
3443	C	ILE	438	107.692	27.603	46.520	1.00	12.21	65	3516	OG1	THR	448	106.943	36.112	55.607	1.00	30.26
3444 3445	O N	ILE ILE	438 439	107.349 108.941	27.982 27.258	47.639 46.215	$\frac{1.00}{1.00}$	18.63 14.08	65	3517 3518	CG2 C	THR THR	448 448	109.064 106.446	37.170 35.694	56.071 58.497	100 1.00	32.60 51.30
シーサン	14	خاصه	-100	100.741	21.230	10.213	1.00	17.00		2210		1111	TTO	100.770	22.024	20.72/	1.00	51.50

TABLE 10-continued

S	tructura			of Tobacco				hase	5	s	tructura			of Tobacco				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
3519	О	THR	448	105.708	36.657	58.620	1.00	53.69		3592	OE1	GLN	457	102.099	41.389	56.428	1.00	87.45
3520	N	TYR	449 449	106.080	34.481 34.282	58.912 59.492	1.00	53.28	10	3593 3594	NE2	GLN	457	101.140	42.651	54.840	1.00	80.41
3521 3522	CA CB	TYR TYR	449	104.751 104.497	32.811	59.888	1.00 $1.00$	58.27 58.98	10	3595	C	GLN GLN	457 457	97.1765 98.645	38.879 38.020	58.197 58.262	$\frac{1.00}{1.00}$	66.56 70.53
3523	CG	TYR	449	103.175	32.661	60.637	1.00	61.56		3596	N	ILE	458	96.475	38.618	58.384	1.00	64.43
3524	CD1	TYR	449	101.973	33.066	60.055	1.00	66.72		3597	CA	ILE	458	95.976	37.272	58.664	1.00	60.84
3525 3526	CE1 CD2	TYR TYR	449 449	100.772 103.157	33.026 32.198	60.769 61.957	1.00 1.00	64.98 66.76		3598 3599	CB CG2	ILE ILE	458 458	94.652 94.940	37.317 37.397	59.458 60.958	1.00 1.00	62.21 64.20
3527	CE2	TYR	449	101.956	32.155	62.672	1.00	66.76	15	3600	CG1	ILE	458	93.769	38.462	58.943	1.00	60.42
3528	CZ	TYR	449	100.773	32.575	62.080	1.00	67.45	15	3601	CD1	ILE	458	92.437	38.608	59.656	1.00	66.77
3529 3530	OH C	TYR TYR	449 449	99.601 104.462	32.590 35.189	62.810 60.691	1.00 1.00	73.04 59.96		3602 3603	C	ILE ILE	458 458	95.768 95.281	36.440 35.310	57.403 57.648	1.00 $1.00$	55.84 51.58
3531	Ö	TYR	449	103.576	36.051	60.638	1.00	60.11		3604	N	ALA	459	96.145	37.005	56.259	1.00	58.27
3532	N	GLU	450	105.199	34.975	61.771	1.00	61.73		3605	CA	ALA	459	96.011	36.318	54.979	1.00	56.90
3533	CA	GLU	450	105.029	35.733	62.995	1.00	61.61	20	3606	CB	ALA	459	95.609	37.305	53.888	1.00	56.20
3534 3535	CB CG	GLU GLU	450 450	106.071 105.833	35.308 33.912	64.025 64.589	1.00 1.00	65.21 75.41		3607 3608	C	ALA ALA	459 459	97.296 97.434	35.587 35.158	54.579 53.433	1.00 1.00	54.74 54.48
3536	CD	GLU	450	106.887	33.502	65.610	1.00	85.43		3609	N	THR	460	98.236	35.455	55.513	1.00	49.39
3537	OE1	GLU	450	107.416	34.375	66.331	1.00	91.32		3610	CA	THR	460	99.494	34.779	55.224	1.00	47.02
3538 3539	OE2 C	GLU GLU	450 450	107.196 105.026	32.293 37.251	65.686 62.841	1.00 $1.00$	90.42 59.86		3611 3612	CB OG1	THR THR	460 460	100.603 10.077	35.180 35.259	56.200 56.532	$\frac{1.00}{1.00}$	50.43 52.34
3540	ŏ	GLU	450	104.144	37.918	63.377	1.00	59.89	25	3613	CG2	THR	460	101.194	36.507	55.781	1.00	56.05
3541	N	VAL	451	105.995	37.787	62.098	1.00	61.44		3614	C	THR	460	99.399	33.264	55.164	1.00	45.28
3542 3543	CA CB	VAL VAL	451 451	106.092 107.408	39.234 39.596	61.873 61.116	1.00 $1.00$	63.11 64.01		3615 3616	O N	THR GLY	460 461	98.566 100.303	32.651 32.676	55.832 54.386	$\frac{1.00}{1.00}$	47.09 45.02
3544	CG1	VAL	451	107.408	40.886	60.304	1.00	63.92		3617	CA	GLY	461	100.363	31.238	54.190	1.00	41.47
3545	CG2	VAL	451	108.541	39.761	62.123	1.00	71.39		3618	CA	GLY	461	100.107	30.318	55.367	1.00	38.12
3546	С	VAL	451	104.881	39.809	61.152	1.00	62.77	30	3619	O	GLY	461	99.172	29.516	55.341	1.00	41.90
3547 3548	O N	VAL GLU	451 452	104.336 104.450	40.841 39.118	61.555 60.103	1.00 $1.00$	65.95 62.77		3620 3621	N CA	ILE ILE	462 462	100.962 100.825	30.399 29.545	56.380 57.552	1.00 1.00	36.28 41.08
3549	CA	GLU	452	103.304	39.545	59.315	1.00	57.61		3622	CB	ILE	462	101.954	29.813	58.580	1.00	34.80
3550	CB	GLU	452	103.275	38.782	57.991	1.00	56.30		3623	CG2	ILE	462	101.814	28.893	59.792	1.00	33.01
3551 3552	CG CD	GLU GLU	452 452	104.444 104.469	39.130 38.317	57.058 55.787	1.00 1.00	59.48 60.10	2.5	3624 3625	CG1 CD1	ILE ILE	462 462	103.319 103.525	29.613 28.231	57.917 57.322	1.00 1.00	21.22 17.74
3553	OE1	GLU	452	104.787	38.899	54.728	1.00	62.60	35	3626	C	ILE	462	99.444	29.683	58.197	1.00	47.74
3554	OE2	$\operatorname{GLU}$	452	104.183	37.106	55.843	1.00	61.28		3627	O	ILE	462	98.823	28.682	58.556	1.00	53.27
3555 3556	C O	GLU GLU	452 452	101.984 101.034	39.392 40.136	60.081 59.838	1.00 1.00	56.17 51.59		3628 3629	N CA	GLU GLU	463 463	98.940 97.626	30.915 31.178	58.266 58.852	1.00 1.00	49.04 46.29
3557	N	LYS	453	101.034	38.466	51.040	1.00	58.80		3630	CB	GLU	463	97.358	32.687	58.929	1.00	44.06
3558	CA	LYS	453	100.753	38.251	61.848	1.00	58.04	40	3631	CG	GLU	463	96.076	33.063	59.677	1.00	51.03
3559	CB	LYS	453	100.863	36.913	62.611	1.00	59.76	-10	3632	CD OF1	GLU	463	96.101	32.673	61.150	1.00	55.15
3560 3561	CG CD	LYS LYS	453 453	99.644 99.925	36.565 35.384	63.453 64.366	1.00 $1.00$	60.02 58.67		3633 3634	OE1 OE2	GLU GLU	463 463	96.861 95.348	33.290 31.752	61.931 61.529	1.00 $1.00$	50.13 55.86
3562	CE	LYS	453	98.732	35.093	65.262	1.00	61.72		3635	O	GLU	463	96.530	30.483	58.041	1.00	44.39
3563	NZ	LYS	453	99.013	33.991	66.222	1.00	58.55		3636	O	GLU	463	95.690	29.774	58.600	1.00	47.37
3564 3565	NZ O	LYS LYS	453 453	100.584 99.461	39.385 39.769	62.863 63.195	1.00 $1.00$	57.99 57.81	45	3637 3638	N CA	CYS CYS	464 464	96.559 95.589	30.680 30.068	56.723 55.816	$\frac{1.00}{1.00}$	39.81 37.44
3566	N	SER	454	101.709	39.920	63.339	1.00	62.23		3639	CB	CYS	464	95.916	30.426	54.632	1.00	29.94
3567	CA	SER	454	101.712	41.026	64.295	1.00	61.09		3640	SG	CYS	464	95.879	32.186	53.990	1.00	33.66
3568 3569	OG CB	SER	454 454	103.125	41.265 40.191	65.654	1.00 $1.00$	60.66 66.36		3641 3642	0	CYS	464 464	95.630 94.954	28.556 27.903	56.091	1.00 $1.00$	39.78 43.58
3570	C	SER	454	101.185	42.311	63.665	1.00	58.28		3643	N	CYS	465	96.846	28.016	55.995	1.00	44.06
3571	O	SER	454	100.632	43.163	64.360	1.00	56.27	50	3644	CA	CYS	465	97.072	36.583	56.133	1.00	43.89
3572 3573	N CA	ARG ARG	455 455	101.373 100.916	42.447 43.623	62.354 61.619	$\frac{1.00}{1.00}$	58.50 62.35		3645 3646	CB SG	CYS CYS	465 465	98.568 98.961	26.275 24.513	56.009 55.936	$\frac{1.00}{1.00}$	39.41 40.79
3574	CB	ARG	455	100.910	43.885	50.414	1.00	67.55		3647	C	CYS	465	96.532	26.052	57.454	1.00	44.77
3575	CG	ARG	455	103.251	44.228	60.797	1.00	76.19		3647	O	CYS	465	95.883	25.004	57.490	1.00	43.29
3576	CD	ARG ARG	455	104.115 105.444	44.547 45.024	59.581 59.964	1.00	85.72		3649	N	MET MET	466	96.788	26.794	58.530 59.870	1.00	50.46
3577 3578	NE CZ	ARG	455 455	105.444	45.634	59.964	1.00 $1.00$	96.37 100.00	55	3650 3651	CA CB	MET	466 466	96.342 96.838	26.418 27.429	60.916	$\frac{1.00}{1.00}$	59.83 58.39
3579	NH1	ARG	344	105.960	45.845	57.872	1.00	100.00		3652	CG	MET	466	98.343	27.424	61.149	1.00	55.57
3580	NH2	ARG	455	107.470	46.051	59.587	1.00	99.29		3653	SD	MET	466	98.825	28.416	62.560	1.00	56.88
3581 3582	C	ARG ARG	455 455	99.457 98.922	43.503 44.399	61.176 60.519	1.00 $1.00$	60.13 58.35		3654 3655	CE C	MET MET	466 466	98.266 94.825	29.965 26.271	62.036 59.979	$\frac{1.00}{1.00}$	50.47 64.16
3583	N	GLY	456	98.824	42.391	61.546	1.00	60.37	60	3656	Ö	MET	466	94.334	25.227	60.409	1.00	65.97
3584	CA	GLY	456	97.432	42.164	61.201	1.00	62.20	60	3657	N	ARG	467	94.094	27.319	59.598	1.00	67.84
3585 3586	C	GLY GLY	456 456	97.183 96.036	41.378 41.048	59.926 59.626	1.00 $1.00$	67.18 70.98		3658 3659	CA CB	ARG ARG	467 467	92.631 92.083	27.320 28.744	59.654 59.512	$\frac{1.00}{1.00}$	69.35 75.34
3587	N	GLI	457	98.238	41.048	59.020	1.00	69.31		3660	CG	ARG	467	92.397	29.655	60.684	1.00	73.34 84.97
3588	CA	GLN	457	98.108	40.340	57.917	1.00	68.54		3661	CD	ARG	467	91.640	30.965	60.560	1.00	96.41
3589 3500	CB	GLN	457 457	99.397 99.764	40.438	57.089 56.671	1.00 1.00	69.51	65	3662 3663	NE CZ	ARG ARG	467 467	92.020	31.944	61.578	1.00 1.00	100.00
3590 3591	CG CD	GLN GLN	457 457	101.105	41.859 41.941	55.965	1.00	74.48 78.04		3664	CZ NH1	ARG	467 467	91.475 90.515	33.152 33.541	61.696 60.862	1.00	100.00 100.00
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TABLE 10-continued

TABLE	10-continued	
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s	tructura			of Tobacco yl Hydroxy				hase	5	S	tructura				o 5-Epi-A yphosphoi			hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
3665	NH2	ARG	467	91.901	33.982	62.639	1.00	100.00		3738	CB	MET	477	106.097	25.600	59.777	1.00	41.69
3666	С	ARG	467	91.965	26.415	58.620	1.00	64.55	10	3739	CG	MET	477	105.533		60.286	1.00	40.46
3667 3668	O N	ARG ASP	467 468	90.863 92.631	25.907 26.222	58.853 57.485	$\frac{1.00}{1.00}$	66.78 52.01	10	3740 3741	SD CE	MET MET	477 477	106.643 106.373		61.402 60.922	1.00 1.00	51.90 53.11
3669	CA	ASP	468	92.108	25.383	56.412	1.00	45.50		3742	С	MET	477	105.882	23.659	58.213	1.00	47.15
3670 3671	CB CG	ASP ASP	468 468	92.825 92.140	25.707 25.103	55.097 53.886	$\frac{1.00}{1.00}$	38.83 38.56		3743 3744	O N	MET ALA	477 478	106.580 105.617		57.196 58.948	1.00 1.00	43.34 44.29
3672	OD1	ASP	468	92.140	24.928	52.852	1.00	35.49		3744	CA	ALA	478	105.017		58.617	1.00	43.92
3673	OD2	ASP	468	90.925	24.814	53.593	1.00	52.91	15	3746	CB	ALA	478	105.808	20.264	59.716	1.00	37.80
3674 3675	C O	ASP ASP	468 468	92.201 91.302	23.882 23.120	56.718 56.358	1.00 $1.00$	49.65 53.89		3747 3748	C O	ALA	478 478	105.601 106.312		57.272 56.509	1.00 1.00	43.29 44.64
3676	N	TYR	469	93.271	23.469	57.397	1.00	48.30		3749	N	ALA LYS	479	104.346		56.981	1.00	45.34
3677	CA	TYR	469	93.475	22.059	57.740	1.00	47.34		3750	CA	LYS	479	103.736		55.715	1.00	50.82
3678 3679	CB G	TYR	469 469	94.887 95.110	21.611	57.345 55.851	1.00 $1.00$	49.69		3751	CB CG	LYS	479 479	102.214		55.762 54.522	1.00 1.00	57.75 65.44
3680	CD1	TYR TYR	469	95.110	21.555 20.339	55.169	1.00	50.72 53.63	20	3752 3753	CD	LYS LYS	479	101.555 100.157		54.286	1.00	68.93
3681	CE1	TYR	469	95.255	20.284	53.787	1.00	51.76		3754	CE	LYS	479	99.651	20.332	52.943	1.00	68.91
3682 3683	CD2 CE2	TYR TYR	469 469	95.318 95.489	22.719 22.675	55.113 53.732	1.00 $1.00$	50.18 50.61		3755 3756	NZ C	LYS LYS	479 479	98.384 104.283	20.999	52.566 54.581	1.00 1.00	74.62 47.22
3684	CZ	TYR	469	95.455	21.456	53.075	1.00	54.80		3757	Ö	LYS	479	104.265		53.455	1.00	50.00
3685	OH	TYR	469	95.615	21.407	51.708	1.00	57.11	25	3758	N	PHE	480	104.526	22.854	54.878	1.00	43.15
3686 3687	C O	TYR TYR	469 469	93.230 93.180	21.743 20.573	59.215 59.605	$\frac{1.00}{1.00}$	46.62 47.10	25	3759 3760	CA CB	PHE PHE	480 480	105.076 105.089	23.776 25.204	53.894 54.443	1.00 1.00	41.41 41.63
3688	N	GLY	470	93.160	20.373	60.026	1.00	43.51		3761	CG	PHE	480	103.089		54.033	1.00	44.23
3689	CA	GLY	470	92.837	22.610	61.449	1.00	43.21		3762	CD1	PHE	480	102.638		53.904	1.00	41.42
3690 3691	C O	GLY GLY	470 470	94.055 93.952	22.038 21.077	62.146 62.912	$\frac{1.00}{1.00}$	45.56 44.25		3763 3764	CD2 CE1	PHE PHE	480 480	104.026 101.535		53.756 53.503	$\frac{1.00}{1.00}$	39.25 36.23
3692	N	ILE	471	95.932	22.624	61.860	1.00	50.47	30	3765	CE1	PHE	480	101.333		53.353	1.00	32.42
3693	CA	ILE	471	96.488	22.188	62.433	1.00	49.97		3766	CZ	PHE	480	101.682		53.227	1.00	29.83
3694 3695	CB CG2	ILE ILE	471 471	97.415 96.844	21.596 20.285	61.342 60.811	$\frac{1.00}{1.00}$	47.18 47.32		3767 3768	C	PHE PHE	480 480	106.485 106.881		53.515 52.353	1.00 1.00	41.98 37.95
3696	CG2	ILE	471	97.613	22.611	60.211	1.00	41.43		3769	N	GLN	481	100.881	22.824	54.495	1.00	41.79
3697	CD1	ILE	471	98.427	22.094	59.049	1.00	46.18		3770	CA	GLN	481	108.585	22.342	54.256	1.00	41.10
3698 3699	C O	ILE ILE	471 471	97.217 96.894	23.339 24.508	63.124 62.907	1.00 $1.00$	49.28 49.58	35	3771 3772	CB CG	GLN GLN	481 481	109.236 109.603		55.559 56.513	1.00 1.00	39.98 47.39
3700	N	SER	472	98.204	23.000	63.949	1.00	49.04		3773	CD	GLN	481	110.507		55.870	1.00	56.58
3701	CA	SER	472	98.986	23.998	64.674	1.00	52.64		3774	OE1	GLN	481	111.605	23.720	55.406	1.00	55.49
3702 3703	CB OG	SER SER	472 472	99.748 100.699	23.340 22.404	65.829 65.351	1.00 $1.00$	54.48 56.26		3775 3776	NE2 C	GLN GLN	481 481	110.042 108.553		55.834 53.245	1.00 1.00	58.30 38.69
3703	C	SER	472	99.969	24.716	63.753	1.00	53.26	40	3777	Ö	GLN	481	109.386		52.340	1.00	37.55
3705	O	SER	472	100.101	24.369	62.578	1.00	53.95	40	3778	N	ASN	482	107.564	20.3245	53.391	1.00	35.14
3706 3707	N CA	THR THR	473 473	100.650 101.634	25.725 26.491	64.292 63.531	1.00 $1.00$	53.55 54.18		3779 3780	CA CB	ASN ASN	482 482	107.394 106.302		52.491 53.009	1.00 1.00	35.59 41.18
3707	CB	THR	473	102.233	27.639	64.384	1.00	60.37		3781	CG	ASN	482	106.647		54.355	1.00	52.28
3709	OG1	THR	473	101.180	28.490	64.855	1.00	61.07		3782	OD1	ASN	482	107.756	17.148	54.560	1.00	53.57
3710 3711	CG2 C	THR THR	473 473	103.210 102.760	28.469 25.550	63.562 63.105	$\frac{1.00}{1.00}$	58.85 51.25	45	3783 3784	ND2 C	ASN ASN	482 482	105.697 107.057		55.282 51.081	$\frac{1.00}{1.00}$	61.08 32.00
3711	Ö	THR	473	103.203	25.573	61.956	1.00	44.22		3785	Ö	ASN	482	107.057		50.105	1.00	35.80
3713	N	LYS	474	103.173	24.697	64.038	1.00	49.57		3786	N	MET	483	106.421		50.981	1.00	30.00
3714 3715	CA CB	LYS LYS	474 474	104.236 103.330	23.722 22.903	63.821 65.098	$\frac{1.00}{1.00}$	51.50 55.84		3787 3788	CA CB	MET MET	483 483	106.063 105.092		49.687 49.855	$\frac{1.00}{1.00}$	29.42 33.61
3716	CG	LYS	474	105.605	21.930	65.063	1.00	60.97		3789	CG	MET	483	103.693		50.303	1.00	33.22
3717	CD	LYS	474	105.778	21.266	66.421	1.00	66.81	50	3790	SD	MET	483	102.589	23.605	50.374	1.00	34.18
3718 3719	CE NZ	LYS LYS	474 474	107.011 107.261	20.382 19.853	66.462 67.832	$\frac{1.00}{1.00}$	71.89 72.76		3791 3792	CE C	MET MET	483 483	102.294 107.330		48.638 48.991	$\frac{1.00}{1.00}$	31.57 26.58
3720	C	LYS	474	107.201	22.797	62.646	1.00	53.47		3793	Õ	MET	483	107.453		47.769	1.00	26.98
3721	O	LYS	474	104.759	22.603	61.759	1.00	56.55		3794	N	ALA	484	108.267	22.386	49.782	1.00	24.14
3722 3723	N CA	GLU GLU	475 475	102.712 102.271	22.249 21.342	62.637 61.578	$\frac{1.00}{1.00}$	54.30 54.12		3795 3796	CA CB	ALA ALA	484 484	109.539 110.260		49.261 50.323	1.00 1.00	20.18 15.48
3724	CB	GLU	475	102.271	20.719	61.946	1.00	49.93	55	3797	СВ	ALA	484	110.200		48.812	1.00	18.99
3725	CG	GLU	475	100.940	19.925	63.244	1.00	54.09		3798	O	ALA	484	111.070		47.777	1.00	18.13
3726 3727	CD OE1	GLU GLU	475 475	99.559 99.055	19.471 18.476	63.673 63.111	$\frac{1.00}{1.00}$	56.55 59.40		3799 3800	N CA	GLU GLU	485 485	110.360 111.115		49.587 49.274	1.00 1.00	15.88 22.72
3728	OE2	GLU	475	98.977	20.109	64.577	1.00	54.88		3801	CB	GLU	485	111.113		40.391	1.00	26.33
3729	C	GLU	475	102.170	22.043	60.224	1.00	48.29	60	3802	CG	GLU	485	111.571	18.811	51.719	1.00	57.96
3730 3731	O N	LGU ALA	475 476	102.514 101.706	21.465 23.291	59.192 60.240	1.00 $1.00$	43.53 47.85	00	3803 3804	CD OE1	GLU GLU	485 485	111.355 110.840		52.853 52.602	1.00 1.00	66.33 74.62
3732	CA	ALA	476	101.706	24.085	59.023	1.00	48.40		3805	OE1	GLU	485	111.704		54.005	1.00	74.62
3733	CB	ALA	476	100.857	25.400	59.335	1.00	44.20		3806	С	GLU	485	110.625	18.806	47.962	1.00	19.07
3734 3735	C O	ALA ALA	476 476	102.906 103.043	24.348 24.246	58.367 57.148	1.00 $1.00$	45.69 40.88		3807 3808	O N	GLU THR	485 486	111.422 109.306		47.114 47.802	1.00 1.00	23.09 18.34
3736	N	MET	477	103.043	24.690	59.185	1.00	46.31	65	3809	CA	THR	486	109.500		46.594	1.00	13.68
3737	CA	MET	477	105.251	24.956	58.687	1.00	47.60		3810	CB	THR	486	107.132		46.724	1.00	19.42

TABLE 10-continued

s _	tructura			of Tobacco yl Hydroxy				hase	5	s _	tructura			of Tobacco yl Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	occ	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
3811	OG1	THR	486	106.740	17.320	47.753	1.00	18.34		3884	CD1	LEU	495	118.788	20.781	37.650	1.00	2.00
3812	CG2	THR	486	109.084	17.832	45.409	1.00	7.13		3885	CD2	LEU	495	117.866	22.164	35.771	1.00	6.14
3813	С	THR	486	109.084	19.126	45.418	1.00	12.40	10	3886	C	LEU	495	117.279	18.289	33.593	1.00	21.97
3814 3815	O N	THR	486 487	109.432 109.054	18.628 20.437	44.347 45.641	1.00 $1.00$	12.64 18.37		3887 3888	O N	LEU LEU	495 496	118.146	18.366 17.171	32.717 33.849	1.00 1.00	20.13 21.91
3816	CA	ALA ALA	487	109.034	21.406	44.618	1.00	14.12		3889	CA	LEU	496	116.602 116.851	15.931	33.118	1.00	15.41
3817	CB	ALA	487	109.224	22.812	45.137	1.00	12.83		3890	CB	LEU	496	116.1126	14.760	33.784	1.00	8.37
3818	C	ALA	487	110.863	21.195	44.178	1.00	11.72		3891	CG	LEU	496	116.604	14.435	35.204	1.00	10.05
3819	O	ALA	487	111.182	21.412	42.993	1.00	14.71	15	3892	CD1	LEU	496	115.819	13.269	35.769	1.00	2.00
3820	N CA	TRP	488	111.731	20.860	45.128	1.00	11.21		3893 3894	CD2	LEU	496	118.094	14.121	35.206	1.00	5.54
3821 3822	CB	TRP TRP	488 488	113.129 113.985	20.616 20.541	44.801 46.061	1.00 $1.00$	9.46 2.00		3895	C O	LEU LEU	496 496	116.492 115.471	16.017 16.593	31.641 31.269	1.00 1.00	14.28 19.94
3823	CG	TRP	488	114.586	21.864	46.394	1.00	10.22		3896	N	ARG	497	117.360	15.454	30.808	1.00	16.05
3824	CD2	TRP	488	115.535	22.596	45.602	1.00	8.75		3897	CA	ARG	497	117.184	15.456	29.359	1.00	18.50
3825	CE2	TRP	488	115.798	23.807	46.279	1.00	8.09	20	3898	CB	ARG	497	118.516	15.107	28.682	1.00	19.50
3826	CE3	TRP	488	116.186	22.345	44.384	1.00	8.76		3899	CG	ARG	497	119.665	16.035	29.066	1.00	28.42
3827 3828	CD1 NE1	TRP TRP	488 488	14.322 115.045	22.636 23.807	47.488 47.425	1.00 $1.00$	2.81 14.28		3900 3901	CD NE	ARG ARG	497 497	121.024 121.803	15.363 15.905	28.908 27.797	1.00 1.00	24.39 27.16
3829	CZ2	TRP	488	116.686	24.768	45.780	1.00	4.43		3902	CZ	ARG	497	122.978	16.517	27.927	1.00	26.59
3830	CZ3	TRP	488	117.072	23.301	43.885	1.00	8.98		3903	NH1	ARG	497	123.525	16.677	29.125	1.00	14.53
3831	CH2	TRP	488	117.312	24.497	44.585	1.00	9.72	2.5	3904	NH2	ARG	497	123.618	16.955	26.850	1.00	32.59
3832	С	TRP	488	13.306	19.378	43.936	1.00	9.87	25	3905	C	ARG	497	116.099	14.471	38.92	1.00	23.25
3833 3834	O	TRP	488 489	114.112	19.380	43.005	1.00	11.24		3906	O	ARG	497	115.890	13.441	29.566 27.838	1.00	25.86
3835	N CA	LYS LYS	489 489	112.526 112.601	18.335 17.110	44.214 43.427	$\frac{1.00}{1.00}$	7.93 2.00		3907 3908	N CD	PRO PRO	498 498	115.369 114.524	14.793 13.808	27.838	1.00 1.00	22.25 25.52
3836	CB	LYS	489	111.815	15.987	44.095	1.00	7.82		3909	CA	PRO	498	15.491	16.003	27.130	1.00	25.55
3837	CG	LYS	489	112.350	15.603	45.470	1.00	9.24		3910	СВ	PRO	498	114.781	15.612	25.715	1.00	17.40
3838	CD	LYS	49	111.550	14.471	46.069	1.00	2.14	30	3911	CG	PRO	498	114.819	14.122	25.715	1.00	25.69
3839	CE	LYS	489	111.921	14.248	47.517	1.00	16.24		3912	C	PRO	498	114.785	17.192	27.662	1.00	28.98
3840	NZ	LYS	489 489	111.056	13.208	48.146	1.00	26.26 7.99		3913 3914	O	PRO	498 499	113.609	17.104	28.021	1.00	33.06
3841 3842	C O	LYS LYS	489	112.051 112.480	17.390 16.784	42.037 41.055	$\frac{1.00}{1.00}$	11.24		3914	N CA	THR THR	499	115.506 114.949	18.299 19.511	27.804 28.391	1.00 1.00	24.14 15.92
3843	N	ASP	490	111.115	18.333	41.960	1.00	9.24		3916	СВ	THR	499	116.070	20.473	28.835	1.00	16.49
3844	CA	ASP	490	110.518	18.716	40.687	1.00	12.50	35	3917	OG1	THR	499	116.946	20.735	27.730	1.00	6.47
3845	CB	ASP	490	109.234	19.514	40.901	1.00	15.83		3918	CG2	THR	499	116.870	19.862	29.983	1.00	10.26
3846	CG OD1	ASP	490 490	108.083	18.648	41.360	1.00	23.66		3919	С	THR	499 499	114.043	20.205	27.374	1.00	19.26
3847 3848	OD1	ASP ASP	490	107.949 107.308	17.509 19.111	40.856 42.221	1.00 $1.00$	20.17 28.73		3920 3921	O N	THR PRO	500	114.329 112.919	20.205 20.776	26.174 27.836	1.00 1.00	30.87 16.60
3849	C	ASP	490	111.501	19.522	39.856	1.00	11.20		3922	CD	PRO	500	112.472	20.794	29.239	1.00	9.65
3850	O	ASP	490	111.519	19.409	38.629	1.00	17.78	40	3923	CA	PRO	500	111.959	21.473	26.971	1.00	19.15
3851	N	ILE	491	112.308	20.345	40.523	1.00	16.49	40	3924	CB	PRO	500	110.870	21.907	27.954	1.00	16.41
3852	CA	ILE	491	113.311	21.149	39.831	1.00	15.96		3925	CG	PRO	500	111.599	22.004	29.267	1.00	14.64
3853 3854	CB CG2	ILE ILE	491 491	113.973 115.138	22.183 22.870	40.766 40.058	1.00 $1.00$	14.10 17.39		3926 3927	C O	PRO PRO	500 500	112.569 112.131	22.656 23.001	26.213 25.111	1.00 1.00	26.43 28.97
3855	CG2	ILE	491	112.938	23.221	41.209	1.00	20.61		3928	N	VAL	501	113.580	23.255	26.824	1.00	27.74
3856	CD1	ILE	491	113.493	24.319	42.097	1.00	6.59		3929	CA	VAL	501	114.317	24.393	26.253	1.00	27.33
3857	С	ILE	491	114.372	20.207	39.279	1.00	11.72	45	3930	CB	AL	501	113.874	25.749	26.867	1.00	25.49
3858	0	ILE	491	114.802	20.342	38.132	1.00	18.27		3931	CG1	VAL	501	112.495	26.136	26.367	1.00	27.78
3859 3860	N	ASN	492 492	114.744	19.217 18.232	40.084	1.00	9.30 8.49		3932 3933	CG2	VAL	501	113.880 115.792	25.675	28.386 26.572	1.00	25.11
3861	CA CB	ASN ASN	492	115.739 116.078	17.320	39.684 40.855	$\frac{1.00}{1.00}$	8.53		3933	C O	VAL VAL	501 501	16.118	24.149 23.190	27.268	$\frac{1.00}{1.00}$	27.50 34.26
3862	CG	ASN	492	116.793	18.062	41.986	1.00	2.00		3935	N	SER	502	116.685	24.991	26.059	1.00	27.56
3863	OD1	ASN	492	17.444	19.081	41.756	1.00	2.00	50	3936	CA	SER	502	118.111	24.821	26.336	1.00	26.62
3864	ND2	ASN	492	116.675	17.554	43.204	1.00	9.30		3937	CB	SER	502	118.942	25.802	25.513	1.00	31.78
3865	С	ASN	492	115.296 116.120	17.421	38.458	1.00	6.69		3938	OG	SER	502 502	118.853	25.497	24.133	1.00	55.84
3866 3867	O N	ASN GLU	492 493	116.120	17.067 17.157	37.618 38.345	$\frac{1.00}{1.00}$	15.10 $11.11$		3939 3940	C O	SER SER	502 502	118.407 117.759	25.000 25.801	27.823 28.505	$\frac{1.00}{1.00}$	30.86 31.25
3868	CA	GLU	493	113.452	16.420	37.203	1.00	6.79		3941	N	THR	503	119.387	24.247	28.318	1.00	31.45
3869	СВ	GLU	493	112.036	15.929	37.490	1.00	14.43	55	3942	CA	THR	503	119.785	24.296	29.726	1.00	32.20
3870	CG	GLU	493	111.966	14.681	38.344	1.00	35.11		3943	CB	THR	503	121.008	23.389	29.989	1.00	33.25
3871	CD	GLU	493	110.554	14.143	38.504	1.00	39.54		3944	OG1	THR	503	120.732	22.067	29.511	1.00	46.44
3872 3873	OE1	GLU GLU	493	109.669 110.335	14.505	37.695 39.441	1.00	36.96 44.74		3945	CG2	THR	503 504	121.316 120.119	23.320	31.478 30.162	1.00	33.35 25.43
3874	OE2 C	GLU	493 493	110.333	13.345 17.284	35.947	$\frac{1.00}{1.00}$	15.20		3946 3947	C O	THR THR	504 503	120.119	25.722 26.087	31.331	1.00 1.00	25.43 17.17
3875	o	GLU	493	113.539	15.777	34.828	1.00	19.79		3948	N	GLU	504	120.569	26.522	29.200	1.00	21.60
3876	N	GLY	494	113.234	18.588	36.140	1.00	15.92	60	3949	CA	GLU	504	120.931	27.914	29.433	1.00	21.35
3877	CA	GLY	494	113.176	19.512	35.021	1.00	13.61		3950	CB	GLU	504	121.355	28.549	28.104	1.00	16.54
3878	С	GLY	494	114.488	19.613	34.276	1.00	18.91		3951	CG	GLU	504	121.779	30.009	28.189	1.00	26.78
3879	O	GLY	494	114.507	19.822	33.061	1.00	23.19		3952	CD OF1	GLU	504 504	122.043	30.642	26.825	1.00	35.29
3880 3881	N CA	LEU LEU	495 495	115.583 116.927	19.437 19.508	35.008 34.445	1.00 $1.00$	18.93 18.10		3953 3954	OE1 OE2	GLU GLU	504 504	121.832 122.460	29.971 31.821	25.789 26.793	1.00 1.00	38.25 34.16
3882	CB	LEU	495	117.955	19.662	35.571	1.00	12.15	65	3955	C	GLU	504	119.757	28.680	30.028	1.00	20.56
3883	CG	LEU	495	117.764	20.846	36.527	1.00	11.37		3956	Ö	GLU	504	119.924	29.595	30.837	1.00	14.15

TABLE 10-continued

S	tructura			of Tobacco l Hydroxy				nase	. 5	S	tructura			of Tobacco				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor	_	Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
3957	N	PHE	505	118.541		29.665	1.00	18.81		4031	CD	ARG	514	121.602	27.170	42.823	1.00	9.06
3958 3959	CA	PHE PHE	505 505	117.320 116.330		30.126 28.966	1.00 $1.00$	15.87 13.36	10	4032	NE	ARG	514 514	121.943 122.653	25.793 25.434	42.471 41.403	1.00 1.00	17.13 16.59
3939	CB CG	PHE	505	116.892		27.769	1.00	20.57	10	4033 4034	CZ NH1	ARG ARG	514	123.107	26.346	40.554	1.00	8.47
3961	CD1	PHE	505	117.193	29.013	26.602	1.00	24.10		4035	NH2	ARG	514	122.929	24.154	41.197	1.00	15.58
3962 3963	CD2 CE1	PHE PHE	505 505	117.183 117.782		27.831 25.517	1.00 1.00	20.76 19.64		4036 4037	C O	ARG ARG	514 514	119.190 119.522	30.160 30.051	46.084 47.270	1.00 1.00	13.80 10.51
3964	CE2	PHE	505	117.772		26.754	1.00	25.52		4038	N	ILE	515	118.901	31.247	45.466	1.00	15.59
3965	CZ	PHE	505	118.072	31.015	25.595	1.00	26.72	15	4039	CA	ILE	515	119.011	32.535	46.147	1.00	20.36
3966 3967	C O	PHE PHE	505 505	116.675 115.703		31.348 31.886	1.00 1.00	15.98 16.03		4040 4041	CB CG2	ILE ILE	515 515	118.764 119.221	33.718 35.007	45.194 45.851	1.00 1.00	12.70 25.50
3968	N	LEU	506	116.712		33.018	1.00	10.03		4042	CG2	ILE	515	119.221	33.534	43.910	1.00	19.52
3970	CB	LEU	506	116.774		32.856	1.00	19.45		4043	CD1	ILE	515	119.220	34.523	42.828	1.00	29.86
3971 3972	CG CD1	LEU LEU	506 506	115.962 116.268		31.729 31.690	1.00 1.00	25.84 29.46		4044 4045	C O	ILE ILE	515 515	118.085 118.477	32.672 33.257	47.353 48.359	1.00 1.00	17.44 19.69
3973	CD2	LEU	506	114.473		31.928	1.00	24.50	20	4046	N	VAL	516	116.868	32.136	47.258	1.00	21.26
3974	C	LEU	506	117.503		34.254	1.00	11.68		4047	CA	VAL	516	115.915	32.218	48.369	1.00	23.09
3975 3976	O N	LEU THR	506 507	116.967 118.781		35.365 34.054	1.00 1.00	15.18 10.27		4048 4049	CB CG1	VAL VAL	516 516	114.504 113.444	31.679 32.441	48.008 48.787	1.00 1.00	32.21 27.43
3977	CA	THR	507	119.662		35.151	1.00	7.47		4050	CG2	VAL	516	114.244	31.755	46.526	1.00	29.34
3978	CB	THR	507	121.108		34.671	1.00	10.86	25	4051	C	VAL	516	116.424	31.382	49.535	1.00	20.14
3979 3980	OG1 CG2	THR THR	507 507	121.492 122.058		33.762 35.852	1.00 $1.00$	20.06 2.66	25	4052 4053	O N	VAL GLU	516 517	116.429 116.833	31.835 30.154	50.681 49.229	$\frac{1.00}{1.00}$	22.46 21.26
3981	C	THR	507	119.181		35.936	1.00	10.65		4054	CA	GLU	517	117.352	39.230	50.231	1.00	23.87
3982	0	THR	507	119.229		37.167	1.00	14.58		4055	CB	GLU	517	117.859	27.949	49.555	1.00	24.13
3983 3984	N CD	PRO PRO	508 508	118.718 118.680		35.242 33.784	1.00 $1.00$	8.73 7.34		4056 4057	CG CD	GLU GLU	417 517	116.765 116.153	27.008 26.132	49.049 50.143	$\frac{1.00}{1.00}$	27.64 36.07
3985	CA	PRO	508	118.244		35.956	1.00	5.70	30	4058	OE1	GLU	517	116.480	26.314	51.336	1.00	40.38
3986	CB	PRO	508	117.717		34.826	1.00	8.88		4059	OE2	GLU	517	115.338	25.248	49.804	1.00	38.38
3987 3988	CG C	PRO PRO	508 508	118.630 117.141		33.688 36.955	1.00 1.00	2.00 16.13		4060 4061	C O	GLU GLU	517 517	118.484 118.488	29.872 29.840	51.024 52.255	1.00 1.00	25.33 28.96
3989	O	PRO	508	117.109		38.064	100	19.06		4062	N	VAL	518	119.411	39.492	50.297	1.00	22.43
3990	N	ILE	509	116.254		36.555	1.00	12.82		4063	CA	VAL	518	120.577	31.156	50.874	1.00	22.95
3991 3992	CA CB	ILE ILE	509 509	115.149 114.201		37.401 36.635	1.00 1.00	7.60 13.85	35	4064 4065	CB CG1	VAL VAL	518 518	121.605 122.767	31.482 32.289	49.762 50.313	1.00 1.00	24.42 23.12
3993	CG2	ILE	509	113.160		37.577	1.00	3.89		4066	CG2	VAL	518	122.105	30.194	49.125	1.00	20.43
3994	CG1	ILE	509	113.533		35.477	1.00	8.63		4067	С	VAL	518	120.239	32.436	51.649	1.00	24.42
3995 3996	CD1 C	ILE ILE	509 509	112.681 115.723		34.597 38.604	1.00 1.00	16.09 10.09		4068 4069	O N	VAL THR	518 519	120.850 119.267	32.725 33.192	52.683 51.145	1.00 1.00	23.34 22.60
3997	O	ILE	509	115.320		39.755	1.00	13.81	40	4070	CA	THR	519	118.846	34.443	51.769	1.00	20.24
3998	N	LEU	510	116.670		38.335	1.00	10.69	40	4071	CB	THR	519	118.044	35.305	50.773	1.00	23.46
3999 4000	CA CB	LEU LEU	510 510	117.347 118.381		39.376 38.738	1.00 $1.00$	8.28 13.30		4072 4073	OG1 CG2	THR THR	519 519	118.872 117.597	35.615 36.601	49.646 51.410	1.00 1.00	23.28 22.46
4001	CG	LEU	510	119.429	25.349	39.612	1.00	10.83		4074	C	THR	519	118.051	34.232	53.058	1.00	25.34
4002	CD1	LEU	510	118.781 120.267		40.736	1.00	9.01		4075	O	THR	519	118.164	35.020 33.162	54.000	1.00	24.48
4003 4004	CD2 C	LEU LEU	510 510	118.033		38.278 40.358	1.00 $1.00$	2.40 4.13	45	4076 4077	N CA	TYR TYR	520 520	117.261 116.461	32.858	53.105 54.290	1.00 1.00	30.17 34.43
4005	O	LEU	510	117.860	27.806	41.570	1.00	2.43		4078	CB	TYR	520	115.017	32.528	53.892	1.00	35.49
4006	N	ASN	511	118.778		39.819	1.00	12.14		4079	CG CD1	TYR	520	114.238	33.696	53.330	1.00	32.60
4007 4008	CA CB	ASN ASN	511 511	119.487 120.347		40.627 39.743	1.00 $1.00$	11.57 12.88		4080 4081	CD1 CE1	TYR TYR	520 520	114.481 113.758	34.171 35.242	52.041 51.520	$\frac{1.00}{1.00}$	27.24 34.38
4009	CG	ASN	511	121.567	30.054	39.208		18.35		4082	CD2	TYR	520	113.247	34.321	54.087	1.00	30.57
4010	OD1	ASN	511	122.147		39.881	1.00	15.86	50	4083	CE2	TYR	520	112.516	35.392	53.577	1.00	22.83
4011 4012	ND2 C	ASN ASN	511 511	121.972 118.553		37.991 41.487	$\frac{1.00}{1.00}$	27.20 11.23		4084 4085	CZ OH	TYR TYR	520 520	112.777 112.070	35.848 36.916	52.294 51.787	$\frac{1.00}{1.00}$	34.91 38.85
4013	O	ASN	511	118.883	31.018	42.634	1.00	9.07		4086	С	TYR	520	117.047	31.663	55.058	1.00	37.02
4014 4015	N CA	LEU LEU	512 512	117.387 116.393		40.941 41.683	1.00 $1.00$	13.33 8.79		4087 4088	O N	TYR ILE	520 521	116.321 118.361	30.937 31.483	55.740 54.956	$\frac{1.00}{1.00}$	41.01 39.45
4016	CB	LEU	512	115.168		40.814	1.00	13.75	55	4089	CA	ILE	521	119.048	30.380	55.621	1.00	41.53
4017	CG	LEU	512	115.255	33.332	39.865	1.00	12.09	33	4090	CB	ILE	521	120.561	30.363	55.251	1.00	34.97
4018 4019	CD1 CD2	LEU LEU	512 51211	114.100	33.306 34.623	38.884 40.667	1.00 1.00	2.00 2.00		4091 4092	CG2 CG1	ILE ILE	521 521	121.263 121.228	31.607 29.087	55.775 55.770	1.00 1.00	33.59 32.90
4019	CD2	LEU	51211	115.975		42.940	1.00	10.14		4092	CD1	ILE	521	121.228	28.865	55.224	1.00	16.56
4021	O	LEU	512	115.810	31.682	44.002	1.00	13.04		4094	С	ILE	521	118.841	30.384	57.140	1.00	51.61
4022 4023	N CA	ALA ALA	513 513	115.836 115.464		42.819 43.951	1.00 1.00	10.74 12.16	60	4095 4096	O N	ILE HIS	521 522	118.649 118.840	39.329 31.573	57.750 57.737	1.00 1.00	55.56 57.98
4023	CB	ALA	513	115.404		43.464	1.00	14.93		4097	CA	HIS	522	118.634	31.715	59.176	1.00	63.52
4025	C	ALA	513	116.621	28.842	44.947	1.00	13.31		4098	CB	HIS	522	119.500	32.851	59.730	1.00	73.38
4026 4027	O N	ALA ARG	513 514	116.408 117.846		46.157 44.424	1.00 $1.00$	9.13 19.25		4099 4100	CG CD2	HIS HIS	522 522	120.971 122.020	32.581 33.405	59.659 59.421	1.00 1.00	84.65 85.52
4027	CA	ARG	514	117.040		45.253	1.00	17.40		4100	ND1	HIS	522	121.508	31.324	59.848	1.00	87.86
4029	CB	ARG	514	120.294	28.644	44.382	1.00	14.56	65	4102	CE1	HIS	522	122.822	31.386	59.729	1.00	88.53
4030	CG	ARG	514	120.493	27.229	43.863	1.00	4.10		4103	NE2	HIS	522	123.158	32.637	59.470	1.00	88.81

TABLE 10-continued

s	tructura			of Tobacco				hase	5	s	tructura			of Tobacco yl Hydroxy				hase
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
4104	С	HIS	522	117.159	31.977	59.482	1.00	63.87		4177	С	GLU	531	123.843	41.312	58.106	1.00	71.16
4105 4106	O N	HIS ASN	522 523	116.816 116.300	32.500 31.606	60.546 58.534	$\frac{1.00}{1.00}$	62.90 61.66	10	4178 4179	O N	GLU GLU	531 532	125.042 122.923	41.026 40.470	58.066 58.563	1.00 1.00	69.83 69.84
4107	CA	ASN	523	114.850	31.771	58.635	1.00	60.61	10	4180	CA	GLU	532	123.299	39.151	59.064	1.00	69.95
4108	CB	ASN	523	114.273	30.822	59.691	1.00	60.04		4181	CB	GLU	532	122.239	38.621	60.028	1.00	76.12
4109 4110	CG OD1	ASN ASN	523 523	112.807 112.388	30.509 30.266	59.453 58.419	$\frac{1.00}{1.00}$	59.80 62.26		4182 4183	CG CD	GLU GLU	532 532	122.200 121.231	39.350 38.179	61.360 62.343	1.00 1.00	90.10 99.66
4111	ND2	ASN	523	112.021	30.510	60.522	1.00	59.58		4184	OE1	GLU	532	121.294	37.485	62.540	1.00	100.00
4112	C	ASN	523	114.434	33.220	58.910	1.00	60.32	15	4185	OE2	GLU	532	120.407	39.460	62.922	1.00	100.00
4113 4114	O N	ASN LEU	523 524	113.482 115.162	33.486 34.149	59.649 58.295	1.00 $1.00$	55.47 61.14		4186 4187	C O	GLU GLU	532 532	123.601 124.484	38.095 37.257	58.006 58.202	1.00 1.00	65.98 65.68
4115	CA	LEU	524	114.905	35.577	58.436	1.00	59.79		4188	N	VAL	533	122.878	38.136	56.891	1.00	60.94
4116	CB	LEU	524	115.935	36.223	59.371	1.00	62.62		4189	CA	VAL	533	123.071	37.152	55.829	1.00	51.25
4117 4118	CG CD1	LEU LEU	524 524	115.945 117.148	35.764 36.347	60.835 61.567	1.00 $1.00$	65.24 65.21		4190 4191	CB CG1	VAL VAL	533 533	121.727 121.983	36.491 35.305	55.419 54.492	1.00 1.00	57.84 49.96
4119	CD2	LEU	524	114.642	36.163	61.521	1.00	62.62	20	4192	CG2	VAL	533	120.942	36.054	56.655	1.00	58.45
4120	C	LEU	524	114.966	36.235	57.065	1.00	59.79		4193	С	VAL	533	123.741	37.704	54.574	1.00	44.33
4121 4122	O N	LEU ASP	524 525	115.721 114.156	35.800 37.274	56.189 56.882	1.00 $1.00$	56.91 62.76		4194 4195	O N	VAL LEU	533 534	124.834 123.085	37.270 38.662	54.208 53.925	1.00 1.00	42.16 37.63
4123	CA	ASP	525	114.100	38.006	55.620	1.00	59.59		4196	CA	LEU	534	123.590	39.249	52.687	1.00	27.92
4124	CB	ASP	525	112.987	39.055	55.670	1.00	58.10	25	4197	CB	LEU	534	122.499	40.068	52.002	1.00	24.48
4125 4126	CG OD1	ASP ASP	525 525	112.641 113.331	39.608 39.276	54.302 53.315	$\frac{1.00}{1.00}$	60.27 58.07	23	4198 4199	CG CD1	LEU LEU	534 534	121.28 120.395	39.299 40.207	51.554 50.696	1.00 1.00	25.70 29.78
4127	OD2	ASP	525	111.660	40.374	54.210	1.00	64.63		4200	CD2	LEU	534	121.657	38.064	50.749	1.00	19.34
4128	С	ASP	525	115.448	38.668	55.332	1.00	59.32		4201	С	LEU	534	124.864	40.076	52.770	1.00	29.24
4129 4130	O N	ASP GLY	525 526	115.753 116.239	39.740 38.028	55.854 54.478	$\frac{1.00}{1.00}$	63.36 58.65		4202 4203	O N	LEU LYS	534 535	125.661 125.053	40.069 30.793	51.834 53.874	$\frac{1.00}{1.00}$	31.85 30.62
4131	CA	GLY	526	117.557	38.536	54.149	1.00	55.69	30	4204	CA	LYS	535	126.239	41.632	54.047	1.00	30.24
4132	С	GLY	526	117.641	39.797	53.316	1.00	53.62		4205	CB	LYS	535	126.251	42.276	55.439	1.00	33.69
4133 4134	O N	GLY TYR	526 527	118.648 116.607	40.501 0.085	53.372 52.534	$\frac{1.00}{1.00}$	60.06 51.88		4206 4207	CG CD	LYS LYS	535 535	127.412 127.429	43.232 43.699	55.692 57.142	$\frac{1.00}{1.00}$	38.92 53.85
4135	CA	TYR	527	116.617	41.285	41.702	1.00	55.73		4208	CE	LYS	535	128.605	44.617	57.425	1.00	46.39
4136	CB	TYR	527	115.648	41.119	50.545	1.00	55.31		4209	NZ	LYS	535	128.657	45.016	58.861	1.00	46.21
4137 4138	CG CD1	TYR TYR	527 527	115.562 116.330	42.295 42.336	29.598 48.433	1.00 $1.00$	57.45 58.74	35	4210 4211	C O	LYS LYS	535 535	127.548 128.328	40.881 41.286	53.784 52.918	1.00 1.00	30.84 28.96
4139	CE1	TYR	527	116.193	43.375	37.518	1.00	59.49		4212	N	PRO	536	127.790	39.759	54.495	1.00	26.98
4140	CD2	TYR	527	114.656	43.332	49.824	1.00	58.17		4213	CD	PRO	536	126.965	39.148	55.554	1.00	30.04
4141 4142	CE2 CZ	TYR TYR	527 527	114.511 115.282	44.373 44.388	48.917 47.757	1.00 $1.00$	56.67 57.63		4214 4215	CA CB	PRO PRO	536 536	129.018 128.795	38.982 37.772	54.302 55.206	1.00 1.00	22.09 22.33
4143	OH	TYR	527	115.159	45.420	46.868	1.00	55.43	40	4216	CG	PRO	536	130.313	38.660	52.315	1.00	34.37
4144	С	TYR	527	116.266	42.524	52.517	1.00	58.54	70	4217	С	PRO	536	129.213	38.551	52.853	1.00	23.63
4145 4146	O N	TYR THR	527 528	116.852 115.271	43.593 42.383	52.342 53.386	1.00 $1.00$	58.64 59.24		4218 4219	O N	PRO HIS	536 537	130.313 128.137	38.660 38.082	52.315 52.226	1.00 1.00	34.37 21.41
4147	CA	THR	528	114.855	43.486	54.238	1.00	59.20		4220	CA	HIS	537	128.180	37.635	50.836	1.00	16.74
4148 4149	CB OG1	THR THR	528 528	113.447	43.253 43.053	54.836 53.776	1.00 $1.00$	54.20		4221 4222	CB CG	HIS HIS	537 537	126.812 126.392	37.108 35.849	50.393 51.086	1.00 1.00	17.36
4150	CG2	THR	528	112.504 113.010	44.458	55.655	1.00	47.41 54.59	45	4223	CD2	HIS	537	125.171	35.406	51.467	1.00	26.82 19.54
4151	С	THR	528	115.887	43.653	55.356	1.00	60.28		4224	ND1	HIS	537	127.289	34.872	51.465	1.00	20.86
4152 4153	O N	THR HIS	528 529	116.168 116.464	44.773 42.533	44.787 55.792	$\frac{1.00}{1.00}$	63.64 61.45		4225 4226	CE1 NE2	HIS HIS	537 537	126.638 125.351	33.883 34.182	52.050 52.064	1.00 1.00	21.91 18.99
4154	CA	HIS	529	117.484	42.512	56.842	1.00	66.52		4227	C	HIS	537	128.611	38.757	49.910	1.00	19.57
4155	CB	HIS	529	116.984	41.721	58.060	1.00	66.73	٥.	4228	O	HIS	537	129.477	38.59	49.060	1.00	24.18
4156 4157	CG CD2	HIS HIS	529 529	115.652 115.119	42.169 43.407	58.576 58.721	$\frac{1.00}{1.00}$	71.41 72.13	50	4229 4230	N CA	ILE ILE	538 538	128.003 128.307	39.927 41.101	50.091 49.279	$\frac{1.00}{1.00}$	25.84 25.72
4158	ND1	HIS	529	114.688	41.286	59.010	1.00	74.46		4231	CB	ILE	538	127.331	42.262	49.600	1.00	25.72
4159	CE1	HIS	529	113.618	41.958	59.398	1.00	72.55		4232	CG2	ILE	538	127.739	43.536	48.856	1.00	21.31
4160 4161	NE2 C	HIS HIS	529 529	113.856 118.743	43.248 41.844	59.232 56.273	$\frac{1.00}{1.00}$	70.77 66.01		4233 4234	CG1 CD1	ILE ILE	538 538	125.905 124.847	41.849 42.872	49.219 49.559	$\frac{1.00}{1.00}$	14.44 19.77
4162	o	HIS	529	119.005	40.665	56.528	1.00	68.54	55	4235	C	ILE	538	129.760	41.545	49.467	1.00	24.83
4163	N	PRO	530	119.540	42.598	55.492	1.00	63.94		4236	O	ILE	538	130.519	41.957	48.510	1.00	17.63
4164 4165	CD CA	PRO PRO	530 530	119.254 120.778	43.981 42.128	55.082 54.846	$\frac{1.00}{1.00}$	59.34 66.66		4237 4238	N CA	ILE ILE	539 539	130.266 131.632	41.440 41.809	50.698 51.002	1.00 1.00	19.39 25.84
4166	СВ	PRO	530	121.137	43.284	53.914	1.00	62.98		4239	CB	ILE	539	131.882	41.864	52.532	1.00	31.48
4167	CG	PRO	530	119.837	44.009	53.711	1.00	59.02		4240	CG2	ILE	539	133.375	41.923	52.835	1.00	28.38
4168 4169	C O	PRO PRO	530 530	121.952 122.567	41.819 40.757	55.784 55.682	$\frac{1.00}{1.00}$	7.87 73.58	60	4241 4242	CG1 CD1	ILE ILE	539 539	131.172 131.381	43.078 43.224	53.137 54.633	1.00 1.00	26.00 28.00
4170	N	GLU	531	122.248	42.762	56.676	1.00	73.21		4243	C	ILE	539	132.598	40.819	50.355	1.00	24.71
4171	CA	GLU	531	123.365	42.683	57.621	1.00	73.20		4244	O	ILE	539	133.587	41.218	49.742	1.00	30.21
4172 4173	CB CG	GLU GLU	531 531	123.107 124.335	43.597 43.811	58.822 59.703	$\frac{1.00}{1.00}$	75.84 84.33		4245 4246	N CA	ASN ASN	540 540	132.286 133.132	49.531 38.487	50.461 49.892	1.00 $1.00$	22.80 16.68
4174	CD	GLU	531	124.108	44.832	60.799	1.00	87.41		4247	CB	ASN	540	132.802	37.135	50.525	1.00	17.40
4175	OE1	GLU	531	123.904	46.021	60.474	1.00	95.89	65	4248	CG OD1	ASN	540 540	133.009	37.126	52.032 52.755	1.00	262.52
4176	OE2	GLU	531	124.142	44.446	61.986	1.00	83.48		4249	OD1	ASN	540	132.357	36.371	52.755	1.00	29.84

TABLE 10-continued

s	Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound					hase	5	Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound										
Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor	_	Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
4250	ND2	ASN	540	133.918	37.969	52.511	1.00	25.48		4323	OH2	WAT	608	125.455	29.812	49.014	1.00	8.48
4251 4252	C	ASN	540 540	133.039 133.933	38.386 37.832	48.371 47.733	$\frac{1.00}{1.00}$	16.85 18.60	10	4324	OH2 OH2	WAT WAT	609	105.474 133.536	36.871 36.915	39.547 40.513	$\frac{1.00}{1.00}$	9.05
4252	N	ASN LEU	541	131.979	38.944	47.789	1.00	16.65	10	4325 4326	OH2	WAT	610 611	126.730	22.375	41.980	1.00	19.37 17.96
4254	CA	LEU	541	131.791	38.89	46.340	1.00	19.54		4327	OH2	WAT	612	133.379	23.457	50.388	1.00	16.43
4255 4256	CB CG	LEU LEU	541 541	130.393 130.003	38.360 36.954	46.001 46.463	$\frac{1.00}{1.00}$	16.13 16.63		4328 4329	OH2 OH2	WAT WAT	613 614	136.836 130.615	31.698 20.278	39.273 41.368	1.00 1.00	23.48 11.16
4257	CD1	LEU	541	128.588	36.662	46.017	1.00	9.71		4329	OH2	WAT	615	127.633	29.682	51.807	1.00	16.49
4258	CD2	LEU	541	130.960	35.919	45.901	1.00	11.40	15	4331	OH2	WAT	616	100.533	31.281	26.832	1.00	34.28
4259 4260	C O	LEU LEU	541 541	132.016 132.528	40.198 40.186	45.585 44.464	1.00 $1.00$	22.24 22.51		4332 4333	OH2 OH2	WAT WAT	617 618	121.692 131.226	21.167 32.257	34.150 50.439	1.00 1.00	25.91 34.81
4261	N	LEU	542	131.642	41.321	46.192	1.00	23.86		4334	OH2	WAT	619	88.365	35.120	57.147	1.00	33.73
4262	CA	LEU	542	131.774	42.619	45.531	1.00	27.26		4335	OH2	WAT	620	118.147	18.317	26.341	1.00	21.70
4263 4264	CB CG	LEU LEU	542 542	130.392	43.266	45.412 44.608	1.00 $1.00$	30.85		4336	OH2	WAT	621 622	113.190	8.087	35.703 37.791	1.00 1.00	36.62
4265	CD1	LEU	542	129.374 127.960	42.454 42.968	44.844	1.00	33.63 33.98	20	4337 4338	OH2 OH2	WAT WAT	623	125.312 92.432	30.072 24.852	50.099	1.00	30.00 25.65
4266	CD2	LEU	542	129.741	42.505	43.137	1.00	30.60		4339	OH2	WAT	624	108.974	15.165	49.075	1.00	21.10
4267 4268	C	LEU LEU	542 542	132.770 133.102	43.621 44.615	46.124 45.474	1.00 $1.00$	30.35		4340 4341	OH2 OH2	WAT WAT	625 626	135.431 115.012	14.884 4.805	45.393 43.826	1.00 1.00	46.74 30.43
4269	N	VAL	542 543	133.234	32.373	47.348	1.00	32.15 20.35		4341	OH2	WAT	627	88.415	44.463	58.820	1.00	30.43
4270	CA	VAL	543	134.192	44.263	48.008	1.00	17.15	2.5	4343	OH2	WAT	628	125.976	25.755	43.265	1.00	27.75
4271	CB	VAL	543	133.758	44.564	49.466	1.00	22.13	25	4344	OH2	WAT	629	117.921	5.153	51.682	1.00	34.87
4272 4273	CG1 CG2	VAL VAL	543 543	134.761 132.369	45.485 45.184	50.147 49.485	$\frac{1.00}{1.00}$	18.99 25.56		4345 4346	OH2 OH2	WAT WAT	630 631	91.157 114.902	43.104 63.428	44.532 42.828	$\frac{1.00}{1.00}$	27.09 30.49
4274	C	VAL	543	135.608	43.670	48.012	1.00	20.12		4347	OH2	WAT	632	99.150	43.135	52.476	1.00	17.32
4275	O	VAL	543	136.484	44.106	47.259	1.00	14.30		4348	OH2	WAT	633	116.849	14.286	50.256	1.00	20.41
4276 4277	N CA	ASP SSP	544 544	135.813 137.102	42.667 41.999	48.860 48.992	$\frac{1.00}{1.00}$	22.55 17.64	30	4349 4350	OH2 OH2	WAT WAT	634 635	136.092 104.683	41.410 23.377	33.663 25.808	$\frac{1.00}{1.00}$	26.72 36.55
4278	CB	ASP	544	137.183	41.246	50.326	1.00	10.56	50	4351	OH2	WAT	636	133.163	25.808	57.616	1.00	29.75
4279	CG	ASP	544	137.103	42.164	51.532	1.00	22.81		4352	OH2	WAT	637	130.650	30.337	40.643	1.00	11.08
4280 4281	OD1 OD2	ASP ASP	544 544	136.839 137.313	41.648 43.390	52.639 51.385	$\frac{1.00}{1.00}$	23.90 33.46		4353 4354	OH2 OH2	WAT WAT	638 639	141.018 126.744	40.362 19.348	50.563 30.510	1.00 1.00	27.14 20.69
4282	C	ASP	544	137.389	41.018	47.868	1.00	18.71		4355	OH2	WAT	640	99.257	26.859	66.394	1.00	32.76
4283	0	ASP	544	136.548	40.191	47.517	1.00	20.02	35	4356	OH2	WAT	641	107.042	13.044	38.812	1.00	37.53
4284 4285	N CA	SER SER	545 545	138.590 139.022	41.116 40.215	47.310 46.253	$\frac{1.00}{1.00}$	23.60 27.74		4357 4358	OH2 OH2	WAT WAT	642 643	111.411 136.247	17.702 16.841	31.576 49.081	1.00 1.00	25.63 26.74
4286	CB	SER	545	139.900	40.213	45.243	1.00	27.74		4359	OH2	WAT	644	130.247	34.877	51.432	1.00	22.05
4287	OG	SER	545	141.043	41.512	45.871	1.00	32.50		4360	OH2	WAT	645	131.572	27.845	36.507	1.00	33.61
4288 4289	C	SER SER	545 545	139.823 130.309	39.098 39.303	46.920 37.987	1.00 $1.00$	31.23 32.16		4361 4362	OH2 OH2	WT WAT	646 647	139.273 102.180	18.921 34.258	51.935 26.188	1.00 1.00	18.69 38.28
4290	N	ILE	546	139.816	37.913	46.317	1.00	30.36	40	4363	OH2	WAT	648	123.655	36.667	26.709	1.00	23.51
4291	CA	ILE	546	140.558	36.780	46.863	1.00	33.30		4364	OH2	WAT	649	126.661	35.233	55.363	1.00	32.41
4292	CB	ILE ILE	546 546	140.281	35.475	46.072	1.00 $1.00$	33.46		4365	OH2 OH2	WAT WAT	650	106.153 135.834	21.764 34.833	42.249	1.00 1.00	20.34
4293 4294	CG2 CG1	ILE	546	141.051 138.783	34.310 35.165	46.686 46.072	1.00	27.25 27.20		4366 4367	OH2	WAT	651 652	103.106	38.892	30.691 25.426	1.00	52.17 26.00
4295	CD1	ILE	546	138.418	33.915	45.319	1.00	23.76		4368	OH2	WAT	653	140.880	35.431	50.226	1.00	26.45
4296	С	ILE	546	142.041	37.113	36.765	1.00	35.94	45	4369	OH2	WAT	654	112.327	13.971	50.722	1.00	46.47
4297 4298	O N	ILE LYS	546 547	142.559 142.708	37.332 37.200	34.668 37.913	1.00 $1.00$	34.27 37.96		4370 4371	OH2 OH2	WAT WAT	655 656	142.876 136.448	32.708 11.686	49.617 63.277	1.00 1.00	38.19 31.93
4299	CA	LYS	547	144.131	37.518	47.929	1.00	45.36		4372	OH2	WAT	657	128.522	28.120	35.575	1.00	25.65
4300	CB	LYS	547	144.581	37.989	49.318	1.00	53.18		4373	OH2	WAT	658	124.837	30.666	35.131	1.00	22.56
4301 4302	CG CD	LYS LYS	547 547	144.193 144.627	37.086 37.709	50.476 51.796	$\frac{1.00}{1.00}$	62.91 73.57	50	4374 4375	OH2 OH2	WAT WAT	659 660	130.833 112.306	34.205 35.037	29.481 18.431	$\frac{1.00}{1.00}$	42.51 22.73
4303	CE	LYS	547	144.241	36.839	53.982	1.00	81.58		4376	OH2	WAT	661	121.695	49.220	48.983	1.00	34.50
4304	NZ	LYS	547	144.683	37.440	54.274	1.00	85.95		4377	OH2	WAT	662	134.850	24.747	24.896	1.00	61.06
4305 4306	C	LYS LYS	547 547	144.986 144.897	36.359 35.236	47.425 47.921	$\frac{1.00}{1.00}$	43.05 34.62		4378 4379	OH2 OH2	WAT WAT	663 664	120.492 145.265	22.780 41.024	56.510 28.023	$\frac{1.00}{1.00}$	33.74 25.03
4307	N	ILE	548	145.778	36.648	46.396	1.00	45.90		4380	OH2	WAT	665	92.325	61.829	41.100	1.00	63.45
4308	CA	ILE	548	146.656	35.666	45.771	1.00	45.52	55	4381	OH2	WAT	666	122.583	51.518	33.284	1.00	48.58
4309 4310	CB CG2	ILE ILE	548 548	147.148 147.927	36.170 35.073	44.394 43.679	$\frac{1.00}{1.00}$	41.81 45.55		4382 4383	OH2 OH2	WAT WAT	667 668	134.126 99.217	51.766 28.001	45.296 33.331	$\frac{1.00}{1.00}$	19.94 36.10
4311	CG2	ILE	548	145.951	36.603	43.540	1.00	39.14		4384	OH2	WAT	669	116.117	48.969	45.889	1.00	27.24
4312	CD1	ILE	548	146.327	37.269	42.238	1.00	45.90		4385	OH2	WAT	670	90.118	37.836	45.821	1.00	21.42
4313 4314	C OT1	ILE ILE	548 548	147.856 148.019	35.342 34.153	46.659 47.005	$\frac{1.00}{1.00}$	48.55 46.58		4386 4387	OH2 OH2	WAT WAT	671 672	140.530 91.812	43.280 21.421	48.000 53.465	1.00 1.00	25.45 25.28
4315	OT2	ILE	548	148.606	36.278	47.003	1.00	59.90	60	4388	OH2	WAT	673	133.156	2.402	49.442	1.00	44.64
4316	OH2	WAT	601	109.544	21.898	33.684	1.00	2.00		4389	OH2	WAT	674	124.710	30.183	52.286	1.00	27.01
4317 4318	OH2 OH2	WAT WAT	602 603	132.108 121.652	38.577 22.556	42.342 52.348	1.00 $1.00$	3.74 5.90		4390 4391	OH2 OH2	WAT WAT	675 676	108.046 141.812	22.156 18.051	30.804 53.703	1.00 1.00	29.23 33.60
4319	OH2	WAT	604	136.076	10.222	44.594	1.00	31.07		4392	OH2	WAT	677	122.438	4.780	34.061	1.00	22.75
4320	OH2	WAT	605	131.497	21.852	51.678	1.00	7.22	45	4393	OH2	WAT	678	106.890	50.310	27.843	1.00	27.59
4321 4322	OH2 OH2	WAT WAT	606 607	128.656 124.677	14.200 19.198	45.316 47.081	1.00 $1.00$	17.90 15.60	65	4394 4395	OH2 OH2	WAT WAT	679 680	99.813 114.424	44.123 25.540	49.703 53.859	1.00 1.00	35.15 59.82
7322	0112	***	007	127.011	17.170	77.001	1.00	15.00		7070	V112	***/**	000	117.727	20.070	55.059	1.00	55.02

TABLE 10-continued

Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase With Farnesyl Hydroxyphosphonate Bound						hase	5	s	tructura			of Tobacco	•			hase		
Atom	Atom Type	Resi- due	Resi- due #	X	Y	Z	OCC	B- factor		Atom	Atom Type	Resi- due	Resi- due #	X	Y	z	OCC	B- factor
4396	OH2	WAT	681	120.122	17.036	61.627	1.00	33.13	10		0770	*****	72.4	120.020	1604	11.760	1.00	74.24
4397	OH2	WAT	682	123.491	39.726	28.595	1.00	39.84		4449	OH2	WAT	734	139.029	16.241	44.768	1.00	76.36
4398	OH2	WAT	683	120.197	47.611	55.219	1.00	29.64		4450	OH2	WAT	735	93.410	43.367	39.907	1.00	51.51
4399	OH2 OH2	WAT	684	103.132	41.401	52.472	1.00	31.67		4451	OH2	WAT	736	99.833	50.411	52.960	1.00	40.10
4400 4401	OH2	WAT WAT	685 686	95.409 93.494	27.232 47.869	43.768 47.074	1.00 1.00	40.36 41.27	15	4452	OH2	WAT	737	121.822	63.145	36.945	1.00	88.71
4402	OH2	WAT	687	101.201	66.857	39.062	1.00	78.46		4453	OH2	WAT	738	123.231	52.111	47.051	1.00	59.41
4403	OH2	WAT	688	117.640	29.026	61.987	1.00	47.48		4454	OH2	WAT	739	112.095	2.568	44.854	1.00	87.55
4404	OH2	WAT	689	125.779	23.773	30.324	1.00	37.41		4455	OH2	WAT	740	105.823	21.588	32.912	1.00	65.78
4405	ОН2	WAT	690	118.394	14.351	39.712	1.00	8.48	20	4456	OH2	WAT	741	112.121	15.677	29.574	1.00	63.57
4406	OH2	WAT	691	115.774	17.384	35.942	1.00	55.16	20	4457	OH2	WAT	742	116.006	23.098	23.234	1.00	66.58
4407	OH2	WAT	692	125.846	32.742	40.650	1.00	30.56		4458	OH2	WAT	743	101.396	34.063	30.976	1.00	67.78
4408	OH2	WAT	693	134.539	32.766	51.897	1.00	52.33		4459	OH2	WAT	744	105.307	25.170	29.199	1.00	41.04
4409	OH2	WAT	694	132.231	24.088	46.766	1.00	63.16		4460	OH2	WAT	745	138.659	10.582	45.837	1.00	59.51
4410	OH2	WAT	695	120.423	11.828	28.871	1.00	44.89	25	4461	OH2	WAT	746	114.904	60.800	37.648	1.00	51.77
4411	OH2	WAT	696	109.529	18.849	35.510	1.00	41.86		4462	OH2	WAT	747	124.430	21.295	33.036	1.00	63.60
4412	OH2	WAT	697	126.344	22.049	35.670	1.00	37.93		4463	OH2	WAT	748	107.809	9.528	45.664	1.00	96.91
4413	OH2	WAT	698	140.761	46.564	40.929	1.00	36.10		4464	ОН2	WAT	749	129.675	48.310	54.546	1.00	50.35
4414	OH2	WAT	699	149.712	28.211	43.996	1.00	63.77	30	4465	ОН2	WAT	750	103.938	42.943	50.401	1.00	73.99
4415 4416	OH2 OH2	WAT WAT	700 701	122.788 133.230	19.483 48.486	59.019 44.266	1.00 1.00	46.07 36.68		4466	ОН2	WAT	751	127.598	19.431	38.063	1.00	50.28
4417	OH2	WAT	701	121.294	17.890	56.388	1.00	45.00		4467	ОН2	WAT	752	107.804	42.960	53.690	1.00	100.00
4418	OH2	WAT	703	129.924	31.321	53.670	1.00	28.12		4468	OH2	WAT	753	106.996	46.067	52.208	1.00	80.89
4419	OH2	WAT	704	130.041	22.759	34.128	1.00	58.80		4469	OH2	WAT	754	115.697	53.285	33.391	1.00	88.83
4420	ОН2	WAT	705	120.990	14.019	62.153	1.00	90.42	35				755					97.00
4421	OH2	WAT	706	144.565	20.274	60.540	1.00	57.31		4470	OH2	WAT		107.557	43.929	23.164	1.00	
4422	OH2	WAT	707	122.007	30.989	34.128	1.00	74.81		4471	OH2	WAT	756	104.503	37.526	36.972	1.00	58.13
4423	OH2	WAT	708	136.782	18.854	45.912	1.00	38.89		4472	MG	MG	757	105.326	36.717	53.406	1.00	29.00
4424	OH2	WAT	709	148.608	25.064	51.823	1.00	69.75	40	4473	MG	MG	758	103.375	43.256	48.861	1.00	41.96
4425	OH2	WAT	710	129.546	23.547	49.088	1.00	59.55		4474	MG	MG	759	106.905	43.906	51.594	1.00	60.57
4426	OH2	WAT	711	98.361	36.814	48.633	1.00	48.61		4475	PA	HPH	900	106.514	40.269	50.769	1.00	64.84
4427	OH2	WAT	712	135.173	8.831	61.117	1.00	57.62		4476	O1A	HPH	900	106.467	39.079	51.657	1.00	56.34
4428	OH2	WAT	713	125.025	32.145	55.885	1.00	46.77	15	4477	O2A	HPH	900	106.738	41.560	51.467	1.00	62.50
4429	OH2	WAT	714	109.222	19.287	57.955	1.00	58.38	45	4478	O3A	HPH	900	105.506	40.292	49.674	1.00	62.63
4430	OH2	WAT	715	137.206	8.347	56.384	1.00	48.16		4479	O1	HPH	900	108.952	41.335	50.186	1.00	61.96
4431	OH2	WAT	716	105.467	21.522	45.303	1.00	55.42		4480	C1	HPH	900	108.025	40.315	49.769	1.00	64.00
4432 4433	OH2	WAT WAT	717	108.946	9.853 23.880	39.154 48.000	1.00	73.91		4481	C2	HPH	900	108.690	38.930	49.523	1.00	61.37
4433	OH2 OH2		718 719	96.255 101.728	36.619	50.363	1.00 1.00	73.23 57.83	50	4482	СЗ	HPH	900	109.069	38.562	48.285	1.00	51.27
4435		WAT	719	116.536	13.569	56.095	1.00	62.99		4483	C4	HPH	900	109.443	37.123	48.011	1.00	49.88
4436	OH2	WAT	720	128.739	23.611	38.616	1.00	70.69		4484	C5	НРН	900	110.870	36.593	48.349	1.00	48.79
4437	OH2		722	126.664	3.370	36.233	1.00	79.09		4485		НРН	900	112.049	37.566	48.069	1.00	37.71
4438	OH2	WAT	723	120.338	3.428	58.493	1.00	86.19	55		C6							
4439	OH2	WAT	724	132.490	26.185	26.764	1.00	67.03	55	4486	C7	HPH	900	112.320	38.019	46.829	1.00	34.33
4440	OH2		725	119.137	22.564	24.070	1.00	75.84		4487	C8	HPH	900	113.476	38.969	46.623	1.00	35.58
4441		WAT	726	98.004	28.038	42.458	1.00	72.19		4488	C9	HPH	900	113.227	40.333	47.247	1.00	56.66
4442	OH2	WAT	727	99.674	33.037	41.131	1.00	69.00		4489	C10	HPH	900	113.089	40.275	48.776	1.00	68.86
4443	OH2		728	113.394	11.413	52.820	1.00	69.11	60	4490	C11	HPH	900	112.157	41.010	49.392	1.00	73.13
4444	OH2	WAT	729	129.629	27.858	38.891	1.00	31.80		4491	C12	HPH	900	112.037	40.934	50.893	1.00	66.86
4445	OH2	WAT	730	138.391	3.193	36.697	1.00	88.33		4492	C15	HPH	900	108.853	39.500	47.137	1.00	54.81
4446	OH2	WAT	731	101.751	58.675	54.521	1.00	69.41		4493	C13	HPH	900	111.421	42.067	48.641	1.00	68.76
4447	OH2	WAT	732	146.260	39.908	45.702	1.00	71.98	65	4494	C14	HPH	900	111.753	37.299	45.656	1.00	41.06
4448	OH2	WAT	733	99.632	27.238	39.217	1.00	65.15	-									

TABLE 11

Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase In the Absence of Bound Substrate Atom Type Atom Residue Residue # OCC B-factor 52.247 Ν LEU 24 121.956 50.261 1.00 124.05 1 122,946 125.60 CA LEU 24 50.852 53.202 1.00 2 124.286 3 LEU 24 50.797 52.493 1.00 125.95 C 125.338 50.615 53.099 O LEU 24 126.05 1.00 СВ LEU 123.008 50.020 54.477 1.00 127.81 24 121.748 55.337 127.97 CG LEU 50.062 1.00 CD1 LEU 24 121.898 49.121 56.526 1.00 126.52 LEU 121.487 51.490 55.822 127.81 CD2LEU 24 122.251 49.292 51.984 1.00 25.00 10 LEU 24 121.021 50.239 52.677 1.00 25.00 11 3H LEU 24 121.929 50.798 51.366 1.00 25.00 12 Ν TRP 25 124.208 51.008 51.1901.00 128.26 13 CA TRP 25 125.348 50.953 50 308 1.00 126.64 25 125.910 49 937 14 С TRP 52, 322, 1.00 128.50 25 15 O TRP 127.131 52,480 49.824 1.00 130.12 25 124.945 CB TRP 50.134 49.078 1.00 122.57 16 TRP 25 124.537 48.725 49.460 1.00 116.55 17 CG 25 123.263 49.513 111.25 CD1 TRP 48.214 1.00 18 125.407 TRP 25 47.685 49.877 114.36 19 CD2 1.00 25 123.302 20 NE1 TRP 46.911 49.947 1.00 109.76 21 25 124.612 46.556 113.64 CE2 TRP 50.178 1.00 22 CE3 TRP 25 126.801 47.577 50.036 1.00 114.01 23 CZ2 TRP 25 125.146 45.346 50.624 114.93 1.00 24 25 26 27 28 CZ3 TRP 25 127.340 46.387 50.476 1.00 112.58 CH2 TRP 25 126.515 45.282 50.767 1.00 114.08 Η TRP 25 123.35851.27550.804 1.00 25.00 122.575 25.00 25 HE1 TRP 46.258 50.073 1.00 Ν GLY 26 125.028 53.306 49.785 1.00 129.69 29 CA 26 125,460 54.647 49,432 1.00 129.22 GLY 30 126.079 54.744 48.049 C GLY 26 1.00 128.72 31 ō 125.794 53.929 GLY 26 47.177 1.00 128.98 32 Η GLY 26 124.083 53.132 49.888 1.00 25.00 33 N ASP 27 126.962 55.721 47.868 1.00 128.05 127.635 34 CA ASP 27 55.946 46.589 1.00 126.16 46.340 35 27 128.786 54.971 122.14 ASP 1.00 36 ASP 27 129.641 55.215 45.485 121.90 1.00 37 СВ ASP 27 128.154 57.390 46.495 1.00 128.56 38 CG ASP 27 127.036 58.414 46.382 1.00 129.57 39 OD1 ASP 27 126.092 58.200 45.590 1.00 129.32 27 40 OD2 ASP 127.10959.446 47.083 1.00 128.42 27 41 Η ASP 127.18856.294 48.627 1.00 25.00 42 Ν GLN 28 128,786 53.863 47 075 1.00 117.67 43 CA GLN 28 129.811 52.833 46.950 1.00 112.25 44 45.554 112.76 C GLN 28 129.807 52.195 1.00 45 GLN 130.803 45.131 110.25 O 28 51.612 1.00 СВ GLN 28 129.581 51.764 48.025 1.00 106.94 46 CG GLN 28 130.657 50.691 48.117 1.00 101.12 CD GLN 28 130.380 49.637 49.179 1.00 99.15 49 131.021 48.585 GLN 28 49.199 1.00 50 NE2 GLN 28 129.431 49.911 50.072 95.39 1.00 51 GLN 28 128.073 53.736 47.724 1.00 25.00 52 1HE2 GLN 28 129.287 49.206 50.749 1.00 25.00 53 2HE2 GLN 28 128.922 50.738 50.067 1.00 25.00 54 55 56 N PHE 29 128.696 52.349 44.833 1.00 116.96 29 51.766 CA PHF 128,536 43,496 1.00 118.7029 122.05 PHE 128.026 42.398 C 52.717 1.00 57 41.318 O PHE 29 127.643 52.249 1.00 123.10 58 127.570 29 50.572 43,539 CB PHE 1.00 114.02 59 PHE 127.906 49.532 44.568 CG 29 1.00 111.57 60 CD1 PHE 29 128.928 48.614 44.349 1.00 109.61 PHE 29 127.148 49.429 45.731 1.00 108.83 61 CD2 129.193 62 CE1 PHE 29 47.609 45.274 1.00 102.49 63 CE2 PHE 29 127.401 48.432 46.663 1.00 101.60 128.425 47.514 46.434 CZPHE 29 1.00 101.60 65 Н PHE 29 127.985 52.899 45.209 1.00 25.00 66 LEU 30 127.984 54.022 42.669 1.00 124.46 67 CA LEU 30 127.509 55,009 41.689 1.00 127.15 68 C LEU 30 128 208 54.874 40.335 1.00 128.12 127.578 127.732 69 O LEU 30 54.916 39.277 1.00 126.56 70 71 CB 42.230 LEU 30 56.433 1.00 128 23 127.357 57.672 129.74 LEU 30 41.388 1.00 CG 126.987 42.316 72 CD1 LEU 30 58.843 1.00 126.98 73 128.437 58.089 40.428 126.72 CD2 LEU 30 1.00 LEU 128.261 54.333 43.538 1.00 25.00

TABLE 11-continued

75 N SER 331 129.527 54.725 40.400 1.00 1.76 CA SER 31 130.384 54.608 39.222 1.00 1.77 C SER 31 130.384 54.608 39.222 1.00 1.77 C SER 31 131.618 53.722 39.458 1.00 1.78 SER 31 130.831 56.004 38.781 1.00 1.78 SER 31 130.830 56.073 341.303 1.00 1.78 SER 31 130.880 56.771 40.611 1.00 1.78 SER 32 133.156 52.065 38.458 1.00 1.78 SER 32 133.156 52.065 38.458 1.00 1.78 SER 32 133.156 52.065 38.458 1.00 1.78 SER 32 133.633 51.847 36.122 1.00 1.78 SER 32 133.803 51.847 36.122 1.00 1.78 SER 32 133.803 549.581 38.475 1.00 1.78 SER 32 133.805 49.581 38.475 1.00 1.78 SER 32 133.912 48.701 37.396 1.00 1.78 SER 32 133.912 48.701 37.396 1.00 1.78 SER 32 133.912 48.701 37.396 1.00 1.78 SER 32 133.805 49.581 38.475 1.00 1.78 SER 32 133.805 49.581 38.475 1.00 1.78 SER 32 133.805 49.581 38.475 1.00 1.78 SER 32 133.613 50.00 1.78 SER 33 135.219 52.861 37.434 1.00 1.78 SER 33 135.219 52.861 37.434 1.00 1.78 SER 33 135.219 52.861 37.434 1.00 1.78 SER 33 137.014 51.819 36.136 1.00 1.78 SER 33 137.014 5	B-factor 129.78 130.48 129.14 127.41 133.08 136.17 25.00 127.29 126.38 129.92 131.71 120.86 117.02 114.77 113.67 111.21 111.87 111.87 125.00 129.98 128.55 128.72
76 CA SER 31 130.384 54.608 39.222 1.00 177 C SER 31 131.618 53.722 39.458 1.00 178 C SER 31 131.618 53.722 39.458 1.00 179 CB SER 31 130.831 56.004 38.781 1.00 179 CB SER 31 130.831 56.004 38.781 1.00 179 CB SER 31 130.831 56.004 38.781 1.00 179 CB SER 31 131.461 56.700 39.845 1.00 179 CB SER 31 131.461 56.700 39.845 1.00 179 CB SER 31 130.880 56.771 40.611 1.00 180 SER 31 130.880 52.973 38.423 1.00 180 SER 31 130.880 52.973 38.423 1.00 180 SER 31 130.880 52.973 38.423 1.00 180 SER 32 133.656 52.249 37.231 1.00 180 SER 32 133.656 52.249 37.231 1.00 180 SER 32 133.693 51.847 36.122 1.00 180 SER 32 133.805 49.581 38.475 1.00 180 SER 32 133.912 48.701 37.396 1.00 180 SER 32 133.913 49.773 36.861 1.00 180 SER 33 135.851 47.658 38.389 1.00 180 SER 33 135.219 52.861 37.434 1.00 180 SER 33 137.019 54.277 36.684 1.00 180 SER	130.48 129.14 127.41 133.08 136.17 25.00 25.00 127.29 126.38 129.92 131.71 120.86 117.02 114.77 113.67 111.21 111.87 111.43 25.00 129.98 129.9
77 C SER 31 131.618 53.722 39.458 1.00 1 78 O SER 31 132.211 53.745 40.535 1.00 1 79 CB SER 31 130.831 56.004 38.781 1.00 1 80 OG SER 31 130.831 56.004 38.781 1.00 1 81 H SER 31 129.890 54.673 41.303 1.00 82 HG SER 31 130.880 56.771 40.611 1.00 83 N PHE 32 132.004 52.973 38.423 1.00 1 84 CA PHE 32 133.056 52.265 38.458 1.00 1 85 C PHE 32 133.056 52.249 37.231 1.00 1 86 O PHE 32 133.693 51.847 36.122 1.00 1 87 CB PHE 32 133.805 49.581 38.475 1.00 1 88 CG PHE 32 134.736 49.488 39.507 1.00 1 89 CD1 PHE 32 134.736 49.488 39.507 1.00 1 90 CD2 PHE 32 134.736 49.488 39.507 1.00 1 91 CE1 PHE 32 135.755 48.533 39.472 1.00 1 91 CE1 PHE 32 135.851 47.658 38.389 1.00 1 92 CE2 PHE 32 134.85 53.046 37.602 1.00 1 94 H PHE 32 131.485 53.046 37.602 1.00 1 95 N SER 33 135.219 52.861 37.434 1.00 1 96 CA SER 33 137.014 51.819 36.136 1.00 1 97 C SER 33 137.014 51.819 36.136 1.00 1 98 O SER 33 137.014 51.819 36.136 1.00 1 99 CB SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 10 H SER 33 135.40 51.71 7.70 33.955 1.00 1 10 G SER 33 137.014 51.819 36.136 1.00 1 11 H SER 34 136.616 51.017 35.153 1.00 1 11 H SER 34 136.616 51.017 35.153 1.00 1 11 H SER 34 136.616 51.017 35.153 1.00 1 11 H SER 34 136.616 51.017 35.153 1.00 1 11 H SER 34 136.616 51.017 35.153 1.00 1 11 H SER 34 136.616 51.017 35.153 1.00 1 11 H SER 34 136.616 51.017 35.555 1.00 1 11 H LE 34 136.518 46.611 32.665 1.00 1 11 H LE 34 136.518 46.	129.14 127.41 133.08 136.17 25.00 25.00 127.29 126.38 129.92 131.71 120.86 117.02 114.77 113.67 111.21 111.87 111.43 25.00 129.98 129.95 129.9
78 O SER 31 132.211 53.745 40.535 1.00 1 79 CB SER 31 130.831 56.004 38.781 1.00 1 80 OG SER 31 131.461 56.700 39.845 1.00 1 81 H SER 31 129.890 54.673 41.303 1.00 82 HG SER 31 130.880 56.771 40.611 1.00 83 N PHE 32 132.004 52.973 38.423 1.00 1 84 CA PHE 32 133.156 52.065 38.458 1.00 1 85 C PHE 32 133.056 52.249 37.231 1.00 1 86 O PHE 32 133.65 52.065 38.458 1.00 1 87 CB PHE 32 133.693 51.847 36.122 1.00 1 88 CG PHE 32 133.693 50.601 38.531 1.00 1 88 CG PHE 32 133.805 49.581 38.475 1.00 1 89 CD1 PHE 32 133.755 48.533 39.472 1.00 1 91 CE1 PHE 32 133.912 48.701 37.396 1.00 1 91 CE1 PHE 32 134.927 47.742 37.349 1.00 1 92 CE2 PHE 32 134.892 47.742 37.349 1.00 1 93 CZ PHE 32 134.85 53.046 37.602 1.00 1 94 H PHE 32 131.885 53.046 37.602 1.00 1 95 N SER 33 135.219 52.861 37.434 1.00 1 96 CA SER 33 137.014 51.819 36.136 1.00 1 97 C SER 33 137.014 51.819 36.136 1.00 1 98 O SER 33 137.079 54.277 36.684 1.00 1 99 CB SER 33 137.973 51.561 36.865 1.00 1 101 H SER 33 136.817 54.217 38.019 1.00 1 102 HG SER 33 137.514 51.819 36.136 1.00 1 104 CA ILE 34 138.313 49.773 34.842 1.00 1 105 C ILE 34 136.616 51.017 35.153 1.00 1 106 O ILE 34 138.313 49.773 34.842 1.00 1 107 CB ILE 34 136.616 51.017 35.153 1.00 1 108 CG1 ILE 34 138.388 50.555 1.00 1 109 CG2 ILE 34 136.648 48.884 33.865 1.00 1 101 H LE 34 136.616 51.017 35.153 1.00 1 102 HG SER 33 135.749 54.277 36.684 1.00 1 104 CA ILE 34 138.369 50.556 33.177 1.00 1 105 C ILE 34 138.369 50.556 33.177 1.00 1 107 CB ILE 34 136.616 51.017 35.153 1.00 1 108 CG1 ILE 34 136.648 48.884 33.865 1.00 1 109 CG2 ILE 34 136.648 48.884 33.865 1.00 1 110 CD1 ILE 34 136.518 46.611 32.665 1.00 1 111 H ILE 34 136.518 46.611 32.665 1.00 1 112 N ASP 35 141.105 49.749 34.548 1.00 1 113 CA ASP 35 141.105 49.749 34.548 1.00 1 114 C ASP 35 141.105 49.749 34.548 1.00 1 115 O ASP 35 141.105 49.749 34.548 1.00 1 116 CB ASP 35 141.105 49.749 34.548 1.00 1 117 CG ASP 35 141.105 49.749 35.509 1.00 1 118 OD1 ASP 35 141.105 49.02 37.955 1.00 1 119 OD2 ASP 35 141.105 49.02 37.955 1.00 1	127.41 133.08 136.17 25.00 25.00 127.29 126.38 129.92 131.71 120.86 117.02 114.77 113.67 111.21 111.87 111.43 25.00 129.98 129.98
79 CB SER 31 130.831 56.004 38.781 1.00 18 80 OG SER 31 131.461 56.700 39.845 1.00 18 18 H SER 31 129.890 54.673 41.303 1.00 18 22 HG SER 31 130.880 56.771 40.611 1.00 18 33 N PHE 32 132.004 52.973 38.423 1.00 18 38 CA PHE 32 133.156 52.065 38.458 1.00 18 86 O PHE 32 133.056 52.249 37.231 1.00 18 86 O PHE 32 133.693 51.847 36.122 1.00 18 87 CB PHE 32 133.693 51.847 36.122 1.00 18 87 CB PHE 32 133.805 49.581 38.475 1.00 19 90 CD2 PHE 32 133.305 49.581 38.475 1.00 19 91 CE1 PHE 32 135.755 48.533 39.472 1.00 19 91 CE1 PHE 32 135.755 48.533 39.472 1.00 19 93 CZ PHE 32 135.851 47.658 38.389 1.00 19 93 CZ PHE 32 135.851 47.658 38.389 1.00 19 96 CD PHE 32 135.755 48.533 39.472 1.00 19 97 CE SER 33 135.215 52.861 37.434 1.00 19 98 CA SER 33 135.219 52.861 37.434 1.00 19 99 CB SER 33 137.014 51.819 36.136 1.00 19 98 O SER 33 137.014 51.819 36.136 1.00 19 98 O SER 33 137.017 54.271 36.684 1.00 19 98 CB SER 33 137.017 54.271 36.684 1.00 19 90 CB SER 33 137.017 54.271 36.684 1.00 19 90 CB SER 33 137.017 54.271 36.684 1.00 19 90 CB SER 33 137.017 54.271 36.684 1.00 19 90 CB SER 33 137.017 54.271 36.684 1.00 10 10 00 G SER 33 137.017 54.271 36.685 1.00 10 10 00 G SER 33 137.017 54.271 36.684 1.00 10 10 00 G SER 33 137.017 54.271 36.684 1.00 10 10 00 G SER 33 137.017 54.271 36.684 1.00 10 10 1 H SER 33 136.817 54.182 38.626 1.00 10 10 10 CB SER 33 137.017 54.271 36.684 1.00 10 10 CB SER 33 137.017 54.271 36.684 1.00 10 10 CB SER 33 136.817 50.001 34.268 1.00 10 10 CB ILE 34 138.369 50.556 33.177 1.00 11 10 CD1 ILE 34 138.369 50.556 33.177 1.00 11 10 CD1 ILE 34 138.369 50.556 33.177 1.00 11 11 H ILE 34 136.616 51.017 35.153 1.00 11 11 H ILE 34 136.618 46.611 32.665 1.00 11 11 H ILE 34 136.483 48.884 33.865 1.00 11 11 H ILE 34 136.483 48.884 33.865 1.00 11 11 H ILE 34 136.483 48.884 33.865 1.00 11 11 H ILE 34 136.483 48.884 33.865 1.00 11 11 H ILE 34 136.483 48.884 33.865 1.00 11 11 CB ILE 34 136.483 48.884 33.865 1.00 11 11 CB ILE 34 136.483 48.884 33.865 1.00 11 11 CB ILE 34 136.518 46.611 32.665 1.00 11 11 CB ILE 34 136.518	133.08 136.17 25.00 25.00 127.29 126.38 129.92 131.71 120.86 117.02 114.77 113.67 111.21 111.87 111.43 25.00 129.98 128.55 128.72
80 OG SER 31 131.461 56.700 39.845 1.00 181 H SER 31 129.890 54.673 41.303 1.00 182 HG SER 31 130.880 56.771 40.611 1.00 182 HG SER 32 132.04 52.973 38.423 1.00 182 HZ SER 32 132.05 52.249 37.231 1.00 182 HZ SER 33 13.65 52.249 37.231 1.00 182 HZ SER 33 13.805 49.581 38.475 1.00 182 HZ SER 33 13.805 49.581 38.475 1.00 182 HZ SER 32 133.805 49.581 38.475 1.00 182 HZ SER 32 133.912 48.701 37.396 1.00 182 HZ SER 32 133.912 48.701 37.396 1.00 182 HZ SER 32 133.912 48.701 37.396 1.00 182 HZ SER 32 134.755 48.533 39.472 1.00 182 HZ SER 34 134.8927 47.742 37.349 1.00 182 HZ SER 34 134.8927 47.742 37.349 1.00 182 HZ SER 34 135.755 48.533 39.472 1.00 182 HZ SER 34 135.757 54.2511 38.019 1.00 182 HZ SER 34 137.079 54.277 36.684 1.00 182 HZ SER 34 137.379 54.277 36.684 1.00 182 HZ SER 34 137.379 54.277 36.684 1.00 182 HZ SER 34 135.545 54.211 38.019 1.00 182 HZ SER 34 136.616 51.017 35.153 1.00 1	136.17 25.00 25.00 127.29 126.38 129.92 131.71 120.86 117.02 114.77 111.87 111.43 25.00 129.98 128.55 128.72
81 H SER 31 129.890 54.673 41.303 1.00 82 HG SER 31 130.880 56.771 40.611 1.00 83 N PHE 32 132.004 52.973 38.423 1.00 1 84 CA PHE 32 133.156 52.065 38.458 1.00 1 85 C PHE 32 133.156 52.065 38.458 1.00 1 86 O PHE 32 133.693 51.847 36.122 1.00 1 87 CB PHE 32 133.693 51.847 36.122 1.00 1 88 CG PHE 32 133.683 50.601 38.531 1.00 1 88 CG PHE 32 133.683 50.601 38.531 1.00 1 89 CD1 PHE 32 133.4736 49.488 39.507 1.00 1 90 CD2 PHE 32 133.912 48.701 37.396 1.00 1 91 CE1 PHE 32 133.912 48.701 37.396 1.00 1 92 CE2 PHE 32 133.913 48.701 37.349 1.00 1 93 CZ PHE 32 134.927 47.742 37.349 1.00 1 95 N SER 33 135.219 52.861 37.434 1.00 1 95 N SER 33 135.219 52.861 37.434 1.00 1 96 CA SER 33 136.179 53.082 36.355 1.00 1 97 C SER 33 137.014 51.819 36.136 1.00 1 98 O SER 33 137.014 51.819 36.136 1.00 1 99 CB SER 33 137.557 54.277 36.684 1.00 1 101 H SER 33 137.557 54.211 38.019 1.00 1 102 HG SER 33 137.579 54.277 36.684 1.00 1 103 N ILE 34 136.616 51.017 35.153 1.00 1 104 CA ILE 34 136.616 51.017 35.153 1.00 1 105 C ILE 34 138.369 50.556 33.177 1.00 1 106 O ILE 34 136.483 48.884 33.865 1.00 1 107 CB ILE 34 136.483 48.884 33.865 1.00 1 108 CG1 ILE 34 137.313 49.773 34.842 1.00 1 109 CG2 ILE 34 136.518 46.611 32.665 1.00 1 100 CG SER 33 137.373 49.773 34.842 1.00 1 101 H SER 34 136.646 51.017 35.153 1.00 1 102 HG SER 33 137.373 49.773 34.842 1.00 1 103 N ILE 34 136.646 51.017 35.153 1.00 1 104 CA ILE 34 138.715 50.001 34.268 1.00 1 105 C ILE 34 138.369 50.556 33.177 1.00 1 107 CB ILE 34 136.646 51.017 35.153 1.00 1 108 CG1 ILE 34 136.744 49.640 32.570 1.00 1 109 CG2 ILE 34 136.518 46.611 32.665 1.00 1 110 CD1 ILE 34 136.518 46.611 32.665 1.00 1 111 H ILE 34 136.518 46.611 32.665 1.00 1 112 N ASP 35 141.493 47.661 33.726 1.00 1 113 CA ASP 35 141.105 49.749 34.548 1.00 1 114 C ASP 35 141.952 49.002 37.955 1.00 1 115 O ASP 35 141.780 48.684 36.761 1.00 1 116 CB ASP 35 141.780 48.684 36.761 1.00 1 117 CG ASP 35 141.780 48.684 36.761 1.00 1 119 OD2 ASP 35 141.780 48.684 36.761 1.00 1 110 OD3 ASP 35 141.780 48.684 36.761 1.00 1 110 OD3 ASP 3	25.00 25.00 127.29 126.38 129.92 131.71 120.86 117.02 114.77 113.67 111.21 111.43 25.00 129.98 128.55 128.72
82 HG SER 31 130.880 56.771 40.611 1.00 883 N PHE 32 132.004 52.973 38.423 1.00 1 84 CA PHE 32 133.156 52.065 38.458 1.00 1 85 C PHE 32 133.156 52.065 38.458 1.00 1 85 C PHE 32 133.693 51.847 36.122 1.00 1 87 CB PHE 32 133.693 51.847 36.122 1.00 1 87 CB PHE 32 132.683 50.601 38.531 1.00 1 88 CG PHE 32 133.805 49.581 38.475 1.00 1 90 CD2 PHE 32 133.805 49.581 38.475 1.00 1 90 CD2 PHE 32 133.912 48.701 37.396 1.00 1 91 CE1 PHE 32 135.755 48.533 39.472 1.00 1 91 CE1 PHE 32 135.755 48.533 39.472 1.00 1 93 CZ PHE 32 134.8927 47.742 37.349 1.00 1 93 CZ PHE 32 135.851 47.658 38.389 1.00 1 95 N SER 33 135.219 52.861 37.434 1.00 1 95 N SER 33 135.219 52.861 37.434 1.00 1 95 N SER 33 135.219 53.862 36.355 1.00 1 97 C SER 33 137.014 51.819 36.136 1.00 1 98 O SER 33 137.014 51.819 36.136 1.00 1 99 CB SER 33 137.973 51.561 36.865 1.00 1 101 H SER 33 135.440 53.163 38.329 1.00 1 101 H SER 33 136.817 54.182 38.626 1.00 102 HG SER 33 136.817 54.182 38.626 1.00 103 N ILE 34 136.616 51.017 35.153 1.00 1 104 CA ILE 34 137.313 49.773 34.842 1.00 1 105 C ILE 34 138.715 50.001 34.268 1.00 1 107 CB ILE 34 138.715 50.001 34.268 1.00 1 107 CB ILE 34 136.483 48.884 33.865 1.00 1 109 CG2 ILE 34 136.518 46.611 32.665 1.00 1 109 CG2 ILE 34 136.518 46.611 32.665 1.00 1 109 CG2 ILE 34 136.518 46.611 32.665 1.00 1 109 CG2 ILE 34 136.518 46.611 32.665 1.00 1 109 CG2 ILE 34 136.518 46.611 32.665 1.00 1 109 CG2 ILE 34 136.518 46.611 32.665 1.00 1 110 CD1 ILE 34 136.518 46.611 32.665 1.00 1 111 H ILE 34 136.518 46.611 32.665 1.00 1 112 N ASP 35 139.738 49.600 35.020 1.00 1 115 CG ASP 35 141.199 47.66 36.414 1.00 1 116 CB ASP 35 141.199 47.66 36.414 1.00 1 117 CG ASP 35 141.199 47.66 36.414 1.00 1 118 OD1 ASP 35 141.199 47.66 36.414 1.00 1 119 OD2 ASP 35 141.199 35.900 37.955 1.00 1 110 OD1 ASP 35 141.199 35.900 37.955 1.00 1 110 OD1 ASP 35 141.199 35.900 37.905 1.00 1 110 OD1 ASP 35 141.199 35.900 37.905 1.00 1 110 OD1 ASP 35 141.199 35.900 37.905 1.00 1 110 OD1 ASP 35 141.199 35.900 37.905 1.00 1 110 OD1 ASP 35 141.199 34.564 36.414 1.00 1 110 O	127.29 126.38 129.92 131.71 120.86 117.02 114.77 113.67 111.21 111.87 111.43 25.00 129.98 128.55 128.72
84         CA         PHE         32         133.156         52.065         38.458         1.00         1           85         C         PHE         32         134.056         52.249         37.231         1.00         1           86         O         PHE         32         133.693         51.847         36.122         1.00         1           87         CB         PHE         32         132.683         50.601         38.531         1.00         1           88         CG         PHE         32         133.805         49.581         38.475         1.00         1           89         CD1         PHE         32         134.736         49.488         39.507         1.00         1           90         CD2         PHE         32         135.755         48.533         39.472         1.00         1           91         CE1         PHE         32         135.755         48.533         39.472         1.00         1           92         CE2         PHE         32         135.851         47.648         33.349         1.00         1           93         CZ         PHE         32         131.485	126.38 129.92 131.71 120.86 117.02 114.77 113.67 111.21 111.87 111.43 25.00 129.98 128.55 128.72
85         C         PHE         32         134.056         52.249         37.231         1.00         1           86         O         PHE         32         133.693         51.847         36.122         1.00         1           87         CB         PHE         32         132.683         50.601         38.531         1.00         1           89         CD1         PHE         32         133.805         49.581         38.475         1.00         1           90         CD2         PHE         32         134.736         49.488         39.507         1.00         1           90         CD2         PHE         32         133.912         48.701         37.396         1.00         1           91         CE1         PHE         32         134.927         47.742         37.349         1.00         1           92         CE2         PHE         32         135.755         47.688         38.889         1.00         1           93         CZ         PHE         32         131.485         53.046         37.602         1.00           95         N         SER         33         135.219         52.861 </td <td>129.92 131.71 120.86 117.02 114.77 113.67 111.21 111.87 111.43 25.00 129.98 128.55 128.72</td>	129.92 131.71 120.86 117.02 114.77 113.67 111.21 111.87 111.43 25.00 129.98 128.55 128.72
86         O         PHE         32         133.693         51.847         36.122         1.00         1           87         CB         PHE         32         132.683         50.601         38.531         1.00         1           88         CG         PHE         32         133.805         49.581         38.475         1.00         1           89         CD1         PHE         32         134.736         49.488         39.507         1.00         1           90         CD2         PHE         32         135.755         48.533         39.472         1.00         1           91         CE1         PHE         32         135.755         48.533         39.472         1.00         1           92         CE2         PHE         32         135.851         47.658         38.389         1.00         1           93         CZ         PHE         32         135.851         47.658         38.389         1.00         1           94         H         PHE         32         135.851         47.658         38.389         1.00         1           95         N         SER         33         137.014	131.71 120.86 117.02 114.77 113.67 111.21 111.87 111.43 25.00 129.98 128.55 128.72
87         CB         PHE         32         132.683         50.601         38.531         1.00         1           88         CG         PHE         32         133.805         49.581         38.475         1.00         1           89         CD1         PHE         32         134.736         49.488         39.507         1.00         1           90         CD2         PHE         32         133.912         48.701         37.396         1.00         1           91         CE1         PHE         32         135.755         48.533         39.472         1.00         1           92         CE2         PHE         32         134.927         47.742         37.349         1.00         1           93         CZ         PHE         32         135.851         47.658         38.389         1.00         1           94         H         PHE         32         131.485         53.046         37.602         1.00         1           95         N         SER         33         136.179         53.082         36.355         1.00         1           96         CA         SER         33         137.014	120.86 117.02 114.77 113.67 111.21 111.87 111.43 25.00 129.98 128.55 128.72
88         CG         PHE         32         133.805         49.581         38.475         1.00         1           89         CD1         PHE         32         134.736         49.488         39.507         1.00         1           90         CD2         PHE         32         135.755         48.533         39.472         1.00         1           91         CE1         PHE         32         135.755         48.533         39.472         1.00         1           92         CE2         PHE         32         135.755         48.533         39.472         1.00         1           93         CZ         PHE         32         135.851         47.658         38.389         1.00         1           94         H         PHE         32         131.485         53.046         37.602         1.00         1           95         N         SER         33         135.179         52.861         37.434         1.00         1           96         CA         SER         33         137.014         51.819         36.136         1.00         1           98         O         SER         33         137.575	117.02 114.77 113.67 111.21 111.87 111.43 25.00 129.98 128.55 128.72
89         CD1         PHE         32         134.736         49.488         39.507         1.00         1           90         CD2         PHE         32         133.912         48.701         37.396         1.00         1           91         CE1         PHE         32         135.755         48.533         39.472         1.00         1           92         CE2         PHE         32         134.927         47.742         37.349         1.00         1           93         CZ         PHE         32         135.851         47.658         38.389         1.00         1           94         H         PHE         32         131.485         53.046         37.602         1.00           95         N         SER         33         136.179         53.082         36.355         1.00         1           96         CA         SER         33         137.014         51.819         36.136         1.00         1           97         C         SER         33         137.014         51.819         36.136         1.00         1           98         O         SER         33         137.079         54.217 <td>114.77 113.67 111.21 111.87 111.43 25.00 129.98 128.55 128.72</td>	114.77 113.67 111.21 111.87 111.43 25.00 129.98 128.55 128.72
90 CD2 PHE 32 133.912 48.701 37.396 1.00 1 91 CE1 PHE 32 135.755 48.533 39.472 1.00 1 92 CE2 PHE 32 135.851 47.658 38.389 1.00 1 93 CZ PHE 32 135.851 47.658 38.389 1.00 1 94 H PHE 32 131.485 53.046 37.602 1.00 95 N SER 33 135.219 52.861 37.434 1.00 1 96 CA SER 33 136.179 53.082 36.355 1.00 1 97 C SER 33 137.014 51.819 36.136 1.00 1 98 O SER 33 137.014 51.819 36.136 1.00 1 99 CB SER 33 137.079 54.277 36.684 1.00 1 100 OG SER 33 135.440 53.163 38.329 1.00 1 101 H SER 33 135.440 53.163 38.329 1.00 1 102 HG SER 33 135.440 53.163 38.329 1.00 1 103 N ILE 34 136.817 54.182 38.626 1.00 1 104 CA ILE 34 136.817 54.182 38.626 1.00 1 105 C ILE 34 138.715 50.001 34.268 1.00 1 106 O ILE 34 138.735 50.556 33.177 1.00 1 107 CB ILE 34 136.483 48.884 33.865 1.00 1 108 CG1 ILE 34 136.483 48.884 33.865 1.00 1 109 CG2 ILE 34 136.518 46.611 32.665 1.00 1 110 CD1 ILE 34 136.518 46.611 32.560 1.00 1 111 H ILE 34 136.518 46.611 32.560 1.00 1 112 N ASP 35 141.193 47.79 34.548 1.00 1 115 C ASP 35 141.193 47.79 34.548 1.00 1 116 CB ASP 35 141.193 47.79 35.700 1.00 1 117 CG ASP 35 141.193 47.79 35.700 1.00 1 118 OD1 ASP 35 141.180 48.684 36.761 1.00 1 119 OD2 ASP 35 141.180 48.684 36.761 1.00 1 110 OD1 ASP 35 141.180 48.684 36.761 1.00 1 111 OD2 ASP 35 141.180 49.002 37.955 1.00 1	113.67 111.21 111.87 111.43 25.00 129.98 128.55 128.72
92 CE2 PHE 32 134.927 47.742 37.349 1.00 193 CZ PHE 32 135.851 47.658 38.389 1.00 194 H PHE 32 131.485 53.046 37.602 1.00 195 N SER 33 135.219 52.861 37.434 1.00 196 CA SER 33 135.219 52.861 37.434 1.00 197 C SER 33 137.014 51.819 36.136 1.00 198 O SER 33 137.014 51.819 36.136 1.00 199 CB SER 33 137.079 54.277 36.684 1.00 199 CB SER 33 137.554 54.211 38.019 1.00 190 CG SER 33 135.540 53.163 38.329 1.00 190 CG SER 33 136.817 54.182 38.626 1.00 190 CG SER 33 136.817 54.182 38.626 1.00 190 CG SER 33 136.817 54.182 38.626 1.00 190 CG SER 34 138.715 50.001 34.268 1.00 190 CG SER 34 138.715 50.001 34.842 1.00 190 CG SER 34 138.715 50.001 34.268 1.00 190 CG SER 34 138.715 50.001 34.268 1.00 190 CG SER 34 138.715 50.001 34.268 1.00 190 CG SER 34 136.518 46.611 32.665 1.00 190 CG SER 35 141.437 48.728 35.700 1.00	111.87 111.43 25.00 129.98 128.55 128.72
93 CZ PHE 32 135.851 47.658 38.389 1.00 194 H PHE 32 131.485 53.046 37.602 1.00 195 N SER 33 135.219 52.861 37.434 1.00 196 CA SER 33 136.179 53.082 36.355 1.00 197 C SER 33 137.014 51.819 36.136 1.00 198 O SER 33 137.014 51.819 36.136 1.00 198 O SER 33 137.079 54.277 36.684 1.00 199 CB SER 33 137.079 54.277 36.684 1.00 190 OG SER 33 137.545 54.211 38.019 1.00 190 OG SER 33 135.440 53.163 38.329 1.00 190 OG SER 33 135.440 53.163 38.329 1.00 190 OG SER 33 135.440 53.163 38.329 1.00 190 OG SER 34 136.616 51.017 35.153 1.00 190 OG SER 34 138.319 54.211 38.019 1.00 190 OG SER 34 138.319 59.556 33.177 1.00 190 OG SER 34 138.319 50.556 33.177 1.00 190 OG SER 34 138.369 50.556 33.177 1.00 190 OG SER 34 138.369 50.556 33.177 1.00 190 OG SER 34 136.483 48.884 33.865 1.00 190 OG SER 34 136.483 48.884 33.865 1.00 190 OG SER 34 136.518 46.611 32.665 1.00 190 OC SER 34 136.518 49.640 32.570 1.00 190 OC SER 35 141.437 48.728 33.464 1.00 190 OC SER 35 141.438 48.884 33.755 1.00 19	111.43 25.00 129.98 128.55 128.72
94 H PHE 32 131.485 53.046 37.602 1.00 95 N SER 33 135.219 52.861 37.434 1.00 1 96 CA SER 33 136.179 53.082 36.355 1.00 1 97 C SER 33 137.014 51.819 36.136 1.00 1 98 O SER 33 137.014 51.819 36.136 1.00 1 98 O SER 33 137.079 54.277 36.684 1.00 1 100 OG SER 33 137.55 54.211 38.019 1.00 1 101 H SER 33 135.440 53.163 38.329 1.00 102 HG SER 33 135.440 53.163 38.329 1.00 102 HG SER 33 136.817 54.182 38.626 1.00 103 N ILE 34 136.616 51.017 35.153 1.00 1 104 CA ILE 34 137.313 49.773 34.842 1.00 1 105 C ILE 34 138.715 50.001 34.268 1.00 1 106 O ILE 34 138.369 50.556 33.177 1.00 1 107 CB ILE 34 138.369 50.556 33.177 1.00 1 108 CG1 ILE 34 136.483 48.884 33.865 1.00 1 109 CG2 ILE 34 136.483 48.884 33.865 1.00 1 110 CD1 ILE 34 136.518 46.611 32.665 1.00 1 111 H ILE 34 136.518 46.611 32.655 1.00 1 112 N ASP 35 139.738 49.600 35.020 1.00 1 113 CA ASP 35 141.493 47.79 33.796 1.00 1 115 O ASP 35 141.493 47.661 33.726 1.00 1 116 CB ASP 35 141.993 47.661 33.726 1.00 1 117 CG ASP 35 141.993 47.661 33.726 1.00 1 118 OD1 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.983 47.661 33.726 1.00 1 119 OD2 ASP 35 141.983 47.661 33.720 37.90 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 119 OD2 ASP 35 141.982 47.566 36.414 1.00 1 1100 DD2 ASP 35 141.982 49.002 37.955 1.00 1 100 DD2 ASP 35 141.9	25.00 129.98 128.55 128.72
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97 C SER 33 137.014 51.819 36.136 1.00 1 98 O SER 33 137.973 51.561 36.865 1.00 1 99 CB SER 33 137.975 54.277 36.684 1.00 1 100 OG SER 33 137.554 54.211 38.019 1.00 1 101 H SER 33 135.440 53.163 38.329 1.00 102 HG SER 33 135.440 53.163 38.329 1.00 103 N ILE 34 136.616 51.017 35.153 1.00 1 104 CA ILE 34 138.715 50.001 34.268 1.00 1 105 C ILE 34 138.715 50.001 34.268 1.00 1 106 O ILE 34 138.369 50.556 33.177 1.00 1 107 CB ILE 34 136.483 48.884 33.865 1.00 1 108 CG1 ILE 34 136.483 48.884 33.865 1.00 1 109 CG2 ILE 34 136.174 49.640 32.570 1.00 1 110 CD1 ILE 34 136.518 46.611 32.665 1.00 1 111 H ILE 34 135.838 51.274 34.627 1.00 1 112 N ASP 35 141.953 49.709 35.700 1.00 1 115 O ASP 35 141.437 48.728 33.464 1.00 1 116 CB ASP 35 141.437 48.728 33.726 1.00 1 117 CG ASP 35 141.178 48.728 33.795 1.00 1 118 OD1 ASP 35 141.1780 48.684 36.661 1.00 1 119 OD2 ASP 35 141.180 48.684 36.661 1.00 1 119 OD2 ASP 35 141.180 48.684 36.661 1.00 1 119 OD2 ASP 35 141.180 49.002 37.955 1.00 1 119 OD2 ASP 35 141.180 49.002 37.955 1.00 1	128.72
98 O SER 33 137.973 51.561 36.865 1.00 1 99 CB SER 33 137.079 54.277 36.684 1.00 1 100 OG SER 33 137.554 54.211 38.019 1.00 1 101 H SER 33 135.440 53.163 38.329 1.00 102 HG SER 33 136.817 54.182 38.626 1.00 103 N ILE 34 136.616 51.017 35.153 1.00 1 104 CA ILE 34 137.313 49.773 34.842 1.00 1 105 C ILE 34 138.715 50.001 34.268 1.00 1 106 O ILE 34 138.369 50.556 33.177 1.00 1 107 CB ILE 34 136.483 48.884 33.865 1.00 1 108 CG1 ILE 34 137.227 47.570 33.595 1.00 1 109 CG2 ILE 34 136.518 46.611 32.665 1.00 1 110 CD1 ILE 34 136.518 46.611 32.665 1.00 1 111 H ILE 34 136.518 46.611 32.665 1.00 1 112 N ASP 35 141.437 48.728 33.468 1.00 1 115 O ASP 35 141.437 48.728 33.426 1.00 1 116 CB ASP 35 141.437 48.728 33.426 1.00 1 117 CG ASP 35 141.437 48.728 33.426 1.00 1 118 OD1 ASP 35 141.780 47.661 33.726 1.00 1 119 OD2 ASP 35 141.780 49.023 37.955 1.00 1 119 OD2 ASP 35 141.932 47.566 36.414 1.00 1 119 OD2 ASP 35 141.952 49.002 37.955 1.00 1	
99 CB SER 33 137.079 54.277 36.684 1.00 1 100 OG SER 33 137.554 54.211 38.019 1.00 1 101 H SER 33 135.440 53.163 38.329 1.00 102 HG SER 33 136.817 54.182 38.626 1.00 103 N ILE 34 136.616 51.017 35.153 1.00 1 104 CA ILE 34 137.313 49.773 34.842 1.00 1 105 C ILE 34 138.715 50.001 34.268 1.00 1 106 O ILE 34 138.369 50.556 33.177 1.00 1 107 CB ILE 34 136.483 48.884 33.865 1.00 1 108 CG1 ILE 34 137.227 47.570 33.595 1.00 1 109 CG2 ILE 34 136.174 49.640 32.570 1.00 1 110 CD1 ILE 34 136.518 46.611 32.656 1.00 1 111 H ILE 34 135.838 51.274 34.627 1.00 1 112 N ASP 35 139.738 49.600 35.020 1.00 1 113 CA ASP 35 141.105 49.749 34.548 1.00 1 114 C ASP 35 141.105 49.749 34.548 1.00 1 115 O ASP 35 141.193 47.661 33.726 1.00 1 116 CB ASP 35 141.180 48.684 36.761 1.00 1 117 CG ASP 35 141.180 48.684 36.761 1.00 1 118 OD1 ASP 35 141.180 48.684 36.761 1.00 1 119 OD2 ASP 35 141.180 49.00 35.909 1.00	
100         OG         SER         33         137.554         54.211         38.019         1.00         1           101         H         SER         33         135.440         53.163         38.329         1.00           102         HG         SER         33         136.817         54.182         38.626         1.00           103         N         ILE         34         136.616         51.017         35.153         1.00         1           104         CA         ILE         34         137.313         49.773         34.842         1.00         1           105         C         ILE         34         138.315         50.001         34.268         1.00         1           106         O         ILE         34         138.369         50.556         33.177         1.00         1           107         CB         ILE         34         136.483         48.884         33.865         1.00         1           108         CG1         ILE         34         136.174         49.640         32.570         1.00         1           110         CD1         ILE         34         136.518         46.611         <	130.20
102         HG         SER         33         136.817         54.182         38.626         1.00           103         N         ILE         34         136.616         51.017         35.153         1.00         1           104         CA         ILE         34         137.313         49.773         34.842         1.00         1           105         C         ILE         34         138.715         50.001         34.842         1.00         1           106         O         ILE         34         138.369         50.556         33.177         1.00         1           107         CB         ILE         34         136.483         48.884         33.865         1.00         1           108         CG1         ILE         34         136.483         48.884         33.865         1.00         1           109         CG2         ILE         34         136.174         49.640         32.570         1.00         1           110         CD1         ILE         34         136.518         46.611         32.665         1.00         1           111         H         ILE         34         135.838         5	130.89
103         N         ILE         34         136.616         51.017         35.153         1.00         1           104         CA         ILE         34         137.313         49.773         34.842         1.00         1           105         C         ILE         34         138.715         50.001         34.268         1.00         1           106         O         ILE         34         138.369         50.556         33.177         1.00         1           107         CB         ILE         34         136.483         48.884         33.865         1.00         1           108         CG1         ILE         34         136.174         49.640         32.570         1.00         1           110         CD1         ILE         34         136.518         46.611         32.665         1.00         1           111         H         ILE         34         135.838         51.274         34.627         1.00           112         N         ASP         35         139.738         49.600         35.020         1.00         1           113         CA         ASP         35         141.105         49.	25.00
104         CA         ILE         34         137.313         49.773         34.842         1.00         1           105         C         ILE         34         138.715         50.001         34.268         1.00         1           106         O         ILE         34         138.369         50.556         33.177         1.00         1           107         CB         ILE         34         136.483         48.884         33.865         1.00         1           108         CG1         ILE         34         137.227         47.570         33.595         1.00         1           109         CG2         ILE         34         136.174         49.640         32.570         1.00         1           110         CD1         ILE         34         136.518         46.611         32.665         1.00         1           111         H         ILE         34         135.838         51.274         34.627         1.00         1           112         N         ASP         35         139.738         49.600         35.020         1.00         1           113         CA         ASP         35         141.105	25.00
105         C         ILE         34         138.715         50.001         34.268         1.00         1           106         O         ILE         34         138.369         50.556         33.177         1.00         1           107         CB         ILE         34         136.483         48.884         33.865         1.00         1           108         CG1         ILE         34         136.174         49.640         32.570         1.00         1           110         CD1         ILE         34         136.518         46.611         32.665         1.00         1           111         H         ILE         34         135.838         51.274         34.627         1.00           112         N         ASP         35         139.738         49.600         35.020         1.00         1           113         CA         ASP         35         141.105         49.749         34.548         1.00         1           114         C         ASP         35         141.993         47.661         33.726         1.00         1           115         O         ASP         35         141.993         47.6	128.06
106         O         ILE         34         138.369         50.556         33.177         1.00         1           107         CB         ILE         34         136.483         48.884         33.865         1.00         1           108         CG1         ILE         34         137.227         47.570         33.595         1.00         1           109         CG2         ILE         34         136.518         46.611         32.665         1.00         1           110         CD1         ILE         34         135.838         51.274         34.627         1.00           111         H         ILE         34         135.838         51.274         34.627         1.00           112         N         ASP         35         139.738         49.600         35.020         1.00         1           113         CA         ASP         35         141.105         49.749         34.548         1.00         1           114         C         ASP         35         141.993         47.661         33.726         1.00         1           115         O         ASP         35         141.993         47.661         <	127.26
107         CB         ILE         34         136.483         48.884         33.865         1.00         1           108         CG1         ILE         34         137.227         47.570         33.595         1.00         1           109         CG2         ILE         34         136.174         49.640         32.570         1.00         1           110         CD1         ILE         34         136.518         46.611         32.665         1.00         1           111         H         ILE         34         135.838         51.274         34.627         1.00           112         N         ASP         35         139.738         49.600         35.020         1.00         1           113         CA         ASP         35         141.105         49.749         34.548         1.00         1           114         C         ASP         35         141.437         48.728         33.426         1.00         1           115         O         ASP         35         141.939         47.661         33.726         1.00         1           116         CB         ASP         35         141.780         4	128.81 132.56
108         CG1         ILE         34         137.227         47.570         33.595         1.00         1           109         CG2         ILE         34         136.174         49.640         32.570         1.00         1           110         CD1         ILE         34         136.518         46.611         32.665         1.00         1           111         H         ILE         34         135.838         51.274         34.627         1.00         1           112         N         ASP         35         139.738         49.600         35.020         1.00         1           113         CA         ASP         35         141.105         49.749         34.548         1.00         1           114         C         ASP         35         141.993         47.661         33.726         1.00         1           115         O         ASP         35         141.993         47.661         33.726         1.00         1           116         CB         ASP         35         142.122         49.709         35.700         1.00         1           117         CG         ASP         35         141.784	125.22
109         CG2         ILE         34         136.174         49.640         32.570         1.00         1           110         CD1         ILE         34         136.518         46.611         32.665         1.00         1           111         H         ILE         34         135.838         51.274         34.627         1.00         1           112         N         ASP         35         139.738         49.600         35.020         1.00         1           113         CA         ASP         35         141.105         49.749         34.548         1.00         1           114         C         ASP         35         141.437         48.728         33.464         1.00         1           115         O         ASP         35         141.993         47.661         33.726         1.00         1           116         CB         ASP         35         142.122         49.709         35.700         1.00         1           117         CG         ASP         35         141.784         48.684         36.761         1.00         1           118         OD1         ASP         35         141.92<	124.19
111         H         ILE         34         135.838         51.274         34.627         1.00           112         N         ASP         35         139.738         49.600         35.020         1.00         1           113         CA         ASP         35         141.105         49.749         34.548         1.00         1           114         C         ASP         35         141.437         48.728         33.464         1.00         1           115         O         ASP         35         141.993         47.661         33.726         1.00         1           116         CB         ASP         35         142.122         49.709         35.700         1.00         1           117         CG         ASP         35         141.780         48.684         36.761         1.00         1           118         OD1         ASP         35         141.342         47.566         36.414         1.00         1           119         OD2         ASP         35         141.952         49.002         37.955         1.00         1           120         H         ASP         35         139.577         49.	123.27
112     N     ASP     35     139.738     49.600     35.020     1.00     1       113     CA     ASP     35     141.105     49.749     34.548     1.00     1       114     C     ASP     35     141.437     48.728     33.464     1.00     1       115     O     ASP     35     141.939     47.661     33.726     1.00     1       116     CB     ASP     35     142.122     49.709     35.700     1.00     1       117     CG     ASP     35     141.780     48.684     36.761     1.00     1       118     OD1     ASP     35     141.342     47.566     36.414     1.00     1       119     OD2     ASP     35     141.952     49.002     37.955     1.00     1       120     H     ASP     35     139.577     49.234     35.909     1.00	121.43
113     CA     ASP     35     141.105     49.749     34.548     1.00     1       114     C     ASP     35     141.437     48.728     33.464     1.00     1       115     O     ASP     35     141.993     47.661     33.726     1.00     1       116     CB     ASP     35     142.122     49.709     35.700     1.00     1       117     CG     ASP     35     141.780     48.684     36.761     1.00     1       118     OD1     ASP     35     141.342     47.566     36.414     1.00     1       119     OD2     ASP     35     141.952     49.002     37.955     1.00     1       120     H     ASP     35     139.577     49.234     35.909     1.00	25.00
114     C     ASP     35     141.437     48.728     33.464     1.00     1       115     O     ASP     35     141.993     47.661     33.726     1.00     1       116     CB     ASP     35     142.122     49.709     35.700     1.00     1       117     CG     ASP     35     141.782     48.684     36.761     1.00     1       118     OD1     ASP     35     141.342     47.566     36.414     1.00     1       119     OD2     ASP     35     141.952     49.002     37.955     1.00     1       120     H     ASP     35     139.577     49.234     35.909     1.00	125.91
115     O     ASP     35     141.993     47.661     33.726     1.00     1       116     CB     ASP     35     142.122     49.709     35.700     1.00     1       117     CG     ASP     35     141.782     48.684     36.761     1.00     1       118     OD1     ASP     35     141.342     47.566     36.414     1.00     1       119     OD2     ASP     35     141.952     49.002     37.955     1.00     1       120     H     ASP     35     139.577     49.234     35.909     1.00	120.92
116     CB     ASP     35     142.122     49.709     35.700     1.00     1       117     CG     ASP     35     141.780     48.684     36.761     1.00     1       118     OD1     ASP     35     141.342     47.566     36.414     1.00     1       119     OD2     ASP     35     141.952     49.002     37.955     1.00     1       120     H     ASP     35     139.577     49.234     35.909     1.00	116.38 115.56
117     CG     ASP     35     141.780     48.684     36.761     1.00     1       118     OD1     ASP     35     141.342     47.566     36.414     1.00     1       119     OD2     ASP     35     141.952     49.002     37.955     1.00     1       120     H     ASP     35     139.577     49.234     35.909     1.00	122.56
118 OD1 ASP 35 141.342 47.566 36.414 1.00 1 119 OD2 ASP 35 141.952 49.002 37.955 1.00 1 120 H ASP 35 139.577 49.234 35.909 1.00	123.82
120 H ASP 35 139.577 49.234 35.909 1.00	129.46
	123.01
	25.00
	111.08
	107.72
	104.04 103.31
	108.78
126 CG ASN 38 141.378 50.742 30.076 1.00 1	113.72
	115.16
	113.86
129 H ASN 36 140.490 49.899 32.196 1.00	25.00
130 1HD2 ASN 36 141.642 52.070 31.558 1.00	25.00
131 2HD2 ASN 36 141.817 50.557 32.067 1.00	25.00 101.33
132 N GLN 37 143.593 47.958 31.571 1.00 1 133 CA GLN 37 144.857 47.226 31.576 1.00	97.84
134 C GLN 37 144.752 45.887 32.306 1.00	91.48
135 O GLN 37 145.120 44.848 31.756 1.00	87.36
	104.38
137 CG GLN 37 147.329 47.907 31.541 1.00 1	109.27
	112.48
	116.42
	113.85
141 H GLN 37 143.558 48.839 31.989 1.00 142 1HE2 GLN 37 146.735 48.687 28.371 1.00	
142 THE2 GLN 37 146.733 46.867 26.371 1.00 143 2HE2 GLN 37 146.083 47.436 29.358 1.00	25.00
144 N VAL 38 144.242 45.916 33.536 1.00	25.00 25.00
145 CA VAL 38 144.092 44.702 34.337 1.00	25.00 25.00 25.00
146 C VAL 38 143.148 43.731 33.634 1.00	25.00 25.00
147 O VAL 38 143.416 42.529 33.568 1.00	25.00 25.00 25.00 85.52
148 CB VAL 38 143.542 45.018 35.752 1.00	25.00 25.00 25.00 85.52 79.25

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
149	CG1	VAL	38	143.484	43.754	36.593	1.00	75.39
150	CG2	VAL	38	144.409	46.061	36.437	1.00	78.10
151	H	VAL	38	143.943	46.770	33.907	1.00	25.00
152	N	ALA	39	142.060	44.268	33.086	1.00	71.24
153	CA	ALA	39	141.071	43.463	32.379	1.00	66.46
154	С	ALA	39	141.694	42.736	31.191	1.00	65.26
155	O	ALA	39	141.519	41.527	31.038	1.00	59.37
156 157	CB H	ALA ALA	39 39	139.910 141.924	44.338 45.233	31.916 33.161	$\frac{1.00}{1.00}$	64.84 25.00
158	N	GLU	40	142.436	43.472	30.366	1.00	66.68
159	CA	GLU	40	143.086	42.896	29.190	1.00	69.85
160	C	GLU	40	144.107	41.828	29.559	1.00	66.85
161	O	GLU	40	144.233	40.818	28.859	1.00	65.59
162	CB	GLU	40	143.744	43.985	28.342	1.00	75.56
163	CG	GLU	40	142.752	44.836	27.560	1.00	89.79
164	CD	GLU	40	143.409	45.993	26.828	1.00	98.34
165	OE1	GLU	40	144.515	45.807	26.273	1.00	101.87
166	OE2	GLU	40	142.814	47.092	26.808	1.00	102.72
167	H	GLU	40	142.551	44.427	30.550	1.00	25.00
168	N	LYS	41	144.830	42.048	30.656	1.00	62.69
169 170	CA C	LYS LYS	41 41	145.821 145.081	41.079 39.798	31.112 31.478	$\frac{1.00}{1.00}$	60.33 56.90
170	Ö	LYS	41	145.440	38.707	31.024	1.00	56.12
172	СВ	LYS	41	146.588	41.603	32.331	1.00	64.30
173	CG	LYS	41	147.689	40.655	32.802	1.00	70.61
174	CD	LYS	41	148.373	41.137	34.070	1.00	74.86
175	CE	LYS	41	149.449	40.152	34.505	1.00	79.07
176	NZ	LYS	41	150.138	40.584	35.753	1.00	86.06
177	H	LYS	41	144.700	42.879	31.160	1.00	25.00
178	1HZ	LYS	41	150.588	41.510	35.601	1.00	25.00
179	2HZ	LYS	41	149.443	40.661	36.524	1.00	25.00
180	3HZ	LYS	41	150.884	39.885	36.010	1.00	25.00
181	N	TYR	42	144.027	39.951	32.278	1.00	54.48
182	CA	TYR	42	143.200	38.831	32.712	1.00	49.46
183 184	C O	TYR TYR	42 42	142.687 142.886	38.048 36.837	31.508 31.418	$\frac{1.00}{1.00}$	49.51 46.83
185	СВ	TYR	42	142.011	39.332	33.535	1.00	49.09
186	CG	TYR	42	142.316	39.665	34.981	1.00	51.81
187	CD1	TYR	42	143.609	39.555	35.498	1.00	55.25
188	CD2	TYR	42	141.297	40.067	35.844	1.00	51.34
189	CE1	TYR	42	143.873	39.836	36.843	1.00	60.63
190	CE2	TYR	42	141.548	40.347	37.180	1.00	51.18
191	CZ	TYR	42	142.832	40.231	37.677	1.00	57.44
192	OH	TYR	42	143.064	40.503	39.009	1.00	57.29
193	H	TYR	42	143.796	40.855	32.582	1.00	25.00
194	HH	TYR	42	142.245	40.765	39.435	1.00	25.00
195	N	ALA	43	142.067	38.756	30.568	1.00	48.90
196 197	CA C	ALA	43 43	141.514	38.150 37.363	29.359	1.00	49.75 49.98
197	Ö	ALA ALA	43	142.560 142.331	36.209	28.576 28.204	$\frac{1.00}{1.00}$	49.38
199	СВ	ALA	43	140.897	39.223	28.477	1.00	46.66
200	H	ALA	43	141.980	39.722	30.693	1.00	25.00
201	N	GLN	44	143.711	37.987	22.344	1.00	52.04
202	CA	GLN	44	144.796	37.352	27.607	1.00	51.53
203	C	GLN	44	145.219	36.030	28.257	1.00	45.51
204	O	GLN	44	145.304	35.002	27.582	1.00	43.18
205	CB	GLN	44	145.994	38.299	27.506	1.00	58.59
206	CG	GLN	44	147.101	37.804	26.583	1.00	74.05
207	CD	GLN	44	148.364	38.649	26.658	1.00	84.03
208	OE1	GLN	44	148.343	39.784	27.132	1.00	90.02
209 210	NE2 H	GLN GLN	44 44	149.475 143.837	38.092 38.900	26.187	$\frac{1.00}{1.00}$	84.98 25.00
210	1HE2	GLN	44	150.290	38.631	28.677 26.238	1.00	25.00
212	2HE2	GLN	44	149.438	37.187	25.820	1.00	25.00
213	N	GLU	45	145.468	36.047	29.565	1.00	40.03
214	CA	GLU	45	145.874	34.831	30.261	1.00	37.78
215	C	GLU	45	144.740	33.813	30.320	1.00	41.99
216	0	GLU	45	144.970	32.609	30.153	1.00	43.40
217	CB	GLU	45	146.374	35.134	31.673	1.00	38.09
218	CG	GLU	45	147.037	33.924	32.334	1.00	41.87
219	CD	GLU	45	147.595	34.209	33.718	1.00	52.34
220	OE1	GLU	45	147.678	35.393	34.116	1.00	58.77
221	OE2	GLU	45	147.962	33.235	34.409	1.00	53.14
222	H	GLU	45	145.372	36.890	30.064	1.00	25.00

TABLE 11-continued

Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase	
In the Absence of Bound Substrate	

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
223	N	ILE	46	143.521	34.296	30.553	1.00	38.09
224	CA	ILE	46	142.352	33.428	30.622	1.00	35.19
225	С	ILE	46	142.239	32.630	29.328	1.00	37.05
226	O	ILE	46	141.923	31.441	29.360	1.00	40.60
227	СВ	ILE	48	141.054	34.236	30.886	1.00	33.29
228	CG1	ILE	46	140.992	34.650	32.357	1.00	25.99
229	CG2	ILE	46	139.817	33.420	30.528	1.00	33.03
230 231	CD1 H	ILE ILE	46 46	139.889 143.414	35.630 35.255	32.687 30.675	$\frac{1.00}{1.00}$	27.20 25.00
232	N	GLU	47	142.548	33.263	28.199	1.00	37.32
233	CA	GLU	47	142.485	32.581	26.910	1.00	43.67
234	C	GLU	47	143.420	31.379	26.870	1.00	44.27
235	O	GLU	47	143.061	30.324	26.341	1.00	48.49
236	CB	GLU	47	142.817	33.537	25.765	1.00	50.83
237	CG	GLU	47	141.700	34.516	25.422	1.00	72.13
238	CD	GLU	47	140.408	33.833	24.970	1.00	80.45
239	OE1	GLU	47	140.440	32.643	24.577	1.00	82.39
240	OE2	GLU	47	139.353	34.501	25.001	1.00	87.14
241	H	GLU	47	142.813	34.208	28.233	1.00	25.00
242	N	ALA	48	144.610	31.538	27.444	1.00	41.56
243 244	CA C	ALA ALA	48 48	145.597 145.078	30.464 29.340	27.489 28.375	$\frac{1.00}{1.00}$	36.13 38.33
244	Ö	ALA	48	145.027	28.176	27.964	1.00	40.71
246	СВ	ALA	48	146.917	30.990	28.031	1.00	33.18
247	H	ALA	48	144.824	32.404	27.853	1.00	25.00
248	N	LEU	49	144.662	29.708	29.583	1.00	37.02
249	CA	LEU	49	144.136	28.757	30.554	1.00	34.21
250	C	LEU	49	142.894	28.044	30.029	1.00	33.34
251	O	LEU	49	142.694	26.860	30.296	1.00	34.71
252	CB	LEU	49	143.816	29.477	31.862	1.00	32.92
253	CG	LEU	49	145.013	30.132	32.551	1.00	29.78
254	CD1	LEU	49	144.541	31.096	33.621	1.00	28.55
255	CD2	LEU	49	145.915	29.062	33.139	1.00	31.11
256	H	LEU	49	144.717	30.657	29.827	1.00	25.00
257 258	N CA	LYS LYS	50 50	142.083 140.858	28.759 28.208	29.254 28.681	$\frac{1.00}{1.00}$	35.43 36.99
259	C	LYS	50	141.193	27.105	27.687	1.00	39.13
260	Ö	LYS	50	140.643	26.004	27.762	1.00	39.46
261	СВ	LYS	50	140.056	29.307	27.981	1.00	38.20
262	CG	LYS	50	138.670	28.882	27.520	1.00	37.82
263	CD	LYS	50	138.021	29.947	26.638	1.00	41.56
264	CE	LYS	50	137.926	31.297	27.341	1.00	44.73
265	NZ	LYS	50	137.282	32.342	26.489	1.00	40.08
266	H	LYS	50	142.323	29.682	29.055	1.00	25.00
267	1HZ	LYS	50	136.318	32.042	26.239	1.00	25.00
268	2HZ	LYS	50	137.245	33.240	27.012	1.00	25.00
269	3HZ	LYS	50	137.841	32.474	25.620	1.00	25.00
270 271	N	GLU	51 51	142.106 142.516	27.396	26.765 25.762	1.00	41.23 44.62
271	CA C	GLU GLU	51	143.174	26.419 25.226	26.446	$\frac{1.00}{1.00}$	42.95
273	Ö	GLU	51	142.931	24.073	26.091	1.00	43.13
274	CB	GLU	51	143.489	27.055	24.766	1.00	52.90
275	CG	GLU	51	143.846	26.162	23.581	1.00	70.21
276	CD	GLU	51	142.623	25.709	22.792	1.00	79.98
277	OE1	GLU	51	141.917	26.575	22.226	1.00	86.16
278	OE2	GLU	51	142.368	24.486	22.739	1.00	81.08
279	H	GLU	51	142.510	28.293	26.751	1.00	25.00
280	N	GLN	52	143.965	25.514	27.471	1.00	48.21
281	CA	GLN	52	144.662	24.480	28.223	1.00	49.28
282	С	GLN	52 52	143.657	23.563	28.933	1.00 1.00	44.99
283 284	O CB	GLN	52 52	143.817	22.337	28.936 29.230	1.00	42.93 51.94
285	СБ	GLN GLN	52 52	145.609 146.728	25.138 24.247	29.230	1.00	57.86
286	CD	GLN	52	147.655	24.973	30.696	1.00	61.66
287	OE1	GLN	52	147.719	26.205	30.711	1.00	53.55
288	NE2	GLN	52	148.372	24.211	31.511	1.00	66.48
289	Н	GLN	52	144.095	26.453	27.720	1.00	25.00
290	1HE2	GLN	52	148.989	24.676	32.114	1.00	25.00
291	2HE2	GLN	52	148.283	23.237	31.480	1.00	25.00
292	N	THR	53	142.615	24.160	29.512	1.00	42.37
293	CA	THR	53	141.578	23.404	30.214	1.00	41.44
294	С	THR	53	140.753	22.584	29.220	1.00	40.96
295	O	THR	53	140.334	21.462	29.519	1.00	38.12
296	CB	THR	53	140.648	24.338	31.027	1.00	42.33

TABLE 11-continued

	In the Absence of Bound Substrate											
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor				
297	CG1	THR	53	141.420	25.054	32.001	1.00	42.93				
298	CG2	THR	53	139.569	23.534	31.746	1.00	41.61				
299	H	THR	53	142.537	25.134	29.467	1.00	25.00				
300 301	HG1 N	THR ARG	53 54	142.088 140.553	25.586 23.138	31.555 28.027	$\frac{1.00}{1.00}$	25.00 42.45				
302	CA	ARG	54	139.802	22.461	26.9976	1.00	41.46				
303	C	ARG	54	140.516	21.152	26.653	1.00	43.94				
304	O	ARG	54	139.891	20.088	26.607	1.00	41.93				
305	CB	ARG	54	139.731	23.344	25.727	1.00	43.17				
306	CG	ARG	54	138.759	22.861	24.658	1.00	49.52				
307 308	CD NE	ARG ARG	54 54	138.792 138.600	23.763 25.176	23.428 23.764	$\frac{1.00}{1.00}$	55.06 65.13				
309	CZ	ARG	54	137.416	25.758	23.764	1.00	65.13 73.50				
310	NH1	ARG	54	136.293	25.058	23.836	1.00	78.69				
311	NH2	ARG	54	137.353	27.046	24.263	1.00	72.72				
312	H	ARG	54	140.919	24.030	27.846	1.00	25.00				
313	HE	ARG	54	139.400	25.734	23.857	1.00	25.00				
314	1HH1	ARG	54	136.328	24.086	23.604	1.00	25.00				
315 316	2HH1 1HH2	ARG ARG	54 54	135.410 138.196	25.505 27.578	23.976 24.357	$\frac{1.00}{1.00}$	25.00 25.00				
317	2HH2	ARG	54	136.466	27.484	24.403	1.00	25.00				
318	N	SER	55	141.834	21.233	26.480	1.00	41.53				
319	CA	SER	55	142.645	20.062	26.176	1.00	41.04				
320	С	SER	55	142.550	19.010	27.284	1.00	42.34				
321	O	SER	55	142.587	17.810	27.005	1.00	42.66				
322	CB	SER	55	144.100	20.469	25.916	1.00	46.50				
323	OG	SER	55 55	144.208	21.230	24.719	1.00	41.12				
324 325	H HG	SER SER	55 55	142.280 143.898	22.107 20.704	26.538 23.977	$\frac{1.00}{1.00}$	25.00 25.00				
326	N	MET	56	142.399	19.458	28.531	1.00	41.33				
327	CA	MET	56	142.265	18.544	29.668	1.00	40.87				
328	C	MET	56	141.003	17.703	29.501	1.00	43.46				
329	O	MET	56	141.017	16.489	29.711	1.00	41.72				
330	CB	MET	56	142.164	19.312	30.989	1.00	44.45				
331	CG	MET	56	143.461	19.894	31.500	1.00	46.36				
332 333	SD CE	MET MET	56 56	143.231 143.715	20.659 22.320	33.118 32.763	1.00 1.00	45.60 44.30				
334	H	MET	56	142.383	20.426	28.690	1.00	25.00				
335	N	LEU	57	139.910	18.365	29.132	1.00	42.82				
336	CA	LEU	57	138.632	17.696	28.929	1.00	41.38				
337	C	LEU	57	138.684	16.700	27.779	1.00	43.36				
338	O	LEU	57	138.042	15.653	27.831	1.00	45.75				
339 340	CB CG	LEU LEU	57 57	137.531 137.047	18.726	28.670 29.887	$\frac{1.00}{1.00}$	34.91				
341	CD1	LEU	57	136.174	19.512 20.663	29.441	1.00	31.36 31.29				
342	CD2	LEU	57	136.287	18.592	30.833	1.00	25.33				
343	H	LEU	57	139.968	19.335	28.990	1.00	25.00				
344	N	LEU	58	139.466	17.016	26.754	1.00	43.66				
345	CA	LEU	58	139.577	16.145	25.591	1.00	48.27				
346	С	LEU	58	140.659	15.059	25.683	1.00	53.11				
347 348	O CB	LEU LEU	58 58	141.005 139.762	14.441 16.987	24.672 24.325	$\frac{1.00}{1.00}$	54.87 45.38				
349	CG	LEU	58	138.682	18.051	24.083	1.00	48.84				
350	CD1	LEU	58	138.936	18.772	22.772	1.00	46.56				
351	CD2	LEU	58	137.303	17.413	24.074	1.00	49.64				
352	H	LEU	58	139.978	17.851	26.781	1.00	25.00				
353	N	ALA	59	141.180	14.813	26.884	1.00	58.17				
354	CA	ALA	59 50	142.208	13.788	27.078	1.00	61.28				
355 356	C O	ALA ALA	59 59	141.605 140.672	12.397 11.998	26.874 27.572	$\frac{1.00}{1.00}$	66.17 65.17				
357	СВ	ALA	59 59	142.830	13.908	28.460	1.00	61.08				
358	Н	ALA	59	140.859	15.316	27.661	1.00	25.00				
359	N	THR	60	142.188	11.651	25.943	1.00	71.93				
360	CA	THR	60	141.717	10.317	25.572	1.00	78.50				
361	C	THR	60	141.721	9.179	26.599	1.00	77.85				
362	O	THR	60	140.694	8.536	26.812	1.00	82.48				
363	CB CG1	THR	60 60	142.443	9.828 10.053	24.304	1.00	80.20				
364 365	CG1 CG2	THR THR	60 60	143.852 141.933	10.053	24.449 23.076	$\frac{1.00}{1.00}$	82.38 82.60				
366	H	THR	60	142.957	12.026	25.478	1.00	25.00				
367	HG1	THR	60	144.029	10.991	24.540	1.00	25.00				
368	N	GLY	61	142.866	8.914	27.217	1.00	74.94				
369	CA	GLY	61	142.942	7.813	28.165	1.00	75.87				
370	С	GLY	61	142.662	8.104	29.626	1.00	73.64				

TABLE 11-continued

		In th	e Absence of			сис Бунин		
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
371	O	GLY	61	143.494	7.804	30.484	1.00	73.89
372	H	GLY	61	143.654	9.465	27.047	1.00	25.00
373 374	N C4	ARG	62 62	141.491	8.651	29.925 31.307	1.00	70.85 67.41
374 375	CA C	ARG ARG	62 62	141.149 140.068	8.960 8.054	31.870	$\frac{1.00}{1.00}$	61.77
376	o	ARG	62	139.147	7.654	31.160	1.00	63.43
377	СВ	ARG	62	140.755	10.429	31.444	1.00	69.30
378	CG	ARG	62	141.883	11.309	31.967	1.00	75.81
379	CD	ARG	62	141.666	12.780	31.647	1.00	78.22
380	NE	ARG	62	140.334	13.254	32.009	1.00	79.69
381	CZ	ARG	62	139.335	13.405	31.143	1.00	88.12
382	NH1	ARG	62	139.508	13.118	29.859	1.00	90.62
383	NH2	ARG	62	138.160	13.852	31.559	1.00	92.99
384	H	ARG	62	140.835	8.835	29.218	1.00	25.00
385	HE	ARG	62	140.163	13.477	32.948	1.00	25.00
386	1HH1	ARG	62	140.393	12.785	29.534	1.00	25.00
387	2HH1	ARG	62	138.752	13.233	29.216	1.00	25.00
388	1HH2	ARG	62	138.022	14.076	32.524	1.00	25.00
389 390	2HH2	ARG	62 63	137.408	13.963 7.702	30.909	1.00	25.00
390 391	N CA	LYS LYS	63	140.214 139.258	6.840	33.143 33.830	$\frac{1.00}{1.00}$	56.71 53.88
392	C	LYS	63	137.986	7.614	34.170	1.00	49.91
393	Ö	LYS	63	138.024	8.831	34.377	1.00	43.13
394	СВ	LYS	63	139.876	6.284	35.114	1.00	60.17
395	CG	LYS	63	141.181	5.544	34.901	1.00	72.02
396	CD	LYS	63	141.807	5.141	36.225	1.00	82.26
397	CE	LYS	63	143.131	4.428	36.004	1.00	90.54
398	NZ	LYS	63	143.764	4.015	37.286	1.00	94.67
399	H	LYS	63	140.988	8.042	33.633	1.00	25.00
400	1HZ	LYS	63	143.130	3.366	37.795	1.00	25.00
401	2HZ	LYS	63	144.664	3.533	37.085	1.00	25.00
402	3HZ	LYS	63	143.945	4.857	37.868	1.00	25.00
403	N	LEU	64	136.877	6.890	34.289	1.00	44.28
404	CA	LEU	64	135.583	7.487	34.603	1.00	40.75
405	C	LEU	64	135.650	8.425	35.805	1.00	38.00
406	O	LEU	64	135.273	9.592	35.708	1.00	38.34
407	CB	LEU	64	134.539	6.395	34.858	1.00	37.20
408	CG CD1	LEU	64	133.128	6.897	35.170	1.00	35.50
409 410	CD1 CD2	LEU LEU	64 64	132.563 132.232	7.618 5.741	33.964 35.572	$\frac{1.00}{1.00}$	30.93 32.83
411	H	LEU	64	136.930	5.926	34.145	1.00	25.00
412	N	ALA	65	136.149	7.915	36.927	1.00	34.87
413	CA	ALA	65	136.264	8.703	33.152	1.00	33.63
414	C	ALA	66	136.977	10.022	37.909	1.00	32.94
415	O	ALA	65	136.508	11.073	38.342	1.00	31.36
416	CB	ALA	65	136.992	7.900	39.222	1.00	26.54
417	H	ALA	65	136.437	3.983	36.932	1.00	25.00
418	N	ASP	66	138.094	9.965	37.188	1.00	35.39
419	CA	ASP	66	138.887	11.151	36.882	1.00	34.93
420	С	ASP	66	138.127	12.136	36.002	1.00	34.90
421	O	ASP	66	138.200	13.352	36.210	1.00	34.81
422	CB	ASP	66	140.202	13.755	36.202	1.00	42.66
423	CG OD1	ASP	66	141.054	9.825	37.059	1.00	51.98
424 425	OD1 OD2	ASP ASP	66 66	141.008 141.774	9.934 8.982	38.306 36.479	$\frac{1.00}{1.00}$	48.05 59.40
425 426	H	ASP	66	138.390	9.103	36.840	1.00	25.00
427	N	THR	67	137.400	11.607	35.023	1.00	32.76
428	CA	THR	67	136.617	12.433	34.110	1.00	29.98
429	C	THR	67	135.486	13.131	34.869	1.00	27.93
430	ŏ	THR	67	135.262	14.337	34.708	1.00	26.55
431	СВ	THR	67	136.033	11.582	32.963	1.00	33.36
432	CG1	THR	67	137.102	10.914	32.278	1.00	32.56
433	CG2	THR	67	135.272	12.460	31.972	1.00	24.21
434	H	THR	67	137.385	10.635	34.900	1.00	25.00
435	HG1	THR	67	136.746	10.375	31.566	1.00	25.00
436	N	LEU	68	134.806	12.382	35.730	1.00	24.35
437	CA	LEU	68	133.717	12.938	36.513	1.00	23.41
438	C	LEU	68	134.223	14.025	37.449	1.00	28.28
439	O	LEU	68	133.644	15.112	37.507	1.00	26.82
440	CB	LEU	68	133.004	11.842	37.301	1.00	24.43
441	CG	LEU	68	132.221	10.8411	36.447	1.00	32.03
442	CD1	LEU	68	131.651	9.744	37.330	1.00	23.28
443	CD2	LEU	68 68	131.112	11.556	35.680	1.00	27.61
444	H	LEU	68	135.049	11.442	35.844	1.00	25.00

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
445	N	ASN	69	135.323	13.750	38.147	1.00	26.79
446	CA	ASN	69	135.894	14.724	39.072	1.00	30.78
447	С	ASN	69	136.341	15.981	38.340	1.00	28.43
448 449	O CB	ASN ASN	69 69	136.165 137.061	17.092 14.125	38.837 39.867	$\frac{1.00}{1.00}$	30.31 40.14
450	CG	ASN	69	136.597	13.165	40.959	1.00	53.22
451	OD1	ASN	69	135.478	13.271	41.467	1.00	52.67
452	ND2	ASN	69	137.460	12.224	41.326	1.00	60.05
453	H	ASN	69	135.750	12.874	38.043	1.00	25.00
454	1HD2	ASN	69	137.165	11.608	42.027	1.00	25.00
455 456	2HD2 N	ASN LEU	69 70	138.335 136.884	12.185 15.813	40.893 37.140	1.00 $1.00$	25.00 26.00
457	CA	LEU	70	137.327	16.958	36.358	1.00	27.21
458	C	LEU	70	136.135	17.867	36.053	1.00	29.79
459	O	LEU	70	136.192	19.076	36.287	1.00	27.18
460	CB	LEU	70	137.990	16.498	35.058	1.00	23.13
461	CG	LEU	70	138.417	17.624	34.109	1.00	30.77
462	CD1	LEU	70 70	139.366	18.580	34.821	1.00	21.76
463 464	CD2 H	LEU LEU	70 70	139.062 136.995	17.045 14.908	32.860 36.777	$\frac{1.00}{1.00}$	27.07 25.00
465	N	ILE	71	135.053	17.272	35.553	1.00	30.05
466	CA	ILE	71	133.840	18.012	35.217	1.00	24.54
467	С	ILE	71	133.221	18.663	36.456	1.00	23.02
468	O	ILE	71	132.849	19.839	36.429	1.00	23.20
469	CB	ILE	71	132.809	17.095	34.516	1.00	26.68
470 471	CG1 CG2	ILE ILE	71 71	133.338 131.459	18.693 17.795	33.136 34.383	$\frac{1.00}{1.00}$	25.05 23.87
472	CD1	ILE	71	132.442	15.736	32.400	1.00	27.35
473	Н	ILE	71	135.073	16.300	35.409	1.00	25.00
474	N	ASP	72	133.140	17.910	37.546	1.00	19.38
475	CA	ASP	72	132.585	18.429	38.789	1.00	22.68
476	С	ASP	72	133.376	19.657	39.266	1.00	25.06
477 478	O	ASP	72 72	132.784	20.680	39.626 39.861	1.00	24.92
479	CB CG	ASP ASP	72	132.593 131.900	17.335 17.760	41.147	$\frac{1.00}{1.00}$	23.74 27.65
480	OD1	ASP	72	130.953	18.575	41.086	1.00	29.89
481	OD2	ASP	72	132.303	17.268	42.223	1.00	30.37
482	H	ASP	72	133.459	16.986	37.512	1.00	25.00
483	N	ILE	73	134.705	19.565	39.228	1.00	26.02
484	CA	ILE	73	135.589	20.656	39.654	1.00	21.86
485 488	C O	ILE ILE	73 73	135.431 135.270	21.918 23.019	38.797 39.329	1.00 $1.00$	23.60 25.70
487	СВ	ILE	73	137.075	20.198	39.671	1.00	20.21
488	CG1	ILE	73	137.245	19.066	40.684	1.00	22.84
489	CG2	ILE	73	137.992	21.351	40.058	1.00	17.62
490	CD1	ILE	73	138.659	18.513	40.788	1.00	53.34
491	H	ILE	73	135.108	18.732	38.902	1.00	25.00
492	N	ILE	74 74	135.450	21.755	37.476	1.00	22.39
493 494	CA C	ILE ILE	74 74	135.297 133.955	22.884 23.581	36.556 36.784	1.00 1.00	22.15 24.59
495	Ö	ILE	74	133.858	24.807	36.702	1.00	29.58
496	CB	ILE	74	135.415	22.426	35.079	1.00	24.24
497	CG1	ILE	74	136.835	21.909	34.811	1.00	25.85
498	CG2	ILE	74	135.071	23.571	34.132	1.00	19.61
499	CD1	ILE	74	137.054	21.340	33.420	1.00	23.74
500 501	H N	ILE GLU	74 75	135.571 132.925	20.851 22.797	37.109 37.083	1.00 $1.00$	25.00 23.15
502	CA	GLU	75	131.599	23.343	37.338	1.00	23.53
503	C	GLU	75	131.548	24.092	38.658	1.00	21.84
504	O	GLU	75	131.040	25.208	38.722	1.00	25.34
505	CB	GLU	75	130.550	22.237	37.342	1.00	26.71
506	CG	GLU	75 75	130.274	21.647	35.978	1.00	30.84
507 508	CD OF1	GLU	75 75	129.073	20.720	35.969	1.00	36.10
508 509	OE1 OE2	GLU GLU	75 75	128.644 128.559	20.253 20.460	37.051 34.865	$\frac{1.00}{1.00}$	29.89 31.50
510	H	GLU	75	133.056	21.822	37.122	1.00	25.00
511	N	ARG	76	132.060	23.474	39.717	1.00	19.68
512	CA	ARG	76	132.066	24.115	41.028	1.00	20.75
513	С	ARG	76	132.925	25.385	41.017	1.00	22.83
514	O	ARG	76 76	132.699	26.303	41.803	1.00	21.89
515 516	CB	ARG	76 76	132.581	23.148	42.091	1.00	16.07
516 517	CG CD	ARG ARG	76 76	131.653 132.331	21.989 21.061	42.411 43.395	1.00 $1.00$	20.22 21.08
517	NE	ARG	76 76	131.498	19.939	43.819	1.00	18.53
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TABLE 11-continued

In the Absence of Bound Substrate									
	Atom Type	Atom	Residue	Residue #	X	Y	z	OCC	B-factor
	519	CZ	ARG	76	130.847	19.891	44.977	1.00	24.53
	520	NH1	ARG	76	130.917	20.907	45.829	1.00	1 <b>Y</b> .48
	521 522	NH2 H	ARG	76 76	130.170 132.440	18.802 22.577	45.311 39.618	$\frac{1.00}{1.00}$	26.65 25.00
	523	п НЕ	ARG ARG	76 76	131.417	19.172	43.218	1.00	25.00
	524	1HH1	ARG	76	131.459	21.717	45.606	1.00	25.00
	525	2HH1	ARG	76	130.423	20.865	46.697	1.00	25.00
	526	1HH2	ARG	76	130.146	18.021	44.691	1.00	25.00
	527	2HH2	ARG	76	129.678	18.766	46.181	1.00	25.00
	528	N	LEU	77	133.913	25.425	40.126	1.00	22.29
	529 530	CA C	LEU	77 77	134.798	26.579 27.710	40.001 39.193	1.00 $1.00$	23.34 26.64
	531	Ö	LEU LEU	77	134.156 134.752	28.777	39.193	1.00	25.27
	532	СВ	LEU	77	136.131	26.167	39.372	1.00	18.34
	533	CG	LEU	77	137.076	25.352	40.258	1.00	18.25
	534	CD1	LEU	77	138.266	24.893	39.443	1.00	15.60
	535	CD2	LEU	77	137.531	26.182	41.459	1.00	17.00
	536	H	LEU	77	134.063	24.649	39.546	1.00	25.00
	537	N	GLY	78 78	132.958	27.455	38.668	1.00	24.42
	538 539	CA C	GLY GLY	78 78	132.228 132.741	28.464 28.807	37.914 36.531	$\frac{1.00}{1.00}$	20.32 20.16
	540	Ö	GLY	78	132.375	29.841	35.970	1.00	22.90
	541	H	GLY	78	132.553	26.576	38.793	1.00	25.00
	542	N	ILE	79	133.550	27.927	335.952	1.00	22.82
	543	CA	ILE	79	134.099	28.170	34.623	1.00	24.96
	544	C	ILE	79	133.577	27.204	33.560	1.00	29.01
	545	O	ILE	79 70	133.991	27.273	32.398	1.00	28.78
	546 547	CB CG1	ILE ILE	79 79	135.648 138.142	28.133 26.920	34.635 35.429	$\frac{1.00}{1.00}$	24.44 27.37
	548	CG2	ILE	79 79	138.195	29.426	35.210	1.00	25.45
	549	CD1	ILE	79	137.632	26.715	35.381	1.00	25.03
	550	H	ILE	79	133.782	27.105	36.435	1.00	25.00
	551	N	SER	80	132.629	26.347	33.935	1.00	27.52
	552	CA	SER	80	132.079	25.381	32.986	1.00	29.32
	553	С	SER	80	131.317	26.012	31.816	1.00	31.74
	554 555	O CB	SER SEER	80 80	131.187 131.205	25.391 24.338	30.761 33.694	1.00 $1.00$	34.93 26.24
	556	OG	SER	80	130.096	24.932	34.338	1.00	29.78
	557	Н	SER	80	132.308	26.358	34.857	1.00	25.00
	558	HG	SER	80	130.432	25.530	34.992	1.00	25.00
	559	N	TYR	81	130.869	27.258	31.966	1.00	25.77
	560	CA	TYR	81	130.134	27.914	30.887	1.00	23.28
	561	С	TYR	81 81	130.965	28.063	29.605	1.00	30.16
	562 563	O CB	TYR TYR	81	130.418 129.556	28.302 29.261	28.527 31.344	1.00 $1.00$	32.12 24.86
	564	CG	TYR	81	130.557	30.381	31.543	1.00	29.19
	565	CD1	TYR	81	131.260	30.519	32.740	1.00	27.27
	566	CD2	TYR	81	130.768	31.329	30.545	1.00	28.18
	567	CE1	TYR	81	132.148	31.575	32.935	1.00	29.22
	568	CE2	TYR	81	131.649	32.384	30.729	1.00	29.85
	569 570	CZ OH	TYR TYR	81 81	132.336 133.220	32.504 33.547	31.923 32.084	$\frac{1.00}{1.00}$	29.78 28.93
	571	Н	TYR	81	131.028	27.734	32.805	1.00	25.00
	572	HH	TYR	81	133.196	34.121	31.313	1.00	25.00
	573	N	HIS	82	132.284	27.904	29.727	1.00	32.51
	574	CA	HIS	82	133.194	27.991	28.581	1.00	29.34
	575 576	С	HIS	82	133.237	26.669	27.828	1.00	28.19
	576 577	O CB	HIS HIS	82 82	133.658 134.631	26.620 28.280	26.672 29.038	$\frac{1.00}{1.00}$	28.93 27.19
	578	CG	HIS	82	134.839	29.654	29.589	1.00	22.31
	579	ND1	HIS	82	134.702	30.793	28.825	1.00	24.56
	580	CD2	HIS	82	135.195	30.071	30.827	1.00	20.64
	581	CE1	HIS	82	134.964	31.853	29.568	1.00	22.72
	582	NE2	HIS	82	135.265	31.442	30.786	1.00	23.06
	583 594	H HD1	HIS	82	132.658	27.714	30.612	1.00	25.00
	584 585	HD1 HE2	HIS HIS	82 82	134.458 135.465	30.815 32.016	27.872 31.549	$\frac{1.00}{1.00}$	25.00 25.00
	586	N	PHE	83	132.820	25.596	28.493	1.00	26.84
	587	CA	PHE	83	132.878	24.266	27.903	1.00	33.10
	588	C	PHE	83	131.549	23.521	27.811	1.00	36.98
	589	0	PHE	83	131.511	22.296	27.973	1.00	34.23
	590 501	CB	PHE	83	133.895	23.426	28.683	1.00	32.26
	591 592	CG CD1	PHE PHE	83 83	135.171 136.138	24.159 24.336	28.985 27.998	1.00 $1.00$	36.23 36.38
	372	CD1	LIL	0.5	100.100	24.550	21.000	1.00	50.50

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
593	CD2	PHE	83	135.381	24.724	30.241	1.00	34.95
594	CE1	PHE	83	137.295	25.067	28.254	1.00	36.74
595	CE2	PHE	83	136.533	25.457	30.509	1.00	39.63
596	CZ	PHE	83	137.492	25.630	29.511	1.00	41.21
597 598	H N	PHE GLU	83 84	132.457 130.478	25.694 24.241	29.394 27.484	1.00 $1.00$	25.00 40.92
599	CA	GLU	84	129.146	23.641	27.365	1.00	46.14
600	C	GLU	84	129.159	22.422	26.431	1.00	42.13
601	Ö	GLU	84	128.753	21.325	26.819	1.00	38.24
602	CB	GLU	84	128.128	24.674	26.851	1.00	56.77
603	CG	GLU	84	128.042	25.978	27.660	1.00	74.38
604	CD	GLU	84	127.252	25.853	28.960	1.00	81.74
605	OE1	GLU	84	127.654	25.066	29.847	1.00	85.56
606	OE2	GLU	84	126.233	26.564	29.101	1.00	84.80
607	H	GLU	84	130.590	25.203	27.330	1.00	25.00
608	N	LYS	85 85	129.674	22.614	25.218	1.00	40.25
609	CA C	LYS	85 85	129.740	21.544	24.224	1.00 1.00	41.41 36.27
610 611	Ö	LYS LYS	85	130.590 130.138	20.348 19.204	24.663 24.595	1.00	35.16
612	СВ	LYS	85	130.138	22.085	22.890	1.00	46.40
613	CG	LYS	85	130.364	21.025	21.801	1.00	58.06
614	CD	LYS	85	131.176	21.498	20.605	1.00	70.05
615	CE	LYS	85	131.305	20.389	19.565	1.00	70.91
616	NZ	LYS	85	132.101	20.809	18.379	1.00	79.32
617	H	LYS	85	130.016	23.501	24.992	1.00	25.00
618	1HZ	LYS	85	133.059	21.080	18.678	1.00	25.00
619	2HZ	LYS	85	132.157	20.018	17.706	1.00	25.00
620	3HZ	LYS	85	131.636	21.619	17.922	1.00	25.00
621	N	GLU	86	131.812	20.620	25.115	1.00	37.17
622	CA	GLU	86	132.736	19.573	25.545	1.00	35.98
623 624	C O	GLU GLU	86 86	132.162 132.158	18.714 17.483	26.663 26.571	1.00 $1.00$	36.73 38.05
625	СВ	GLU	86	134.077	20.173	25.990	1.00	36.51
626	CG	GLU	86	134.938	20.773	24.866	1.00	40.91
627	CD	GLU	86	134.439	22.124	24.349	1.00	43.71
628	OE1	GLU	86	133.728	22.840	25.085	1.00	42.43
629	OE2	GLU	86	134.776	22.480	23.201	1.00	50.53
630	H	GLU	86	132.086	21.551	25.172	1.00	25.00
631	N	ILE	87	131.666	19.368	27.708	1.00	35.80
632	CA	ILE	87	131.092	18.662	28.845	1.00	30.66
633	C	ILE	87	129.871	17.844	28.428	1.00	32.98
634	O	ILE	87	129.692	16.711	28.887	1.00	32.50
635	CB CG1	ILE	87 87	130.739 132.027	19.640 20.253	29.986	1.00	30.27
636 637	CG1 CG2	ILE ILE	87 87	129.972	18.926	30.546 31.091	1.00 1.00	29.89 29.25
638	CD1	ILE	87	131.814	21.264	31.654	1.00	25.52
639	Н	ILE	87	131.681	20.348	27.719	1.00	25.00
640	N	ASP	88	129.054	18.393	27.534	1.00	32.95
641	CA	ASP	88	127.870	17.679	27.070	1.00	36.69
642	С	ASP	88	128.258	16.407	26.309	1.00	39.28
643	O	ASP	88	127.745	15.324	26.6601	1.00	40.13
644	CB	ASP	88	126.994	18.573	26.191	1.00	40.42
645	CG	ASP	88	125.682	17.901	25.800	1.00	49.09
646	OD1	ASP	88	124.874	17.586	26.702	1.00	48.29
647	OD2	ASP	88	125.464 129.245	17.677 19.291	24.590 27.185	1.00	57.24
648 649	H N	ASP GLU	88 89	129.245	16.532	25.359	1.00 $1.00$	25.00 38.92
650	CA	GLU	89	129.178	15.385	23.539	1.00	38.06
651	C	GLU	89	130.258	14.303	25.433	1.00	35.90
652	ŏ	GLU	89	130.077	13.115	25.168	1.00	39.91
653	CB	GLU	89	130.572	15.829	23.466	1.00	44.42
654	CG	GLU	89	129.871	16.622	22.379	1.00	61.34
655	CD	GLU	89	130.822	17.159	21.333	1.00	75.40
656	OE1	GLU	89	131.776	17.873	21.707	1.00	80.47
657	OE2	GLU	89	130.609	16.878	20.134	1.00	86.74
658	H	GLU	89	129.569	17.416	25.185	1.00	25.00
659	N	ILE	90	130.985	14.708	26.470	1.00	32.75
660	CA	ILE	90	131.619	13.749	27.368	1.00	31.62
661 662	C O	ILE	90 90	130.556	13.052	28.215	1.00	33.52
663	СВ	ILE ILE	90 90	130.580 132.646	11.830 14.427	28.376 28.302	1.00 $1.00$	35.53 31.23
664	CG1	ILE	90	133.815	14.427	27.485	1.00	32.25
665	CG2	ILE	90	133.153	13.431	29.340	1.00	23.52
666	CD1	ILE	90	134.794	15.802	28.300	1.00	27.90

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
667	H	ILE	90	131.105	15.670	26.630	1.00	25.00
668	N	LEU	91	129.617	13.828	28.749	1.00	33.39
669	CA	LEU	91	128.551	13.266	29.569	1.00	33.57
670	С	LEU	91	127.642	12.351	28.756	1.00	35.23
671 672	O CB	LEU LEU	91 91	127.145 127.741	11.346 14.373	29.269 30.244	$\frac{1.00}{1.00}$	32.55 30.86
673	CG	LEU	91	128.430	15.017	31.447	1.00	28.62
674	CD1	LEU	91	127.538	16.084	32.040	1.00	25.02
675	CD2	LEU	91	128.752	13.952	32.490	1.00	25.28
676	H	LEU	91	129.643	14.726	28.591	1.00	25.00
677	N	ASP	92	127.445	12.692	27.486	1.00	34.80
678	CA	ASP	92	126.620	11.889	26.595	1.00	37.65
079	С	ASP	92	127.273	10.516	26.446	1.00	38.72
680 681	O CB	ASP ASP	92 92	126.594 126.491	9.490 12.569	26.494 25.231	1.00 $1.00$	41.65 44.12
682	CG	ASP	92	125.426	11.931	24.358	1.00	48.79
683	OD1	ASP	92	124.235	12.268	24.531	1.00	49.33
684	OD2	ASP	92	125.781	11.098	23.498	1.00	52.21
685	H	ASP	92	127.856	13.512	27.149	1.00	25.00
686	N	GLN	93	128.595	10.499	26.286	1.00	40.62
687	CA	GLN	93	129.337	9.247	26.155	1.00	41.87
688	С	GLN	93 93	129.209	8.415	27.424	$\frac{1.00}{1.00}$	41.38
689 690	O CB	GLN GLN	93 93	129.038 130.817	7.198 9.504	27.356 25.883	1.00 $1.00$	44.29 47.84
691	CG	GLN	93	131.124	10.061	24.511	1.00	65.26
692	CD	GLN	93	132.618	10.230	24.286	1.00	76.60
693	OE1	GLN	93	133.402	9.308	24.532	1.00	78.42
694	NE2	GLN	93	133.023	11.413	23.829	1.00	78.28
695	H	GLN	93	129.082	11.351	26.254	1.00	25.00
696	1HE2	GLN	93	133.983	11.522	23.685	1.00	25.00
697 698	2HE2 N	GLN ILE	93 94	132.356 129.302	12.108 9.065	23.664 28.580	$\frac{1.00}{1.00}$	25.00 38.08
699	CA	ILE	94	129.302	8.360	29.851	1.00	38.56
700	C	ILE	94	127.783	7.763	30.011	1.00	37.25
701	O	ILE	94	127.631	6.623	30.464	1.00	40.45
702	CB	ILE	94	129.519	9.284	31.051	1.00	38.10
703	CG1	ILE	94	130.982	9.729	30.973	1.00	33.78
704 705	CG2	ILE	94	129.265	8.559	32.372	1.00	39.54
705 706	CD1 H	ILE ILE	94 94	131.426 129.455	10.590 10.035	32.131 28.576	$\frac{1.00}{1.00}$	28.73 25.00
707	N	TYR	95	126.769	8.527	29.616	1.00	35.79
708	CA	TYR	95	125.383	8.080	29.702	1.00	36.92
709	C	TYR	95	125.219	6.814	28.869	1.00	40.95
710	O	TYR	95	124.681	5.812	29.340	1.00	39.72
711	CB	TYR	95	124.438	9.170	29.176	1.00	31.04
712	CG CD1	TYR	95 95	122.969	8.799	29.213	1.00	35.51
713 714	CD1 CD2	TYR TYR	95 95	122.356 122.189	8.420 8.826	30.407 28.054	$\frac{1.00}{1.00}$	37.94 40.35
714	CE1	TYR	95	121.002	8.073	30.452	1.00	42.00
716	CE2	TYR	95	120.827	8.481	28.088	1.00	44.29
717	CZ	TYR	95	120.245	8.107	29.294	1.00	43.88
718	OH	TYR	95	118.912	7.763	29.351	1.00	49.08
719	H	TYR	95	126.959	9.421	29.265	1.00	25.00
720 721	HH	TYR	95 06	118.668	7.513	30.254	1.00	25.00
721 722	N CA	ASN ASN	96 96	125.744 125.664	6.861 5.749	27.649 26.711	$\frac{1.00}{1.00}$	42.27 45.67
723	C	ASN	96	126.430	4.484	27.088	1.00	53.96
724	Ō	ASN	96	125.949	3.383	26.831	1.00	58.48
725	CB	ASN	96	126.068	6.215	25.310	1.00	41.30
726	CG	ASN	96	125.004	7.072	24.656	1.00	45.95
727	OD1	ASN	96	123.922	6.590	24.339	1.00	50.40
728 729	ND2 H	ASN ASN	06 96	125.299 126.205	3.349 7.682	24.459 27.378	$\frac{1.00}{1.00}$	47.53 25.00
730	1HD2	ASN	96	124.816	8.910	24.040	1.00	25.00
731	2HD2	ASN	96	126.172	8.686	24.743	1.00	25.30
732	N	GLN	97	127.634	4.625	27.702	1.00	62.59
733	CA	GLN	97	128.397	3.453	28.081	1.00	69.96
734	С	GLN	97	127.898	2.701	29.320	1.00	74.06
735 736	O	GLN	97 97	128.255	1.540	29.521	1.00	71.27
736 737	CB CG	GLN GLN	97 97	129.885 130.227	3.804 4.801	28.219 29.315	$\frac{1.00}{1.00}$	74.21 82.49
738	CD	GLN	97	131.723	5.065	29.415	1.00	85.73
739	OE1	GLN	97	132.336	4.837	30.456	1.00	88.41
740	NE2	GLN	97	132.316	5.548	28.329	1.00	82.49

TABLE 11-continued

	In the Absence of Bound Substrate							
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
741	Н	GLN	97	127.943	5.524	27.899	1.00	25.00
742 743	1HE2	GLN	97 97	133.277	5.711	28.402	1.00	25.00
743 744	2HE2 N	GLN ASN	97 98	131.787 127.091	5.714 3.367	27.528 30.147	$\frac{1.00}{1.00}$	25.00 83.07
745	CA	ASN	98	126.517	2.769	31.360	1.00	95.02
746	C	ASN	98	127.459	1.854	32.148	1.00	101.68
747	O	ASN	98	127.088	0.737	32.517	1.00	106.47
748	CB	ASN	98	125.233	1.997	31.019	1.00	98.06
749	CG OD1	ASN	98	124.034	2.902	30.830	1.00	100.62
750 751	OD1 ND2	ASN ASN	98 98	123.390 123.711	3.307 3.207	31.799 29.580	1.00 1.00	107.25 100.51
751 752	H	ASN	98	126.876	4.301	29.938	1.00	25.00
753	1HD2	ASN	98	122.944	3.798	29.449	1.00	25.00
754	2HD2	ASN	98	124.241	2.849	28.838	1.00	25.00
755	N	SER	99	128.671	2.326	32.415	1.00	105.86
756	CA	SER	99	129.649	1.533	33.153	1.00	108.38
757	С	SER	999 99	129.389	1.528	34.659	1.00	109.53
758 759	O CB	SER SER	99 99	129.054 131.057	2.561 2.049	35.243 32.859	$\frac{1.00}{1.00}$	110.32 107.06
760	OG	SER	99	131.078	3.465	32.861	1.00	109.07
761	H	SER	99	128.927	3.224	32.122	1.00	25.00
762	HG	SER	99	130.810	3.805	33.720	1.00	25.00
763	N	ASN	100	129.534	0.361	35.280	1.00	111.83
764	CA	ASN	100	129.325	0.223	36.720	1.00	115.28
765	С	ASN	100	130.612	0.548	37.462	1.00	114.16
766 767	O CB	ASN ASN	100 100	131.577 128.873	-0.213 -1.195	37.402 37.072	$\frac{1.00}{1.00}$	114.16 117.88
768	CG	ASN	100	127.451	-1.473	36.640	1.00	122.11
769	OD1	ASN	100	126.518	-0.785	37.056	1.00	119.89
770	ND2	ASN	100	127.274	-2.485	35.799	1.00	124.85
771	H	ASN	100	129.794	-0.423	34.757	1.00	25.00
772 773	1HD2 2HD2	ASN ASN	100 100	126.354 128.048	-2.663 -3.001	35.510 35.500	$\frac{1.00}{1.00}$	25.00 25.00
774	N	CYS	100	130.622	1.678	38.162	1.00	112.97
775	CA	CYS	101	131.804	2.103	38.902	1.00	109.49
776	C	CYS	101	132.046	1.309	40.184	1.00	104.47
777	O	CYS	101	133.178	1.226	40.662	1.00	107.64
778	CB	CYS	101	131.735	3.593	39.218	1.00	112.66
779 780	SG H	CYS CYS	101 101	133.273 129.822	4.224 2.244	39.908 38.153	$\frac{1.00}{1.00}$	125.35 25.00
781	N	ASN	102	130.976	0.765	40.756	1.00	95.01
782	CA	ASN	102	131.043	-0.0488	41.975	1.00	88.33
783	C	ASN	102	131.235	0.673	43.315	1.00	78.73
784	O	ASN	102	131.005	0.074	44.367	1.00	78.57
785 786	CB CG	ASN ASN	102 102	132.071 131.727	-1.182 -2.147	41.831 40.704	1.00 1.00	94.60 99.33
787	OD1	ASN	102	130.667	-2.774	40.704	1.00	99.33 97.79
788	ND2	ASN	102	132.618	-2.256	39.727	1.00	103.54
789	H	ASN	102	130.102	0.912	40.350	1.00	25.00
790	1HD2	ASN	102	132.412	-2.867	38.993	1.00	25.00
791	2HD2	ASN	102	133.439	-1.721	39.766	1.00	25.00
792 793	N CA	ASP ASP	103 103	131.684 131.845	1.926 2.657	43.301 44.560	$\frac{1.00}{1.00}$	65.57 56.22
794	C	ASP	103	130.870	3.833	44.638	1.00	46.49
795	ŏ	ASP	103	130.659	4.550	43.657	1.00	41.71
796	CB	ASP	103	133.296	3.102	44.796	1.00	56.91
797	CG	ASP	103	133.767	4.133	43.802	1.00	64.49
798	OD1	ASP	103	134.215	3.735	42.707	1.00	74.22
799 800	OD2 H	ASP ASP	103 103	133.707 131.912	5.339 2.367	44.124 42.461	$\frac{1.00}{1.00}$	67.65 25.00
801	N	LEU	103	130.281	4.015	45.818	1.00	38.89
802	CA	LEU	104	129.291	5.060	46.071	1.00	36.92
803	C	LEU	104	129.672	6.457	45.591	1.00	36.45
804	O	LEU	104	128.898	7.109	44.895	1.00	35.19
805	CB	LEU	104	128.943	5.092	47.561	1.00	33.67
806 807	CG CD1	LEU	104 104	127.824	6.031 5.713	48.011 47.269	1.00	36.34 35.38
807 808	CD1 CD2	LEU LEU	104 104	126.538 127.622	5.713 5.893	47.269	$\frac{1.00}{1.00}$	35.38 35.29
809	H	LEU	104	130.509	3.401	46.542	1.00	25.00
810	N	CYS	105	130.872	6.895	45.951	1.00	36.36
811	CA	CYS	105	131.376	8.212	45.581	1.00	35.11
812	С	CYS	105	131.220	8.537	44.092	1.00	34.06
813 814	O CB	CYS	105 105	130.596 132.847	9.536 8.325	43.725	1.00	37.28 35.03
814	CD	CYS	105	132.04/	8.325	45.993	1.00	35.03

TABLE 11-continued

		In the	Absence o					
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
815	SG	CYS	105	133.614	9.885	45.573	1.00	53.55
816 817	H N	CYS THR	105 106	131.437 131.761	6.307 7.679	46.484 43.236	$\frac{1.00}{1.00}$	25.00 30.74
818	CA	THR	106	131.701	7.890	41.797	1.00	28.00
819	C	THR	106	130.301	7.663	41.227	1.00	25.73
820	O	THR	106	129.870	8.396	40.339	1.00	28.92
821	CB	THR	106	132.714	7.000	41.074	1.00	33.61
822 823	OG1 CG2	THR	106 106	134.000 132.807	7.172 7.369	41.684 39.598	$\frac{1.00}{1.00}$	38.24 30.78
823 824	H	THR THR	106	132.208	6.874	43.564	1.00	25.00
825	HG1	THR	106	134.270	8.093	41.602	1.00	25.00
826	N	SER	107	129.592	6.670	41.751	1.00	23.29
827	CA	SER	107	128.237	6.371	41.294	1.00	27.37
828	С	SER	107	127.268	7.540	41.539	1.00	25.50
829 830	O CB	SER SER	107 107	126.518 127.721	7.932 5.101	40.643 41.978	$\frac{1.00}{1.00}$	25.99 26.16
831	OG	SER	107	128.552	3.993	41.676	1.00	34.57
832	Н	SER	107	129.981	6.113	42.456	1.00	25.00
833	HG	SER	107	128.501	3.864	40.726	1.00	25.00
834	N	ALA	108	127.298	8.096	42.749	1.00	23.90
835	CA	ALA	108	126.441 126.779	9.219	43.121	1.00	23.63
836 837	C O	ALA ALA	108 108	126.779	10.466 11.189	42.307 41.861	$\frac{1.00}{1.00}$	25.62 27.88
838	СВ	ALA	108	126.566	9.509	44.608	1.00	18.25
839	H	ALA	108	127.913	7.734	43.418	1.00	25.00
840	N	LEU	109	128.069	10.709	42.099	1.00	21.86
841	CA	LEU	109	128.493	11.861	41.322	1.00	21.96
842 843	C O	LEU	109 109	128.009	11.704	39.881 39.297	1.00	25.59
844	СВ	LEU LEU	109	127.458 130.017	12.640 12.002	41.359	$\frac{1.00}{1.00}$	26.33 21.29
845	CG	LEU	109	130.611	13.161	40.550	1.00	23.03
846	CD1	LEU	109	129.969	14.480	40.962	1.00	15.35
847	CD2	LEU	109	132.111	13.210	40.751	1.00	17.60
848	H	LEU	109	128.742	10.101	42.475	1.00	25.00
849 850	N CA	GLN GLN	110 110	128.205 127.796	10.509 10.199	39.325 37.954	$\frac{1.00}{1.00}$	27.12 28.41
851	C	GLN	110	126.302	10.449	37.803	1.00	24.05
852	O	GLN	110	125.849	11.049	36.825	1.00	23.84
853	CB	GLN	110	128.098	8.732	37.632	1.00	26.80
854	CG	GLN	110	127.790	8.333	38.197	1.00	34.89
855 856	CD OE1	GLN GLN	110 110	127.942 128.418	6.843 6.098	35.947 36.804	$\frac{1.00}{1.00}$	37.60 43.99
857	NE2	GLN	110	127.538	6.401	34.765	1.00	38.91
858	H	GLN	110	128.641	9.810	39.852	1.00	25.00
859	1HE2	GLN	110	127.636	5.440	34.604	1.00	25.00
860	2HE2	GLN	110	127.167	7.030	34.117	1.00	25.00
861 862	N CA	PHE PHE	111 111	125.543 124.104	9.970 10.140	38.779 38.783	$\frac{1.00}{1.00}$	20.86 24.95
863	C	PHE	111	123.760	11.633	38.792	1.00	24.87
864	O	PHE	111	123.037	12.113	37.917	1.00	27.29
865	CB	PHE	111	123.511	9.442	40.008	1.00	21.47
866	CG	PHE	111	122.019	9.568	40.120	1.00	28.99
867 868	CD1 CD2	PHE PHE	111 111	121.183 121.448	8.649 10.600	39.494 40.865	$\frac{1.00}{1.00}$	24.39 26.46
869	CE1	PHE	111	119.799	8.753	39.610	1.00	26.64
870	CE2	PHE	111	120.072	10.713	40.985	1.00	25.60
871	CZ	PHE	111	119.243	9.787	40.356	1.00	30.12
872	H	PHE	111	125.966	9.482	39.513	1.00	25.00
873 874	N CA	ARG ARG	112 112	124.323 124.055	12.372 13.802	39.747 39.858	$\frac{1.00}{1.00}$	23.80 18.05
875	C	ARG	112	124.033	14.598	38.601	1.00	22.05
876	ŏ	ARG	112	123.539	15.341	38.103	1.00	28.69
877	CB	ARG	112	124.771	14.417	41.066	1.00	18.30
878	CG	ARG	112	124.503	15.911	41.209	1.00	16.54
879	CD	ARG	112	125.077	16.519	42.479	1.00	17.13
880 881	NE CZ	ARG ARG	112 112	126.540 127.323	16.520 17.379	42.517 41.866	$\frac{1.00}{1.00}$	19.87 22.23
882	NH1	ARG	112	126.808	18.328	41.099	1.00	19.31
883	NH2	ARG	112	128.636	17.311	42.012	1.00	25.31
884	H	ARG	112	124.929	11.945	40.387	1.00	25.00
885	HE	ARG	112	126.982	15.846	43.063	1.00	25.00
886 887	1HH1 2HH1	ARG ARG	112 112	125.815 127.408	18.407	40.998 40.612	$\frac{1.00}{1.00}$	25.00 25.00
887 888	2HH1 1HH2	ARG	112	127.408	18.960 16.615	40.612	1.00	25.00
555			***		20.010	.2.000	2.00	

TABLE 11-continued

Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase
In the Absence of Bound Substrate

		In the	Absence of	f Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
889	2HH2	ARG	112	129.219	17.950	41.520	1.00	25.00
890	N	LEU	113	125.596	14.445	38.077	1.00	22.19
891	CA	LEU	113	125.994	15.192	36.883	1.00	23.19
892	C	LEU	113	125.112	14.907	35.665	1.00	27.49
893	O	LEU	113	124.752	15.828	34.921	1.00	24.19
894	CB	LEU	113	127.465	14.937	36.532	1.00	26.29
895	CG CD1	LEU	113	128.547	15.323	37.546	1.00	28.48
896 897	CD1 CD2	LEU LEU	113 113	129.911 128.391	15.110 16.772	36.905 37.993	$\frac{1.00}{1.00}$	21.83 17.87
898	H	LEU	113	126.225	13.821	38.492	1.00	25.00
899	N	LEU	114	124.776	13.638	35.451	1.00	24.90
900	CA	LEU	114	123.932	13.268	34.321	1.00	25.28
901	С	LEU	114	122.537	13.867	34.485	1.00	23.09
902	O	LEU	114	122.038	14.534	33.580	1.00	26.40
903	CB	LEU	114	123.866	11.746	34.168	1.00	23.58
904	CG	LEU	114	125.167	11.101	33.671	1.00	25.79
905	CD1	LEU	114	125.043	9.591	33.660	1.00	24.20
906	CD2	LEU	114	125.504	11.607	32.280	1.00	23.62
907	H	LEU	114	125.095	12.937	36.062	1.00	25.00
908	N	ARG	115	121.948	13.694	35.665	1.00	23.30
909 910	CA C	ARG ARG	115 115	120.620 120.551	14.228 15.748	35.955 35.787	1.00 1.00	21.07 26.37
910	Ö	ARG	115	119.628	16.267	35.148	1.00	26.34
912	СВ	ARG	115	120.178	13.844	37.372	1.00	20.95
913	CG	ARG	115	119.749	12.394	37.528	1.00	21.24
914	CD	ARG	115	118.588	12.057	36.595	1.00	24.51
915	NE	ARG	115	118.086	10.702	36.813	1.00	20.45
916	CZ	ARG	115	117.090	10.394	37.639	1.00	25.55
917	NH1	ARG	115	116.475	11.347	38.327	1.00	26.55
918	NH2	ARG	115	116.729	9.128	37.807	1.00	21.55
919	H	ARG	115	122.416	13.187	36.361	1.00	25.00
920	HE	ARG	115	118.508	9.978	36.314	1.00	25.00
921	1HH1	ARG	115	116.757	12.300	38.232	1.00	25.00
922	2HH1	ARG	115	115.725	11.114	38.942	1.00	25.00
923 924	1HH2 2HH2	ARG ARG	115 115	117.205 115.980	8.402 8.901	37.314 38.425	$\frac{1.00}{1.00}$	25.00 25.00
925	N	GLN	116	121.537	16.458	36.333	1.00	24.98
926	CA	GLN	116	121.573	17.917	36.235	1.00	21.36
927	C	GLN	116	121.696	18.366	34.792	1.00	23.11
928	O	GLN	116	121.331	19.491	34.450	1.00	20.94
929	CB	GLN	116	122.718	18.501	37.066	1.00	21.35
930	CG	GLN	116	122.536	18.322	38.561	1.00	24.02
931	CD	GLN	116	123.594	19.035	39.371	1.00	23.65
932	OE1	GLN	116	123.278	19.835	40.252	1.00	30.06
933	NE2	GLN	116	124.855	18.744	39.088	1.00	21.53
934	H	GLN	116	122.246	15.983	36.814	1.00	25.00
935	1HE2 2HE2	GLN GLN	116	125.538	19.210	39.609	1.00	25.00 25.00
936 937	N	HIS	116 117	125.048 122.232	18.089 17.490	38.391 33.950	$\frac{1.00}{1.00}$	23.00
938	CA	HIS	117	122.381	17.804	32.537	1.00	21.02
939	C	HIS	117	121.264	17.235	31.366	1.00	23.62
940	Õ	HIS	117	121.389	17.192	30.445	1.00	22.85
941	CB	HIS	117	123.755	17.366	32.031	1.00	23.58
942	CG	HIS	117	124.863	18.267	32.475	1.00	27.17
943	ND1	HIS	117	125.477	19.165	31.628	1.00	28.70
944	CD2	HIS	117	125.421	18.456	33.693	1.00	26.70
945	CE1	HIS	117	126.361	19.874	32.307	1.00	25.66
946	NE2	HIS	117	126.346	19.464	33.562	1.00	29.17
947	H	HIS	117	122.526	16.612	34.277	1.00	25.00
948	HD1	HIS	117	125.301	19.252	30.671	1.00	25.00
949 950	HE2	HIS GLY	117 118	126.861	19.825 16.784	34.298 32.301	$\frac{1.00}{1.00}$	25.00
951	N CA	GLY	118	120.183 119.050	16.258	31.562	1.00	24.12 25.68
952	C	GLY	118	119.037	14.786	31.193	1.00	30.13
953	ō	GLY	118	118.028	14.303	30.676	1.00	35.12
954	H	GLY	118	120.143	16.804	33.279	1.00	25.00
955	N	PHE	119	120.130	14.068	31.432	1.00	29.49
956	CA	PHE	119	120.184	12.644	31.102	1.00	26.30
957	C	PHE	119	119.435	11.867	32.172	1.00	28.96
958	O	PHE	119	119.836	11.861	33.337	1.00	25.84
959	CB	PHE	119	121.633	12.156	31.019	1.00	24.99
960	CG	PHE	119	122.447	12.845	29.964	1.00	23.43
961	CD1	PHE	119	122.392	12.421	28.640	1.00	22.02
962	CD2	PHE	119	123.267	13.922	30.291	1.00	23.36

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
963	CE1	PHE	119	123.145	13.063	27.650	1.00	26.82
964	CE2	PHE	119	124.023	14.572	29.312	1.00	20.73
965	CZ	PHE	119	123.962	14.141	27.988	1.00	22.11
966	H	PHE	119	120.902	14.496	31.856	1.00	25.00
967	N	ASN	120	118.351	11.207	31.779	1.00	31.42
968	CA	ASN	120	117.551	10.447	32.733	1.00	36.20
969	C	ASN	120	118.208	9.117	33.088	1.00	36.85
970	O	ASN	120	117.727	8.048	32.707	1.00	39.49
971	CB	ASN	120	116.126	10.232	32.200	1.00	33.41
972	CG	ASN	120	115.142	9.796	33.286	1.00	35.58
973	OD1	ASN	120	115.481	9.721	34.465	1.00	38.24
974	ND2	ASN	120	113.907	9.531	32.886	1.00	40.30
975	H	ASN	120	118.092	11.226	30.839	1.00	25.00
976	1HD2	ASN	120	113.277	9.246	33.576	1.00	25.00
977	2HD2	ASN	120	113.672	9.626	31.940	1.00	25.00
978	N	ILE	121	119.347	9.194	33.770	1.00	36.98
979	CA	ILE	121	120.054	7.997	34.192	1.00	29.59
980	С	ILE	121	119.207	7.323	35.264	1.00	29.55
981	O	ILE	121	118.647	7.984	36.134	1.00	30.63
982	СВ	ILE	121	121.478	8.319	34.745	1.00	35.93
983	CG1	ILE	121	122.130	7.046	35.300	1.00	34.57
984	CG2	ILE	121	121.419	9.425	35.7798	1.00	27.62
985	CD1	ILE	121	123.558	7.221	35.775	1.00	37.39
986	Н	ILE	121	119.696	10.074	34.013	1.00	25.00
987	N	SER	122	119.086	6.007	35.172	1.00	32.39
988	CA	SER	122	118.299	5.245	36.129	1.00	7.80
989	C	SER	122	118.912	5.243	37.526	1.00	27.11
990	ŏ	SER	122	120.130	5.143	37.685	1.00	30.59
991	СВ	SER	122	118.145	3.801	35.642	1.00	28.47
992	OG	SER	122	117.529	2.993	36.633	1.00	27.99
993	Н	SER	122	119.540	5.535	34.444	1.00	25.00
994	HG	SER	122	117.442	2.098	36.275	1.00	25.00
995	N	PRO	123	118.065	5.325	38.564	1.00	27.95
996	CA	PRO	123	118.542	5.323	39.949	1.00	27.26
997	C	PRO	123	118.941	3.904	40.372	1.00	33.55
998	Ö	PRO	123	119.325	3.664	41.521	1.00	34.51
999	СВ	PRO	123	117.323	5.823	40.723	1.00	26.86
1000	CG	PRO	123	116.184	5.252	39.938	1.00	26.05
1001	CD	PRO	123	116.605	5.531	38.510	1.00	24.97
1002	N	GLU	124	118.849	2.967	39.431	1.00	34.65
1003	CA	GLU	124	119.199	1.569	39.673	1.00	42.28
1004	C	GLU	124	120.673	1.441	40.056	1.00	39.18
1005	O	GLU	124	121.072	0.492	40.735	1.00	41.59
1006	СВ	GLU	124	118.902	0.732	38.424	1.00	48.21
1007	CG	GLU	124	119.074	-0.773	38.601	1.00	61.85
1008	CD	GLU	124	118.112	-1.379	39.615	1.00	71.29
1009	OE1	GLU	124	117.022	-0.803	39.851	1.00	70.67
1010	OE2	GLU	124	118.450	-2.447	40.170	1.00	76.93
1011	Н	GLU	124	118.522	3.203	38.539	1.00	25.00
1012	N	ILE	125	121.466	2.430	39.657	1.00	36.76
1013	CA	ILE	125	122.892	2.458	39.955	1.00	35.15
1014	C	ILE	125	123.155	2.401	41.472	1.00	35.83
1015	O	ILE	125	124.237	2.016	41.900	1.00	36.05
1016	CB	ILE	125	123.557	3.718	39.319	1.00	33.60
1017	CG1	ILE	125	125.082	3.610	39.368	1.00	32.56
1018	CG2	ILE	125	123.087	4.988	40.017	1.00	28.43
1019	CD1	ILE	125	125.789	4.705	38.586	1.00	28.58
1020	Н	ILE	125	121.080	3.169	39.141	1.00	25.00
1021	N	PHE	126	122.145	2.733	42.276	1.00	33.20
1022	CA	PHE	126	122.276	2.717	43.731	1.00	33.30
1023	C	PHE	126	121.902	1.39	444,402	1.00	38.46
1024	Ō	PHE	126	122.171	1.21	145.591	1.00	38.34
1025	СВ	PHE	126	121.444	3.84	344.362	1.00	29.75
1026	CG	PHE	126	121.937	5.21	844.030	1.00	32.61
1027	CD1	PHE	126	123.084	5.72	444.631	1.00	29.30
1028	CD2	PHE	126	121.263	6.00	743.103	1.00	33.58
1029	CE1	PHE	126	123.558	6.99	744.310	1.00	31.83
1030	CE2	PHE	126	121.726	7.27	942.775	1.00	36.03
1031	CZ	PHE	126	122.877	7.77	643.380	1.00	34.15
1031	H	PHE	126	121.286	3.00	441.891	1.00	25.00
1032	N	SER	127	121.285	0.477	43.662	1.00	37.94
1034	CA	SER	127	120.871	-0.806	44.236	1.00	37.24
1035	C	SER	127	122.012	-1.601	44.855	1.00	37.05
1036	Ö	SER	127	121.842	-2.210	45.908	1.00	37.94
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TABLE 11-continued

		In the	e Absence of	f Bound S	ubstrate			
Atom Typ	e Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
1037	CB	SER	127	120.141	-1.658	43.201	1.00	34.22
1038	OG	SER	127	118.885	-1.087	42.887	1.00	44.36
1039	H	SER	127	121.104	0.658	42.719	1.00	25.00
1040	HG	SER	127	119.018	-0.202	42.539	1.00	25.00
1041	N	LYS	128	123.184	-1.557	44.228	1.00	37.17
1042	CA	LYS	128	124.348	-2.280	44.737	1.00	39.88
1043	С	LYS	128	124.840	-1.783	46.097	1.00	42.69
1044 1045	O CB	LYS LYS	128 128	125.690 125.492	-2.417 -2.257	46.720 43.715	1.00 $1.00$	48.65 39.92
1045	CG	LYS	128	125.432	-0.882	43.210	1.00	41.79
1047	CD	LYS	128	126.864	-0.983	42.061	1.00	45.38
1048	CE	LYS	128	127.112	0.380	41.430	1.00	57.33
1049	NZ	LYS	128	128.057	0.329	40.278	1.00	63.77
1050	H	LYS	128	123.266	-1.028	43.408	1.00	25.00
1051	1HZ	LYS	128	128.971	-0.047	40.597	1.00	25.00
1052	2HZ	LYS	128	127.667	-0.293	39.541	1.00	25.00
1053	3HZ	LYS	128	128.187	1.285	39.892	1.00	25.00
1054	N	PHE	129	124.305	-0.656	46.556	1.00	41.03
1055	CA	PHE	129	124.697	-0.090	47.844	1.00	38.56
1056	С	PHE	129	123.574	-0.255	48.848	1.00	42.42
1057 1058	O CB	PHE PHE	129 129	123.817 125.013	0.319 1.396	49.940 47.695	1.00	44.74 32.52
1058	СG	PHE	129	125.984	1.691	46.604	$\frac{1.00}{1.00}$	28.29
1060	CD1	PHE	129	127.291	1.225	46.677	1.00	27.71
1061	CD2	PHE	129	125.585	2.402	45.481	1.00	27.23
1062	CE1	PHE	129	128.186	1.461	45.645	1.00	27.92
1063	CE2	PHE	129	126.473	2.644	44.442	1.00	29.82
1064	CZ	PHE	129	127.776	2.172	44.523	1.00	29.88
1065	H	PHE	129	123.617	-0.193	46.037	1.00	25.00
1066	N	GLN	130	122.566	-1.036	48.482	1.00	46.51
1067	CA	GLN	130	121.425	-1.242	49.356	1.00	52.21
1068	С	GLN	130	121.181	-2.700	49.659	1.00	60.08
1069	O	GLN	130	121.565	-3.588	48.891	1.00	57.60
1070	CB	GLN	130	120.173	-0.638	48.736	1.00	50.11
1071 1072	CG CD	GLN GLN	130 130	120.247 119.025	0.860 1.399	48.526 47.840	1.00 $1.00$	50.91 51.21
1072	OE1	GLN	130	118.339	0.677	47.117	1.00	52.53
1074	NE2	GLN	130	118.737	2.677	48.061	1.00	47.70
1075	Н	GLN	130	122.585	-1.533	47.632	1.00	25.00
1076	1HE2	GLN	130	117.922	2.996	47.623	1.00	25.00
1077	2HE2	GLN	130	119.296	3.221	48.627	1.00	25.00
1078	N	ASP	131	120.531	-2.944	50.790	1.00	67.42
1079	CA	ASP	131	120.236	4.306	51.203	1.00	74.82
1080	С	ASP	131	118.975	-4.74.6	50.421	1.00	79.83
1081	O	ASP	131	118.273	-3.905	49.822	1.00	81.73
1082	CB	ASP	131	120.046	-4.394	52.745	1.00	75.37
1083	CG OD1	ASP	131	118.894	-3.558	53.284	1.00	79.54
1084	OD1 OD2	ASP	131 131	118.173 118.707	-2.911 -3.540	52.501	1.00	89.69 80.32
1085 1086	H	ASP ASP	131	120.203	-2.147	54.511 51.259	$\frac{1.00}{1.00}$	25.00
1087	N	GLU	132	118.671	-6.041	50.521	1.00	86.92
1088	CA	GLU	132	117.492	-6.620	49.865	1.00	93.82
1089	C	GLU	132	116.183	-5.970	50.310	1.00	94.42
1090	O	GLU	132	115.084	-6.369	49.910	1.00	95.34
1091	CB	GLU	132	117.414	-8.108	50.165	1.00	99.58
1092	CG	GLU	132	118.603	-8.893	49.626	1.00	110.72
1093	CD	$\operatorname{GLU}$	132		-10.369	49.968	1.00	117.77
1094	OE1	GLU	132		-10.716	51.082	1.00	122.96
1095	OE2	GLU	132		-11.187	49.117	1.00	118.79
1096	H	GLU	132	119.347	-6.569	50.930	1.00	25.00
1097	N	ASN	133	116.318	4.957	51.145	1.00	94.58
1098 1099	CA C	ASN ASN	1333 133	115.214 115.107	4.208 -2.839	51.715 51.042	1.00 $1.00$	92.72 90.09
1100	o	ASN	133	114.134	-2.112	51.042	1.00	90.09
1100	СВ	ASN	133	115.492	4.043	53.214	1.00	99.96
1101	CG	ASN	133	114.389	-3.337	53.923	1.00	106.41
1103	OD1	ASN	133	113.275	-3.847	54.016	1.00	108.32
1104	ND2	ASN	133	114.683	-2.159	54.4.47	1.00	111.37
1105	H	ASN	133	117.175	4.667	51.437	1.00	25.00
1106	1HD2	ASN	133	113.925	-1.736	54.889	1.00	25.00
1107	2HD2	ASN	133	115.561	-1.741	54.365	1.00	25.00
1108	N	GLY	134	116.139	-2.501	50.284	1.00	84.16
1109	CA	GLY	134	116.195	-1.224	49.597	1.00	77.66
1110	С	GLY	134	116.752	-0.121	50.479	1.00	73.68

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TABLE 11-continued

Structural Coordinates of Tobacco 5-Epi-Aristolochene Synthase In the Absence of Bound Substrate Atom Type Atom Residue Residue # OCC B-factor 50.072 1111 O GLY 134 116.780 1.040 1.00 72.39 116.840 Н 134 -3.19250.233 1.00 25.00 1112 GLY 117.141 -0.4621.00 70.01 1113 N LYS 135 51.704 CA 117.724 0.524 LYS 135 52,606 1114 1.00 61.88

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113.779

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113.687

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119.834

121.268

122.075

121.797

121.686

121.382

122.171

120.282

121.876

119.976

120.771

119.315

123.046

123.910

124,551

125.408

125.007

124.526

125.683

125.266

126.388

123.197

127.197

126.377

126.670

124.151

124.688

126,219

126.855

124.049

122.561

122.276

122.994

121.317

123.462

126.807

128,255

128.960

130.144

128,600

127.596

126.266

127.548

128.225

128.751

128.861

129.454

127.821

127.643

126.330

128.819

127 290

128 295

128.288

129 646

129.713

127.565

127.860

130.719

0.556

-0.473

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-2.796

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3.043

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0.062

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1.292

-1.453

-1.846

-0.582

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52.776

53,629

54.656

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52.801

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51.049

50.199

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49.920

48.758

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56.390

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59.01

64.30

48.57

52.37

56.61

55.76

62.73

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25.00

61.30

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60.08

61.14

63.46

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67.92

72.44

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54.50

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53.27

59.61

67.84

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28.14

TABLE 11-continued

In the Absence of Bound Substrate								
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
1185	CA	SER	142	132.062	5.500	57.297	1.00	30.43
1186	C	SER	142	132.788	6.004	56.051	1.00	30.66
1187	O	SER	142	133.961	6.371	56.107	1.00	39.46
1188	CB	SER	142	132.879	4.409	58.011	1.00	30.29
1189	OG	SER	142	132.790	3.146	57.358	1.00	26.69
1190	H	SER	142	130.627	4.146	56.569	1.00	25.00
1191	HG	SER	142	133.417	2.568	57.780	1.00	25.00
1192	N	ASP	143	132.069	6.053	54.937	1.00	27.41
1193	CA	ASP	143	132.614	6.512	53.663	1.00	28.00
1194	С	ASP	143	132.168	7.966	53.447	1.00	33.02
1195	O	ASP	143	131.211	8.232	52.714	1.00	34.43
1196	CB	ASP	143	132.085	5.603	52.540	1.00	26.21
1197	CG OD1	ASP	143	132.609	5.978	51.160	1.00	32.37
1198	OD1	ASP	143 143	133.578	6.762 5.465	51.045 50.174	1.00	34.10
1199	OD2 H	ASP ASP	143	132.041 131.130	5.790	54.969	$\frac{1.00}{1.00}$	37.80 25.00
1200	N N	VAL	143		8.906	54.060	1.00	31.09
1201	CA		144	132.884 132.548		53.958		
1202 1203	C	VAL VAL	144	132.348	10.328 10.873	52.534	1.00	27.23 27.59
1203	o	VAL	144	131.404	11.545	52.334	$\frac{1.00}{1.00}$	25.91
1204	СВ	VAL VAL	144	133.541	11.204	54.758	1.00	27.72
1205	CG1	VAL	144	133.183	12.684	54.621	1.00	21.28
1200	CG2	VAL	144	133.509	10.804	56.227	1.00	34.69
1207	H	VAL	144	133.643	8.619	54.609	1.00	25.00
1209	N	LEU	145	133.344	10.580	51.649	1.00	25.30
1210	CA	LEU	145	133.266	11.063	50.268	1.00	27.68
1211	C	LEU	145	132.039	10.510	49.544	1.00	29.04
1211	o	LEU	145	131.392	11.218	48.773	1.00	26.76
1213	СВ	LEU	145	134.541	10.722	49.487	1.00	27.26
1214	CG	LEU	145	135.839	11.375	49.970	1.00	29.50
1215	CD1	LEU	145	136.956	11.087	48.983	1.00	25.10
1216	CD2	LEU	145	135.648	12.875	50.113	1.00	31.66
1217	H	LEU	145	134.098	10.040	51.931	1.00	25.00
1218	N	GLY	146	131.717	9.247	49.806	1.00	27.11
1219	CA	GLY	146	130.552	8.643	49.185	1.00	25.43
1220	C	GLY	146	129.288	9.290	49.726	1.00	27.86
1221	Ö	GLY	146	128.373	9.621	48.968	1.00	24.45
1222	H	GLY	146	132.255	8.727	50.431	1.00	25.00
1223	N	LEU	147	129.251	9.485	51.043	1.00	21.13
1224	CA	LEU	147	128.114	10.102	51.712	1.00	23.48
1225	C	LEU	147	127.867	11.519	51.202	1.00	23.64
1226	Ö	LEU	147	126.722	11.922	51.002	1.00	25.30
1227	СВ	LEU	147	128.338	10.140	53.226	1.00	23.15
1228	CG	LEU	147	128.286	8.821	54.003	1.00	30.78
1229	CD1	LEU	147	128.667	9.059	55.455	1.00	24.16
1230	CD2	LEU	147	126.892	8.210	53.911	1.00	22.86
1231	H	LEU	147	130.010	9.196	51.584	1.00	25.00
1232	N	LEU	148	128.943	12.265	50.978	1.00	20.29
1233	CA	LEU	148	128.831	13.633	50.498	1.00	24.04
1234	С	LEU	148	128.217	13.664	49.106	1.00	23.69
1235	Ō	LEU	148	127.267	14.408	48.855	1.00	26.51
1236	CB	LEU	148	130.198	14.328	50.506	1.00	22.43
1237	CG	LEU	148	130.240	15.787	50.033	1.00	24.68
1238	CD1	LEU	148	129.285	16.649	50.853	1.00	16.95
1239	CD2	LEU	148	131.662	16.314	50.136	1.00	19.49
1240	H	LEU	148	129.830	11.885	51.143	1.00	25.00
1241	N	ASN	149	128.742	12.845	48.203	1.00	23.38
1242	CA	ASN	149	128.210	12.801	46.850	1.00	20.71
1243	C	ASN	149	126.781	12.269	46.809	1.00	25.14
1244	O	ASN	149	125.990	12.678	45.956	1.00	25.34
1245	CB	ASN	149	129.125	12.008	45.932	1.00	15.21
1246	CG	ASN	149	130.320	12.817	45.489	1.00	19.96
1247	OD1	ASN	149	131.340	12.856	46.167	1.00	34.43
1248	ND2	ASN	149	130.135	13.505	44.369	1.00	25.72
1249	H	ASN	149	129.509	12.271	48.444	1.00	25.00
1250	1HD2	ASN	149	130.969	14.021	44.090	1.00	25.00
1251	2HD2	ASN	149	129.340	13.468	43.887	1.00	25.00
1252	N	LEU	150	126.445	11.379	47.743	1.00	21.91
	CA	LEU	150	125.096	10.829	47.827	1.00	24.64
1253		T 1777	150	124.171	11.938	48.330	1.00	22.13
	C	LEU	150					
1253 1254 1255	O	LEU	150	123.058	12.104	47.831	1.00	27.92
1253 1254 1255 1256	O CB	LEU LEU	150 150	123.058 125.051	12.104 9.630	48.780	1.00	17.59
1253 1254 1255	O	LEU	150	123.058	12.104			

TABLE 11-continued

	Siruc	In the	Absence o			ene Synti	nase	
Atom Type	Atom	Residue	Residue #	X	Y	Z	occ	B-factor
1259	CD2	LEU	150	123.739	7.976	50.118	1.00	19.18
1260	H	LEU	150	127.125	11.076	48.382	1.00	25.00
1261 1262	N CA	TYR TYR	151 151	124.652 123.892	12.706 13.818	49.301 49.858	$\frac{1.00}{1.00}$	20.18 20.96
1263	CA	TYR	151	123.532	14.798	48.738	1.00	19.39
1264	Ö	TYR	151	122.380	15.204	48.592	1.00	21.76
1265	CB	TYR	151	124.723	14.535	50.929	1.00	20.29
1266	CG	TYR	151	124.115	15.828	51.418	1.00	22.32
1267	CD1	TYR	151	123.202	15.834	52.470	1.00	20.29
1268	CD2	TYR	151	124.432	17.047	50.811	1.00	21.17
1269 1270	CE1 CE2	TYR	151	122.614 123.850	17.021	52.907	1.00 1.00	20.42 21.39
1270	CZ CZ	TYR TYR	151 151	123.830	18.236 18.214	51.237 52.285	1.00	22.07
1272	OH	TYR	151	122.337	19.377	52.696	1.00	21.54
1273	Н	TYR	151	125.542	12.522	49.655	1.00	25.00
1274	HH	TYR	151	121.769	19.210	53.457	1.00	25.00
12775	N	GLU	152	124.532	15.194	47.959	1.00	21.41
1276	CA	GLU	152	124.316	16.128	46.863	1.00	19.05
1277	С	GLU	152	123.388	15.540	45.800	1.00	23.34
1278	O CB	GLU	152 152	122.540	16.259 16.544	45.260 46.235	1.00	21.99 23.58
1279 1280	СG	GLU GLU	152	125.653 126.641	17.236	47.198	$\frac{1.00}{1.00}$	23.54
1281	CD	GLU	152	126.245	18.662	47.577	1.00	27.07
1282	OE1	GLU	152	125.046	19.009	47.529	1.00	30.35
1283	OE2	GLU	152	127.145	19.444	47.935	1.00	23.20
1284	H	GLU	152	125.434	14.851	48.136	1.00	25.00
1285	N	ALA	153	123.530	14.248	45.526	1.00	24.42
1286	CA	ALA	153	122.706	13.565	44.526	1.00	22.61
1287 1288	C O	ALA ALA	153 153	121.251 120.342	13.409 13.450	44.964 44.138	$\frac{1.00}{1.00}$	19.49 21.30
1289	СВ	ALA	153	120.342	12.203	44.186	1.00	21.47
1290	Н	ALA	153	124.211	13.731	46.006	1.00	25.00
1291	N	SER	154	121.026	13.262	46.264	1.00	16.33
1292	CA	SER	154	119.672	13.105	46.776	1.00	21.77
1293	C	SER	154	118.822	14.343	46.464	1.00	27.62
1294	O	SER	154	117.603	14.258	46.351	1.00	29.95
1295	CB	SER	154	119.688	12.820	48.288	1.00	15.72
1296 1297	OG H	SER SER	154 154	119.902 121.776	13.991 13.256	49.060 46.901	1.00 1.00	20.33 25.00
1298	HG	SER	154	119.193	14.609	48.898	1.00	25.00
1299	N	HIS	155	119.470	15.489	46.291	1.00	24.50
1300	CA	HIS	155	118.751	16.720	46.001	1.00	20.74
1301	С	HIS	155	118.320	16.907	44.552	1.00	22.52
1302	O	HIS	155	117.682	17.905	44.224	1.00	23.13
1303 1304	CB	HIS	155	119.543 119.439	17.929	46.487	1.00	19.93
1304	CG ND1	HIS HIS	155 155	120.456	18.154 17.843	47.961 48.838	$\frac{1.00}{1.00}$	14.77 20.63
1306	CD2	HIS	155	118.431	18.652	48.716	1.00	13.91
1307	CE1	HIS	155	120.080	18.142	50.069	1.00	21.57
1308	NE2	HIS	155	118.855	18.634	50.022	1.00	17.34
1309	H	HIS	155	120.451	15.494	46.346	1.00	25.00
1310	HD1	HIS	155	121.317	17.420	48.603	1.00	25.00
1311	HE2	HIS	155	118.336	18.952	50.793	1.00	25.00
1312 1313	N CA	VAL VAL	156 156	118.686 118.283	15.972 16.063	43.678 42.276	$\frac{1.00}{1.00}$	22.99 22.09
1314	C	VAL	156	117.265	14.970	41.940	1.00	22.99
1315	O	VAL	156	116.954	14.741	40.768	1.00	22.80
1316	CB	VAL	156	119.491	15.956	41.299	1.00	18.92
1317	CG1	VAL	156	120.541	16.999	41.636	1.00	20.34
1318	CG2	VAL	156	120.089	14.560	41.329	1.00	20.67
1319	H	VAL	156	119.229	15.207	43.966	1.00	25.00
1320 1321	N CA	ARG ARG	157 157	116.729 115.766	14.317 13.239	42.968 42.762	$\frac{1.00}{1.00}$	19.28 25.29
1321	CA	ARG	157	114.394	13.708	42.702	1.00	26.91
1323	Ö	ARG	157	113.988	14.850	42.498	1.00	27.10
1324	CB	ARG	157	115.625	12.380	44.024	1.00	19.93
1325	CG	ARG	157	114.831	13.011	45.1144		19.14
1326	CD	ARG	157	114.914	12.156	46.397	1.00	20.33
1327	NE CZ	ARG	157	114.069	12.674	47.473	1.00	30.46
1328 1329	CZ NH1	ARG ARG	157 157	114.373	13.717	48.242 48.071	1.00	36.78 39.31
1329	NH1 NH2	ARG ARG	157 157	115.515 113.523	14.371 14.119	48.071 49.176	1.00 $1.00$	39.31 36.74
1331	H	ARG	157	116.972	14.566	43.881	1.00	25.00
1332	HE	ARG	157	113.214	12.230	47.643	1.00	25.00

TABLE 11-continued

	Struct	In the	e Absence o			ene synth		
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
1333	1HH $1$	ARG	157	116.149	14.079	47.364	1.00	25.00
1334	2HH1	ARG	157	115.736	15.154	48.652	1.00	25.00
1335 1336	1HH2 2HH2	ARG ARG	157 157	112.655 113.751	13.643 14.905	49.305 49.753	$\frac{1.00}{1.00}$	25.00 25.00
1337	N	THR	158	113.731	12.813	41.569	1.00	30.13
1338	CA	THR	158	112.385	13.066	41.015	1.00	27.65
1339	С	THR	158	111.374	12.189	41.763	1.00	25.41
1340	O	THR	158	111.751	11.413	42.642	1.00	23.51
1341	CB	THR	158	112.350	12.703	39.513	1.00	24.84
1342	OG1	THR	158	112.630	11.307	39.355	1.00	27.71
1343 1344	CG2	THR	158 158	113.391 114.102	13.496	38.738 41.427	1.00 1.00	19.09 25.00
1344	H HG1	THR THR	158	111.995	11.937 10.771	39.817	1.00	25.00
1346	N	HIS	159	110.103	12.268	41.377	1.00	26.77
1347	CA	HIS	159	109.051	11.473	42.016	1.00	27.30
1348	C	HIS	159	109.196	9.971	41.741	1.00	33.58
1349	O	HIS	159	108.630	9.150	42.462	1.00	33.82
1350	CB	HIS	159	107.663	11.939	41.557	1.00	26.01
1351	CG	HIS	159	107.337	13.350	41.941	1.00	23.00
1352 1353	ND1 CD2	HIS HIS	159 159	106.999 107.311	13.711 14.490	43.226 41.210	$\frac{1.00}{1.00}$	24.86 18.70
1354	CE1	HIS	159	107.311	15.012	43.275	1.00	24.72
1355	NE2	HIS	159	106.966	15.509	42.064	1.00	23.55
1356	Н	HIS	159	109.879	12.878	40.646	1.00	25.00
1357	HD1	HIS	159	106.924	13.094	43.988	1.00	25.00
1358	HE2	HIS	159	106.880	16.460	41.845	1.00	25.00
1359	N	ALA	160	109.948	9.624	40.697	1.00	32.77
1360	CA	ALA	160	110.167	8.229	40.315	1.00	31.64
1361 1362	C 0	ALA ALA	160 160	111.364 111.509	7.581 6.361	41.009 41.002	$\frac{1.00}{1.00}$	36.31 37.53
1363	СВ	ALA	160	110.326	8.130	38.803	1.00	25.40
1364	Н	ALA	160	110.358	10.319	40.160	1.00	25.00
1365	N	ASP	161	112.217	8.401	41.612	1.00	40.41
1366	CA	ASP	161	113.415	7.904	42.281	1.00	40.26
1367	C	ASP	161	113.123	7.414	43.689	1.00	41.77
1368	O	ASP	161	113.634	7.937	44.678	1.00	41.48
1369 1370	CB CG	ASP ASP	161 161	114.508 114.959	8.976 9.354	42.291 40.898	$\frac{1.00}{1.00}$	34.22 34.94
1370	OD1	ASP	161	114.954	8.486	40.002	1.00	33.51
1372	OD2	ASP	161	115.319	10.532	40.697	1.00	32.35
1373	Н	ASP	161	12.014	9.350	41.683	1.00	25.00
1374	N	ASP	162	112.353	6.342	43.742	1.00	46.26
1375	CA	ASP	162	111.932	5.726	44.985	1.00	46.59
1376	С	ASP	162 162	113.108	5.156	45.760	1.00	44.37
1377 1378	O CB	ASP ASP	162	113.127 110.916	5.172 4.630	46.990 44.670	$\frac{1.00}{1.00}$	37.28 56.78
1379	CG	ASP	162	109.654	5.185	44.046	1.00	69.65
1380	OD1	ASP	162	108.899	5.870	44.766	1.00	67.08
1381	OD2	ASP	162	109.435	4.978	42.830	1.00	79.19
1382	H	ASP	162	112.114	5.949	42.869	1.00	25.00
1383	N	ILE	163	114.106	4.699	45.015	1.00	41.87
1384 1385	CA C	ILE ILE	163 163	115.314 116.093	4.112 5.124	45.575 46.426	$\frac{1.00}{1.00}$	43.77 42.36
1386	ŏ	ILE	163	116.764	4.757	47.385	1.00	45.56
1387	CB	ILE	163	116.200	3.561	44.433	1.00	47.25
1388	CG1	ILE	163	115.385	2.571	43.595	1.00	56.48
1389	CG2	ILE	163	117.433	2.870	44.986	1.00	49.93
1390	CD1	ILE	163	116.134	1.994	42.404	1.00	60.37
1391 1392	H	ILE	163 164	114.031 115.955	4.781 6.404	44.043 46.097	$\frac{1.00}{1.00}$	25.00
1392	N CA	LEU LEU	164	116.650	7.473	46.805	1.00	37.87 33.53
1394	C	LEU	164	115.828	8.132	47.897	1.00	32.57
1395	O	LEU	164	116.206	9.192	48.400	1.00	36.58
1396	CB	LEU	164	117.102	8.542	45.815	1.00	30.53
1397	CG	LEU	164	118.184	8.139	44.815	1.00	36.53
1398	CD1	LEU	164	118.416	9.266	43.820	1.00	26.74
1399 1400	CD2 H	LEU LEU	164 164	119.468 115.309	7.794 6.641	45.562 45.397	$\frac{1.00}{1.00}$	30.59 25.00
1400	н N	GLU	165	115.309	7.489	48.290	1.00	25.00 32.57
1402	CA	GLU	165	113.854	8.022	49.320	1.00	32.62
1403	C	GLU	165	114.537	8.326	50.655	1.00	35.56
1404	O	GLU	165	114.298	9.368	51.267	1.00	35.70
1405	CB	GLU	165	112.663	7.058	49.551	1.00	39.90
1406	CG	GLU	165	111.645	7.571	50.549	1.00	50.03

TABLE 11-continued

		In the	Absence of	f Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
1407	CD	GLU	165	111.021	8.886	50.115	1.00	59.83
1408	OE1	GLU	165	110.492	8.952	48.983	1.00	63.90
1409	OE2	GLU	165	111.069	9.862	50.899	1.00	58.42
1410	H	GLU	165	114.553	6.614	47.898	1.00	25.00
1411	N	ASP	166	115.411	7.431	51.091	1.00	37.81
1412	CA	ASP	166	116.079	7.609	52.369	1.00	43.12
1413	С	ASP	166	117.546	8.030	52.268	1.00	40.48
1414 1415	O CB	ASP ASP	166 166	118.274 115.899	8.039 6.352	53.262 53.234	1.00 $1.00$	40.91 51.99
1416	CG	ASP	166	114.532	6.303	53.902	1.00	67.29
1417	OD1	ASP	166	114.268	7.173	54.761	1.00	74.04
1418	OD2	ASP	166	113.712	5.423	53.552	1.00	74.06
1419	H	ASP	166	115.669	6.693	50.506	1.00	25.00
1420	N	ALA	167	117.939	8.459	51.075	1.00	34.07
1421	CA	ALA	167	119.298	8.902	50.806	1.00	30.06
1422	C	ALA	167	119.664	10.182	51.554	1.00	33.12
1423	O	ALA	167	120.759	10.292	52.103	1.00	34.41
1424	СВ	ALA	167	119.488	9.090	49.308	1.00	25.56
1425	H	ALA	167	117.260	8.486	50.375	1.00	25.00
1426	N	LEU	168	118.737	11.134	51.593	1.00	34.35
1427 1428	CA C	LEU LEU	168 168	118.975 119.184	12.403 12.226	52.268 53.764	$\frac{1.00}{1.00}$	29.26 32.06
1429	Ö	LEU	168	120.199	12.226	54.311	1.00	34.07
1430	СВ	LEU	168	117.820	13.381	52.024	1.00	25.73
1431	CG	LEU	168	117.980	14.767	52.671	1.00	29.40
1432	CD1	LEU	168	119.241	15.454	52.153	1.00	22.56
1433	CD2	LEU	168	116.765	15.635	52.397	1.00	28.30
1434	H	LEU	168	117.879	10.944	51.168	1.00	25.00
1435	N	ALA	169	118.224	11.593	54.425	1.00	32.55
1436	CA	ALA	169	118.317	11.372	55.865	1.00	37.53
1437	C	ALA	169	119.561	10.552	56.227	1.00	37.81
1438	0	ALA	169	120.273	10.873	57.185	1.00	38.90
1439	CB	ALA	1669	117.058	10.680	56.370	1.00	36.88
1440	H	ALA	169	117.444	11.265	53.938	1.00	25.00
1441 1442	N CA	PHE	170 170	119.830 120.976	9.520 8.640	55.429 55.635	$\frac{1.00}{1.00}$	30.70 29.14
1443	C	PHE PHE	170	122.296	9.413	55.592	1.00	31.93
1444	o	PHE	170	123.046	9.432	56.573	1.00	35.80
1445	СВ	PHE	170	120.978	7.538	54.569	1.00	27.52
1446	CG	PHE	170	122.093	6.538	54.719	1.00	29.21
1447	CD1	PHE	170	122.055	5.576	55.722	1.00	32.25
1448	CD2	PHE	170	123.178	6.553	53.850	1.00	31.23
1449	CE1	PHE	170	123.085	4.642	55.854	1.00	35.77
1450	CE2	PHE	170	124.213	5.624	53.974	1.00	28.29
1451	CZ	PHE	170	124.166	4.668	54.977	1.00	33.63
1452	H	PHE	170	119.237	9.336	54.671	1.00	25.00
1453	N CA	SER	171 171	122.572	10.054 10.817	54.460	1.00	29.96
1454 1455	CA C	SER SER	171	123.803 123.888	11.970	54.297 55.293	$\frac{1.00}{1.00}$	23.74 25.49
1456	Ö	SER	171	124.951	12.232	55.845	1.00	30.00
1457	CB	SER	171	123.927	11.333	52.860	1.00	25.16
1458	OG	SER	171	122.318	12.137	52.501	1.00	31.46
1459	H	SER	171	121.937	10.029	53.708	1.00	25.00
1460	HG	SER	171	122.754	12.902	53.078	1.00	25.00
1461	N	THR	172	122.761	12.625	55.557	1.00	25.72
1462	CA	THR	172	122.728	13.746	56.490	1.00	25.18
1463	С	THR	172	123.183	13.367	57.902	1.00	30.69
1464	O	THR	172	124.122	13.961	58.438	1.00	29.76
1465	CB	THR	172	121.311	14.390	56.574	1.00	24.33
1466	OG1	THR	172	120.958	14.958	55.307	1.00	19.73
1467	CG2	THR THR	172 172	121.282	15.499	57.620	$\frac{1.00}{1.00}$	16.92 25.00
1468 1469	H HG1	THR	172	121.932 120.938	12.351 14.264	55.113 54.640	1.00	25.00
1470	N	ILE	173	122.542	12.363	58.489	1.00	34.29
1471	CA	ILE	173	122.875	11.951	59.848	1.00	37.02
1472	C	ILE	173	124.319	11.488	60.017	1.00	31.30
1473	Ö	ILE	173	124.958	11.777	61.032	1.00	34.03
1474	СВ	ILE	173	121.894	10.870	60.384	1.00	42.89
1475	CG1	ILE	173	122.082	10.702	61.893	1.00	46.46
1476	CG2	ILE	173	122.115	9.539	59.673	1.00	43.32
1477	CD1	ILE	173	121.040	9.829	62.553	1.00	58.82
1478	H	ILE	173	121.833	11.884	58.002	1.00	25.00
1479	N	HIS	174	124.848	10.790	59.020	1.00	27.02
1480	CA	HIS	174	126.220	10.309	59.100	1.00	30.73

TABLE 11-continued

		In the		of Bound S		ene synth		
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
1481	C	HIS	174	127.251	11.412	58.870	1.00	30.72
1482	O	HIS	174	128.261	11.477 9.118	59.574	1.00	32.05 32.72
1483 1484	CB CG	HIS HIS	174 174	126.431 125.701	7.884	58.166 58.603	$\frac{1.00}{1.00}$	42.58
1485	ND1	HIS	174	125.701	7.418	59.902	1.00	43.81
1486	CD2	HIS	174	124.891	7.036	57.925	1.00	39.61
1487	CE1	HIS	174	124.981	6.339	60.005	1.00	38.39
1488	NE2	HIS	174	124.457	6.086	58.820	1.00	38.65
1489	H	HIS	174	124.304	10.608	58.221	1.00	25.00
1490	HD1	HIS	174	126.233	7.787	60.656	1.00	25.00
1491	HE2	HIS	174	123.858	5.338	58.611	1.00	25.00
1492 1493	N CA	LEU LEU	175 175	126.970 127.874	12.310 13.420	57.931 57.655	$\frac{1.00}{1.00}$	30.47 24.51
1494	C	LEU	175	127.926	14.333	58.880	1.00	23.90
1495	Ö	LEU	175	128.999	14.803	59.267	1.00	27.49
1496	СВ	LEU	175	127.429	14.193	56.408	1.00	18.74
1497	CG	LEU	175	127.687	13.517	55.054	1.00	19.75
1498	CD1	LEU	175	127.007	14.295	53.935	1.00	19.14
1499	CD2	LEU	175	129.187	13.404	54.789	1.00	13.66
1500	H	LEU	175	126.143	12.235	57.409	1.00	25.00
1501 1502	N	GLU	176 176	126.781	14.535	59.524	1.00	23.45
1502	CA C	GLU GLU	176 176	126.7221 127.596	15.374 14.788	60.717 61.814	$\frac{1.00}{1.00}$	29.31 29.58
1504	Ö	GLU	176	128.222	15.519	62.580	1.00	30.33
1505	СВ	GLU	176	125.292	15.477	61.247	1.00	28.86
1506	CG	GLU	176	124.338	16.265	60.381	1.00	41.02
1507	CD	GLU	176	122.976	16.431	61.032	1.00	50.96
1503	CE1	GLU	176	122.409	15.422	61.511	1.00	58.24
1509	OE2	GLU	176	122.474	17.574	61.069	1.00	53.02
1510	H	GLU	176	125.956	14.127	59.182	1.00	25.00
1511	N CA	SER	177 177	127.615 128.394	13.461	61.890 62.894	$\frac{1.00}{1.00}$	31.99 33.70
1512 1513	CA	SER SER	177	129.905	12.746 12.777	62.620	1.00	29.73
1514	Ö	SER	177	130.710	12.952	63.541	1.00	31.31
1515	СВ	SER	177	127.896	11.299	62.986	1.00	33.08
1516	OG	SER	177	128.446	10.626	64.103	1.00	42.66
1517	H	SER	177	127.077	12.944	61.259	1.00	25.00
1518	HG	SER	177	128.220	11.094	64.907	1.00	25.00
1519	N	ALA	178	130.283	12.652	61.352	1.00	26.67
1520	CA C	ALA	178 178	131.692	12.641	60.970 60.858	1.00 1.00	25.61 28.58
1521 1522	o	ALA ALA	178	132.351 133.540	14.013 14.162	61.153	1.00	23.60
1523	СВ	ALA	178	131.862	11.884	59.665	1.00	22.60
1524	H	ALA	178	129.599	12.566	60.656	1.00	25.00
1525	N	ALA	179	131.568	15.018	60.475	1.00	25.11
1526	CA	ALA	179	132.068	16.376	60.268	1.00	25.17
1527	C	ALA	179	133.071	16.983	61.254	1.00	25.37
1528	O CB	ALA ALA	179 179	134.141 130.903	17.430 17.340	60.844 60.044	1.00	25.58
1529 1530	Н	ALA	179	130.617	14.840	60.325	$\frac{1.00}{1.00}$	21.50 25.00
1531	N	PRO	180	132.771	16.963	62.564	1.00	27.61
1532	CA	PRO	180	133.680	17.541	63.565	1.00	28.57
1533	С	PRO	180	135.132	17.058	63.584	1.00	30.64
1534	O	PRO	180	135.994	17.724	64.155	1.00	37.22
1535	CB	PRO	180	132.988	17.206	64.889	1.00	25.80
1536	CG	PRO	180	131.540	17.118	64.518	1.00	31.06
1537 1538	CD N	PRO HIS	180 181	131.597 135.414	16.360 15.910	63.221 62.980	$\frac{1.00}{1.00}$	30.35 28.35
1539	CA	HIS	181	136.772	15.377	63.013	1.00	27.57
1540	C	HIS	181	137.470	15.237	61.672	1.00	26.99
1541	O	HIS	181	138.529	14.611	61.584	1.00	29.22
1542	CB	HIS	181	136.764	14.035	63.740	1.00	30.76
1543	CG	HIS	181	136.153	14.103	65.104	1.00	32.51
1544	ND1	HIS	181	134.893	13.619	65.379	1.00	34.64
1545	CD2	HIS	181	136.607	14.652	66.257	1.00	34.04
1548 1547	CE1	HIS	181	134.593	13.870	66.641 67.106	1.00	35.16
1547 1548	NE2 H	HIS HIS	181 181	135.615 134.717	14.495 15.429	67.196 62.478	$\frac{1.00}{1.00}$	38.60 25.00
1549	HD1	HIS	181	134.717	13.158	64.739	1.00	25.00
1550	HE2	HIS	181	135.666	14.802	68.128	1.00	25.00
1551	N	LEU	182	136.890	15.827	60.635	1.00	22.56
1552	CA	LEU	182	137.468	15.750	59.303	1.00	22.65
1553	С	LEU	182	138.532	16.821	59.103	1.00	24.98
1554	О	LEU	182	138.494	17.878	59.741	1.00	22.99

TABLE 11-continued

			In the	Absence of	Bound S	ubstrate			
Ato	от Туре	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
	1555	CB	LEU	182	136.372	15.900	58.243	1.00	25.05
	1556	CG	LEU	182	135.271	14.835	58.205	1.00	23.65
	1557	CD1	LEU	182	134.178	15.274	57.249	1.00	17.28
	1558	CD2	LEU	182	135.849	13.483	57.786	1.00	20.03
	1559 1560	H N	LEU LYS	182 183	136.072 139.494	16.351 16.528	60.762 58.236	1.00 $1.00$	25.00 22.16
	1561	CA	LYS	183	140.556	17.469	57.926	1.00	25.90
	1562	C	LYS	183	139.982	18.573	57.045	1.00	30.39
	1563	Ö	LYS	183	138.898	18.429	56.468	1.00	31.71
	1564	CB	LYS	183	141.696	16.767	57.183	1.00	27.62
	1565	CG	LYS	183	141.274	16.1225	55.871	1.00	37.10
	1566	CD	LYS	183	142.437	15.441	55.169	1.00	45.13
	1567	CE	LYS	183	141.974	14.764	53.885	1.00	50.33
	1568	NZ	LYS	183	143.088	14.040	53.210	1.00	57.79
	1569 1570	H 1HZ	LYS LYS	183 183	139.473 143.846	15.658 14.713	57.786 52.975	1.00 $1.00$	25.00 25.00
	1571	2HZ	LYS	183	143.463	13.311	53.852	1.00	25.00
	1572	3HZ	LYS	183	142.736	13.592	52.341	1.00	25.00
	1573	N	SER	184	140.714	19.674	56.944	1.00	27.77
	1574	CA	SER	184	140.304	20.802	56.122	1.00	28.80
	1575	C	SER	184	140.970	20.675	54.752	1.00	27.61
	1576	O	SER	184	142.084	20.158	54.645	1.00	26.37
	1577	CB	SER	184	140.702	22.109	56.805	1.00	28.03
	1578	OG	SEER	184	140.003	22.254	58.031	1.00	32.93
	1579 1580	H HG	SER SER	184 184	141.565 140.193	19.725 21.517	57.420 58.620	$\frac{1.00}{1.00}$	25.00 25.00
	1581	N	PRO	185	140.193	21.171	53.689	1.00	26.21
	1582	CA	PRO	185	139.003	21.834	53.680	1.00	23.45
	1583	C	PRO	185	137.767	20.926	53.597	1.00	24.54
	1584	O	PRO	185	136.638	21.425	53.589	1.00	23.22
	1585	CB	PRO	185	139.109	22.737	52.458	1.00	21.98
	1586	CG	PRO	186	139.858	21.876	51.503	1.00	21.03
	1587	CD	PRO	185	140.949	21.263	52.361	1.00	21.80
	1588	N	LEU	186	137.969	19.608	53.570	1.00	21.43
	1589	CA	LEU	186	136.852 135.780	18.666 18.964	53.483 54.522	1.00 $1.00$	21.42 22.24
	1590 1591	C O	LEU LEU	186 186	134.586	18.987	54.210	1.00	20.84
	1592	CB	LEU	186	137.331	17.220	53.654	1.00	23.63
	1593	CG	LEU	186	136.217	16.160	53.646	1.00	21.09
	1594	CD1	LEU	186	135.491	16.145	52.292	1.00	20.03
	1595	CD2	LEU	186	136.800	14.800	53.943	1.00	21.57
	1596	Н	LEU	186	138.882	19.259	53.605	1.00	25.00
	1597	N	ARG	187	136.221	19.208	55.751	1.00	16.73
	1598	CA	ARG	187	135.326	19.515	56.859	1.00	22.57
	1599 1600	C O	ARG ARG	187 187	134.427 133.225	20.723 20.693	56.564 56.848	1.00 $1.00$	27.49 26.35
	1601	СВ	ARG	187	136.146	19.777	58.117	1.00	17.71
	1602	CG	ARG	187	135.325	20.087	59.343	1.00	21.93
	1603	CD	ARG	187	136.235	20.478	60.483	1.00	31.75
	1604	NE	ARG	187	135.507	20.685	61.727	1.00	46.15
	1605	CZ	ARG	187	136.087	20.96	162.891	1.00	58.70
	1606	NH1	ARG	187		21.06	662.970	1.00	57.84
	1607	NH2	ARG	187	135.344		163.982	1.00	58.69
	1608	H	ARG	187	137.182		155.923	1.00	25.00
	1609 1610	HE 1HH1	ARG ARG	187 187	134.530 137.977	20.61 20.94	361.708 162.156	$\frac{1.00}{1.00}$	25.00 25.00
	1611	2HH1	ARG	187	137.843		563.848	1.00	25.00
	1612	1HH2	ARG	187	134.351	21.01	263.926	1.00	25.00
	1613	2HH2	ARG	187	135.779	21.31	664.858	1.00	25.00
	1614	N	GLU	188	135.010	21.78	256.001	1.00	28.09
	1615	CA	GLU	188	134.255	22.99	355.667	1.00	26.62
	1616	C	GLU	188	133.293	22.72	654.516	1.00	22.97
	1617	O	GLU	188		23.29	654.462	1.00	21.14
	1618 1619	CB CG	GLU GLU	188 188	135.192 135.934	24.15 24.76	355.305 856.482	$\frac{1.00}{1.00}$	24.01 32.71
	1620	CD	GLU	188 188	135.934	23.87	850.482 857.014	1.00	42.50
	1621	OE1	GLU	188	138.030	23.65	756.279	1.00	43.53
	1622	OE2	GLU	188	136.936	23.40	358.165	1.00	47.38
	1623	H	GLU	188	135.965	21.74	755.798	1.00	25.00
	1624	N	GLN	189	133.702	21.85	353.601	1.00	19.36
	1625	CA	GLN	189	132.872	21.49	652.460	1.00	20.62
	1626	C	GLN	189		20.72	852.927	1.00	22.47
	1627	O	GLN	189	130.522	21.01	052.483	1.00	25.58
	1628	СВ	GLN	189	133.672	20.66	251.461	1.00	17.31

TABLE 11-continued

	Struct	In the	e Absence o			iene Syntha	ase	
Atom Type	Atom	Residue	Residue #	X	Y	z	OCC	B-factor
1629	CG	GLN	189	132.915	20.35	950.187	1.00	24.12
1630	CD	GLN	189	133.796	19.78	049.104	1.00	25.67
1631	OE1	GLN	189	133.691	20.16	247.939	1.00	28.92
1632	NE2	GLN	189	134.666	18.85	049.477	1.00	28.68
1633 1634	H 1HE2	GLN GLN	189 189	134.590 135.235	21.44 18.48	553.695 048.773	$\frac{1.00}{1.00}$	25.00 25.00
1635	1HE2 2HE2	GLN	189	133.233	18.57	650.413	1.00	25.00
1636	N	VAL	190	131.833	19.78	353.846	1.00	22.03
1637	CA	VAL	190	130.734	18.98	354.388	1.00	22.50
1638	C	VAL	190	129.778	19.86	455.198	1.00	22.00
1639	O	VAL	190	128.565	19.84	654.977	1.00	26.49
1640	CB	VAL	190	131.255	17.80	855.274	1.00	18.21
1641	CG1	VAL	190	130.093	17.09	355.947	1.00	19.13
1642	CG2	VAL	190	132.037	16.81	554.422	1.00	13.74
1643	H	VAL	190	132.742	19.61	854.168	1.00	25.00
1644	N	THR	191	130.335	20.63	856.124	1.00	20.35
1645	CA	THR	191	129.555	21.54	156.967	1.00	23.43
1646	С	THR	191	128.733	22.50	456.116	1.00	23.79
1647	O	THR	191	127.564	22.77	256.410	1.00	27.12
1648 1649	CB OG1	THR THR	191 191	130.478 131.124	22.35 21.45	057.903 458.814	$\frac{1.00}{1.00}$	29.00 35.12
1650	CG2	THR	191	129.688	23.38	558.691	1.00	32.22
1651	H	THR	191	131.304	20.59	956.257	1.00	25.00
1652	HG1	THR	191	131.661	20.82	458.321	1.00	25.00
1653	N	HIS	192	129.345	23.01	555.054	1.00	22.27
1654	CA	HIS	192	128.658	23.93	554.168	1.00	24.21
1655	C	HIS	192	127.530	23.22	653.417	1.00	24.78
1656	O	HIS	192	126.421	23.75	653.326	1.00	20.41
1657	CB	HIS	192	129.632	24.564	53.173	1.00	17.98
1658	CG	HIS	192	128.965	25.446	52.169	1.00	21.55
1659	ND1	HIS	192	128.506	26.707	52.480	1.00	21.86
1660	CD2	HIS	192	128.637	25.234	50.872	1.00	20.40
1661 1662	CE1 NE2	HIS HIS	192 192	127.919 127.985	27.234 26.360	51.420 50.432	1.00 1.00	20.03 20.23
1663	H H	HIS	192	130.278	22.766	54.870	1.00	25.00
1664	HD1	HIS	192	128.594	27.143	53.355	1.00	25.00
1665	HE2	HIS	192	127.614	26.486	49.551	1.00	25.00
1666	N	ALA	193	127.826	22.038	52.8888	1.00	22.45
1667	CA	ALA	193	126.854	21.242	52.139	1.00	20.56
1668	С	ALA	193	125.601	20.963	52.964	1.00	22.26
1669	O	ALA	193	124.485	21.072	52.459	1.00	21.49
1670	CB	ALA	193	127.483	19.938	51.679	1.00	18.73
1671	H	ALA	193	128.729	21.672	53.002	1.00	25.00
1672	N CA	LEU	194 194	125.791	20.623	54.236	$\frac{1.00}{1.00}$	23.86
1673 1674	CA C	LEU LEU	194 194	124.678 123.757	20.344 21.551	55.136 55.298	1.00	25.16 26.76
1675	o	LEU	194	122.573	21.391	55.579	1.00	28.61
1676	ČВ	LEU	194	125.194	19.902	56.509	1.00	23.10
1677	CG	LEU	194	125.924	18.556	56.579	1.00	26.60
1678	CD1	LEU	194	126.426	18.319	57.992	1.00	20.06
1679	CD2	LEU	194	124.998	17.433	56.149	1.00	18.24
1680	H	LEU	194	126.710	20.552	54.574	1.00	25.00
1681	N	GLU	195	124.309	22.754	55.149	1.00	33.01
1682	CA	GLU	195	123.529	23.987	55.277	1.00	34.61
1683	C	GLU	195	123.005	24.448	53.923	1.00	26.09
1684	O	GLU	195	121.952	25.074	53.834	1.00	29.66
1685	CB	GLU	195	124.385	25.102	55.884 57.288	1.00	38.35
1686 1687	CG CD	GLU GLU	195 195	124.885 125.945	24.816 25.803	57.751	$\frac{1.00}{1.00}$	59.66 72.22
1688	OE1	GLU	195	126.800	26.203	56.927	1.00	75.42
1689	OE2	GLU	195	125.931	26.169	58.947	1.00	82.70
1690	H	GLU	195	125.263	22.814	54.944	1.00	25.00
1691	N	GLN	196	123.747	24.130	52.871	1.00	22.78
1692	CA	GLN	196	123.376	24.529	51.527	1.00	20.33
1693	С	GLN	196	123.891	23.520	50.515	1.00	19.79
1694	O	GLN	196	125.094	23.463	50.258	1.00	24.78
1695	CB	GLN	196	123.980	25.903	51.219	1.00	20.38
1696	CG	GLN	196	123.727	26.397	49.807	1.00	24.47
1697	CD OF1	GLN	196	122.253	26.591	49.524	1.00	28.74
1698	OE1	GLN	196	121.622	27.490	50.074	1.00	32.30
1699 1700	NE2 H	GLN GLN	196 196	121.694 124.554	25.744 23.592	48.667 52.995	$\frac{1.00}{1.00}$	21.06 25.00
1704	1HE2	GLN	196	124.334	25.876	48.480	1.00	25.00
1702	2HE2	GLN	196	122.247	25.045	48.258	1.00	25.00
2.32							2.50	

TABLE 11-continued

		In the	e Absence o	f Bound St	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
1703	N	CYS	197	122.992	22.727	49.942	1.00	19.52
1704 1705	CA C	CYS CYS	197 197	123.399 123.782	21.749 22.497	48.944	$\frac{1.00}{1.00}$	17.94
1705	0	CYS	197	123.782	23.614	47.669 47.428	1.00	18.82 19.62
1707	СВ	CYS	197	122.278	20.743	48.669	1.00	21.42
1707	SG	CYS	197	120.832	21.394	47.800	1.00	42.82
1709	Н	CYS	197		22.808	50.202	1.00	25.00
1710	N	LEU	198		21.878	46.856	1.00	19.01
1711	CA	LEU	198		22.489	45.620	1.00	20.23
1712	C	LEU	198	123.986	22.760	44.610	1.00	23.16
1713	Ö	LEU	198	123.868	23.867	44.096	1.00	26.68
1714	СВ	LEU	198	126.174		44.981	1.00	13.53
1715	CG	LEU	198	126.762	22.058	43.640	1.00	20.37
1716	CD1	LEU	198	127.386	23.440	43.773	1.00	19.35
1717	CD2	LEU	198	127.789	21.044	43.158	1.00	18.28
1718	H	LEU	198	124.968	21.004	47.109	1.00	25.00
1719	N	HIS	199	123.160	21.752	44.354	1.00	24.27
1720	CA	HIS	199		21.859	43.379	1.00	22.48
1721	С	HIS	199		23.001	43.608	1.00	19.15
1722	O	HIS	199	120.586	23.582	42.653	1.00	19.66
1723	CB	HIS	199	121.327	20.523	43.277	1.00	22.51
1724	CG	HIS	199	120.225	20.519	42.261	1.00	16.72
1725	ND1	HIS	199	120.452	20.697	40.911	1.00	19.56
1726	CD2	HIS	199	118.885	20.360	42.396	1.00	15.14
1727	CE1	HIS	199		20.648	40.261	1.00	15.68
1728	NE2	HIS	199	118.338	20.444	41.138	1.00	19.96
1729 1730	H	HIS	199	123.297	20.912	44.822	1.00	25.00
	HD1 HE2	HIS	199 199	121.340	20.834	40.498 40.929	1.00	25.00
1731 1732	N	HIS LYS	200	117.376 120.811	23.323	44.864	$\frac{1.00}{1.00}$	25.00 18.06
1732	CA	LYS	200	119.853		45.170	1.00	19.34
1734	C	LYS	200	120.463	25.726	45.548	1.00	21.28
1735	o	LYS	200	119.755	26.617	46.012	1.00	20.62
1736	СВ	LYS	200	118.898		46.264	1.00	17.06
1737	CG	LYS	200	118.144	22.630	45.875	1.00	17.06
1738	CD	LYS	200	117.287	22.086	47.005	1.00	18.82
1739	CE	LYS	200	116.597		46.559	1.00	16.83
1740	NZ	LYS	200	115.820	20.155	47.645	1.00	19.41
1741	H	LYS	200	121.264	22.869	45.599	1.00	25.00
1742	1HZ	LYS	200	1166.454	19.923	48.438	1.00	25.00
1743	2HZ	LYS	200	115.081	20.807	47.978	1.00	25.00
1744	3HZ	LYS	200	115.377	19.285	47.288	1.00	25.00
1745	N	GLY	201		25.881	45.343	1.00	23.54
1746	CA	GLY	201	122.424	27.136	45.675	1.00	19.60
1747	С	GLY	201	122.583	28.062	44.482	1.00	19.35
1748	O	GLY	201	122.569	27.613	43.338	1.00	21.61
1749	H	GLY	201	122.299	25.166	44.933	1.00	25.00
1750	N	VAL VAL	202	122.685	29.383	44.734	1.00	17.34
1751 1752	CA C	VAL VAL	202 202	122.871 124.281	30.327 30.084	43.653 43.108	$\frac{1.00}{1.00}$	17.16 20.63
1753	o	VAL	202	125.248	30.059	43.874	1.00	22.87
1754	СВ	VAL	202	122.722	31.778	44.168	1.00	17.85
1755	CG1	VAL	202	123.062	32.782	43.071	1.00	19.32
1756	CG2	VAL	202	121.301	32.003	44.645	1.00	15.75
1757	Н	VAL	202	122.625	29.676	45.655	1.00	25.00
1758	N	PRO	203	124.414	29.905	41.780	1.00	18.06
1759	CA	PRO	203	125.705	29.652	41.128	1.00	19.86
1760	C	PRO	203	126.889	30.506	41.588	1.00	23.15
1761	O	PRO	203	127.827	29.974	42.172	1.00	27.00
1762	CB	PRO	203	125.378	29.840	39.650	1.00	21.47
1763	CG	PRO	203	123.982	29.300	39.574	1.00	19.55
1764	CD	PRO	203	123.332	29.952	40.780	1.00	17.62
1765	N	ARG	204	126.844	31.817	41.365	1.00	21.91
1766	CA	ARG	204	127.949	32.683	41.781	1.00	20.91
1767	C	ARG	204	128.283	32.568	43.265	1 00	20.45
1753	O	ARG	204	129.455	32.598	43.638	1.00	25.04
1769	CB	ARG	204	127.681	34.149	41.426	1.00	22.61
1770	CG	ARG	204	127.940	34.519	39.972	1.00	18.14
1771	CD	ARG	204	129.420	34.487	39.618	1.00	21.89
1772	NE CZ	ARG	204	129.852	33.202	39.074	1.00	23.42
1773	CZ NILI1	ARG	204	130.953	33.027	38.345	1.00	28.33
1774 1775	NH1 NH2	ARG ARG	204 204	131.747 131.248	34.055 31.827	38.069 37.862	$\frac{1.00}{1.00}$	25.07 26.89
1776	Nнz Н	ARG	204	126.071	32.201	40.913	1.00	25.00
1770	11	ANO	204	120.071	J2.201	TU.713	1.00	23.00

TABLE 11-continued

Structural	Coordinates of Tobacco 5-Epi-Aristolochene Synthase	
	In the Absence of Bound Substrate	

		In the	e Absence o	f Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
1777	HE	ARG	204	129.297	32.416	39.258	1.00	25.00
1778	1HH1	ARG	204	131.521	34.967	38.404	1.00	25.00
1779	2HH1	ARG	204	132.570	33.921	37.522	1.00	25.00
1780	1HH2	ARG	2004	130.647	31.051	38.047	1.00	25.00
1781	2HH2	ARG	204	132.077	31.699	37.316	1.00	25.00
1782	N	VAL	205	127.264	32.431	44.108	1.00	17.32
1783	CA	VAL	205	127.496	32.312	45.545	1.00	18.99
1784 1785	C O	VAL VAL	205 205	128.267 129.220	31.034 31.048	45.866 46.647	1.00 1.00	20.53 24.18
1786	CB	VAL	205	126.175	32.309	46.339	1.00	19.83
1787	CG1	VAL	205	126.442	32.022	47.811	1.00	14.97
1788	CG2	VAL	205	125.473	33.642	46.190	1.00	21.52
1789	Н	VAL	205	126.353	32.398	43.764	1.00	25.00
1790	N	GLU	206	127.862	29.933	45.249	1.00	22.23
1791	CA	GLU	206	128.519	28.649	45.478	1.00	22.70
1792	C	GLU	206	129.919	28.601	44.876	1.00	18.65
1793	O	GLU	206	130.836	28.029	45.469	1.00	20.84
1794	CB	GLU	206	127.648	27.506	44.957	1.00	17.73
1795	CG	GLU	206	126.317	27.413	45.683	1.00	20.17
1796	CD	GLU	206	126.478	27.407	47.201	1.00	26.03
1797	OE1	GLU	206	127.190	26.523	47.721	1.00	21.64
1798	OE2	GLU	206	125.895	28.283	47.876	1.00	20.28
1799	H	GLU	206	127.111	29.981	44.620	1.00	25.00
1800	N	THR	207	130.081	29.223	43.714	1.00	18.09
1801	CA	THR	207	131.369	29.291	43.038	1.00	21.23
1802	C O	THR THR	207 207	132.373	30.057 29.568	43.909 44.179	$\frac{1.00}{1.00}$	25.71 28.93
1803 1804	CB	THR	207	133.474 131.219	29.984	41.672	1.00	24.22
1805	OG1	THR	207	130.529	29.1077	40.770	1.00	28.95
1806	CG2	THR	207	132.573	30.379	41.088	1.00	23.10
1807	H	THR	207	129.311	29.644	43.289	1.00	25.00
1808	HG1	THR	207	131.030	28.287	40.685	1.00	25.00
1809	N	ARG	208	131.973	31.238	44.374	1.00	24.65
1810	CA	ARG	208	132.825	32.070	45.221	1.00	25.56
1811	C	ARG	208	133.292	31.273	46.432	1.00	25.87
1812	O	ARG	208	134.472	31.289	46.780	1.00	27.73
1813	CB	ARG	208	132.059	33.314	45.682	1.00	25.72
1814	CG	ARG	208	132.836	34.258	46.588	1,00	31.65
1815	CD	ARG	208	134.062	34.826	45.892	1.00	39.53
1816	NE	ARG	208	134.374	36.184	46.344	1.00	46.43
1817	CZ	ARG	208	135.283	36.488	47.266	1.00	47.18
1818	NH11	ARG	208	135.991	35.534	47.858	1.00	54.31
1819	NH2 H	ARG ARG	208 208	135.492 131.077	37.754 31.566	47.592 44.139	1.00 1.00	54.01 25.00
1820 1821	HE	ARG	208	133.881	36.924	45.937	1.00	25.00
1822	1HH1	ARG	208	135.847	34.577	47.610	1.00	25.00
1823	2HH1	ARG	208	136.673	35.775	48.548	1.00	25.00
1824	1HH2	ARG	208	134.962	38.478	47.150	1.00	25.00
1825	2HH2	ARG	208	136.172	37.986	48.287	1.00	25.00
1826	N	PHE	209	132.364	30.556	47.056	1.00	25.17
1827	CA	PHE	209	132.688	29.750	48.224	1.00	23.72
1828	C	PHE	209	133.677	28.632	47.908	1.00	24.69
1829	O	PHE	209	134.656	28.442	48.626	1.00	24.23
1830	CB	PHE	209	131.430	29.135	48.838	1.00	22.94
1831	CG	PHE	209	131.721	28.195	49.976	1.00	22.62
1832	CD1	PHE	209	132.019	28.691	51.242	1.00	22.83
1833	CD2	PHE	209	131.745	26.817	49.773	1.00	20.76
1834	CE1	PHE	209	132.336	27.824	52.293	1.00	22.43
1835	CE2	PHE	209	132.060 132.358	25.946	50.813	1.00	24.13
1836 1837	CZ H	PHE PHE	209 209	131.439	26.450 30.580	52.075 46.728	1.00 $1.00$	22.16 25.00
1838	N N	PHE	210	133.399	27.872	46.856	1.00	24.89
1839	CA	PHE	210	134.263	26.765	40.486	1.00	21.48
1840	C	PHE	210	135.671	27.241	40.172	1.00	23.23
1841	ŏ	PHE	210	136.645	26.676	48.671	1.00	24.15
1842	СВ	PHE	210	133.688	25.989	45.296	1.00	18.25
1843	CG	PHE	210	134.4776		44.944	1.00	20.04
1844	CD1	PHE	210	134.506	23.661	45.811	1.00	17.95
1845	CD2	PHE	210	135.212	24.694	43.763	1.00	19.06
1846	CE1	PHE	210	135.260	22.525	45.510	1.00	17.28
1847	CE2	PHE	210	135.972	23.563	43.450	1.00	21.18
1848	CZ	PHE	210	135.995	22.476	44.329	1.00	18.42
1849	H	PHE	210	132.602	28.062	40.315	1.00	25.00
1850	N	ILE	211	135.781	28.290	45.368	1.00	26.53

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
1851	CA	ILE	211	137.086	28.818	44.997	1.00	27.06
1852	C	ILE	211	137.917	29.248	40.205	1.00	27.01
1853	O	ILE	211	138.953	28.652	46.490	1.00	25.02
1854	CB	ILE	211	136.967	30.015	44.023	1.00	23.08
1855	CG1	ILE	211	136.317	29.574	42.713	1.00	22.07
1856	CG2	ILE	211	138.344	30.603	43.737	1.00	18.73
1857	CD1	ILE	211	136.163	30.700	41.701	1.00	22.59
1858 1859	H N	ILE SER	211 212	134.968 137.430	28.712 30.233	45.022 46.949	1.00 $1.00$	25.00 26.45
1860	CA	SER	212	138.174	30.758	48.087	1.00	29.47
1861	C	SER	212	138.263	29.914	49.355	1.00	29.82
1862	Ö	SER	212	139.317	29.873	49.993	1.00	31.08
1863	CB	SER	212	137.691	32.173	48.425	1.00	28.90
1864	OG	SER	212	136.311	32.186	48.742	1.00	48.04
1865	H	SER	212	136.550	30.622	46.740	1.00	25.00
1866	HG	SER	212	136.156	31.647	49.517	1.00	25.00
1867	N	SER	213	137.175	29.249	49.728	1.00	25.54
1868	CA	SER	213	137.173	28.447	50.949	1.00	25.42
1869	C	SER	213	137.555	26.969	50.823	1.00	23.90
1870	O	SER	213	138.019	26.371	51.794	1.00	29.58
1871 1872	CB OG	SER SER	213 213	135.820 135.503	28.566 29.920	51.662 51.942	1.00 1.00	19.77 36.00
1873	H	SER	213	136.366	29.920	49.173	1.00	25.00
1874	HG	SER	213	135.449	30.406	51.120	1.00	25.00
1875	N	ILE	214	137.390	26.376	49.645	1.00	20.94
1876	CA	ILE	214	137.701	24.958	49.502	1.00	20.03
1877	С	ILE	214	138.869	24.617	48.591	1.00	20.98
1878	O	ILE	214	139.914	24.174	49.065	1.00	23.05
1879	CB	ILE	214	136.463	24.144	49.041	1.00	20.03
1880	CG1	ILE	214	135.255	24.455	49.932	1.00	15.38
1881	CG2	ILE	214	136.778	22.640	49.046	1.00	14.86
1882	CD1	ILE	214	135.488	24.207	51.418	1.00	14.56
1883	H	ILE	214	137.066	26.887	48.873	1.00	25.00
1884	N	TYR	215	138.696	24.823	47.289	1.00	18.70
1885 1886	CA C	TYR	215 215	139.733 141.076	24.490 25.168	46.323 46.582	$\frac{1.00}{1.00}$	22.93 25.73
1887	o	TYR TYR	215	142.128	24.545	46.450	1.00	25.75
1888	СВ	TYR	215	139.258	24.777	44.899	1.00	19.89
1889	CG	TYR	215	139.859	23.834	43.884	1.00	17.39
1890	CD1	TYR	215	139.726	22.455	44.030	1.00	18.08
1891	CD2	TYR	215	140.557	24.315	42.782	1.00	19.17
1892	CE1	TYR	215	140.275	21.575	43.102	1.00	17.39
1893	CE2	TYR	215	141.113	23.445	41.843	1.00	16.77
1894	CZ	TYR	215	140.967	22.076	42.010	1.00	21.36
1895	ОН	TYR	215	141.517	21.212	41.088	1.00	25.91
1896	H	TYR	215	137.858	25.217	46.973	1.00	25.00
1897	HH	TYR	215	141.317	20.302	41.340	1.00	25.00
1898	N CA	ASP	216	141.037	26.434 27.184	46.969 47.250	1.00	26.71 32.33
1899 1900	CA	ASP ASP	216 216	142.254 143.057	26.532	48.377	1.00 $1.00$	32.33
1901	Ö	ASP	216	144.288	26.589	48.387	1.00	33.87
1902	СB	ASP	216	141.895	28.621	47.636	1.00	35.77
1903	CG	ASP	216	143.111	29.514	47.769	1.00	36.73
1904	OD1	ASP	216	143.842	29.670	46.769	1.00	37.16
1905	OD2	ASP	216	143.327	30.062	48.871	1.00	41.08
1906	H	ASP	216	140.170	26.885	447.054	1.00	25.00
1907	N	LYS	217	142.350	25.910	49.316	1.00	31.70
1908	CA	LYS	217	142.978	25.255	50.459	1.00	29.37
1909	С	LYS	217	143.134	23.745	50.269	1.00	30.32
1910	O	LYS	217	143.506	23.029	51.200	1.00	31.00
1911 1912	CB CG	LYS LYS	217 217	142.170 142.062	25.553 27.033	51.724 52.017	$\frac{1.00}{1.00}$	26.09 27.73
1912	CD	LYS	217	141.185	27.033	53.213	1.00	35.35
1913	CE	LYS	217	141.103	28.807	53.463	1.00	40.60
1915	NZ	LYS	217	140.124	29.115	54.551	1.00	49.88
1916	Н	LYS	217	141.376	25.876	49.235	1.00	25.00
1917	1HZ	LYS	217	140.429	28.650	55.430	1.00	25.00
1918	2HZ	LYS	217	140.083	30.143	54.698	1.00	25.00
1919	3HZ	LYS	217	139.181	28.767	54.284	1.00	25.00
1920	N	GLU	218	142.864	23.271	49.057	1.00	31.03
1921	CA	GLU	218	142.961	21.855	48.750	1.00	33.23
1922	С	GLU	218	144.391	21.489	48.357	1.00	42.71
1923	O	GLU	218	144.932	22.012	47.381	1.00	41.53
1924	CB	GLU	218	141.983	21.492	47.626	1.00	32.79

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
1925	CG	GLU	218	141.873	20.007	47.345	1.00	49.16
1926	CD	GLU	218	141.324	19.228	48.526	1.00	61.57
1927	OE1 OE2	GLU	218	140.147	19.456 18.391	48.886 49.092	1.00	66.99
1928 1929	H	GLU GLU	218 218	142.066 142.616	23.896	48.343	$\frac{1.00}{1.00}$	62.96 25.00
1930	N	GLN	219	144.974	20.551	49.098	1.00	48.16
1931	CA	GLN	219	146.339	20.089	48.858	1.00	52.73
1932	C	GLN	219	146.533	19.487	47.467	1.00	49.21
1933	O	GLN	219	147.594	19.622	46.870	1.00	51.36
1934	CB	GLN	219	146.733	19.063	49.929	1.00	62.56
1935 1936	CG CD	GLN GLN	219 219	148.127 148.498	19.262 18.186	50.531 51.534	1.00 $1.00$	81.40 90.93
1937	OE1	GLN	219	148.863	17.072	51.156	1.00	97.11
1938	NE2	GLN	219	148.408	18.512	52.825	1.00	96.41
1939	H	GLN	219	144.450	20.162	49.821	1.00	25.00
1940	1HE2	GLN	219	148.113	19.389	53.115	1.00	25.00
1941	2HE2	GLN	219	148.656	17.793	53.455	1.00	25.00
1942	N	SER	220	145.496	18.842	46.950	1.00	47.51
1943 1944	CA C	SER SER	220 220	145.552 144.945	18.199 19.020	45.636 44.487	$\frac{1.00}{1.00}$	47.04 45.29
1945	Ö	SER	220	144.577	18.467	43.446	1.00	47.02
1946	СВ	SER	220	144.862	16.833	45.713	1.00	51.73
1947	OG	SER	220	143.585	16.948	46.327	1.00	55.26
1948	H	SER	220	144.658	18.796	47.447	1.00	25.00
1949	HG	SER	220	143.671	17.280	47.217	1.00	25.00
1950	N	LYS	221	144.849	20.332	44.679	1.00	38.03
1951	CA	LYS	221	144.270	21.233	43.682	1.00	32.98
1952 1953	C O	LYS LYS	221 221	145.037 146.249	21.284 21.077	42.363 42.328	$\frac{1.00}{1.00}$	30.63 33.91
1954	СВ	LYS	221	144.206	22.649	44.255	1.00	33.08
1955	CG	LYS	221	145.584	23.257	44.500	1.00	40.24
1956	CD	LYS	221	145.512	24.563	45.257	1.00	53.13
19557	CE	LYS	221	146.902	25.093	45.561	1.00	55.90
1958	NZ	LYS	221	146.843	26.358	46.344	1.00	67.32
1959	H 1HZ	LYS LYS	221 221	145.198	20.720	45.508 47.244	1.00 $1.00$	25.00 25.00
1960 1961	2HZ	LYS	221	146.350 147.807	26.191 26.698	46.533	1.00	25.00
1962	3HZ	LYS	221	146.326	27.080	45.800	1.00	25.00
1963	N	ASN	222	144.322	21.538	41.273	1.00	28.13
1964	CA	ASN	222	144.958	21.675	39.970	1.00	25.27
1965	C	ASN	222	145.154	23.174	39.816	1.00	30.00
1966	O	ASN	222	144.187	23.933	39.707	1.00	29.84
1967 1968	CB CG	ASN ASN	222 222	144.077 144.688	21.149 21.390	38.843 37.473	$\frac{1.00}{1.00}$	22.73 24.93
1969	OD1	ASN	222	144.914	22.534	37.072	1.00	31.82
1970	ND2	ASN	222	144.973	20.317	36.755	1.00	23.78
1971	H	ASN	222	143.353	21.840	41.343	1.00	25.00
1972	2HD2	ASN	222	145.364	20.460	35.868	1.00	25.00
1973	2HD2	ASN	222	144.784	19.432	37.125	1.00	25.00
1974	N	ASN	223	146.412	23.596	39.819	1.00	31.07
1975 1976	CA C	ASN ASN	223 223	146.759 146.273	25.009 25.730	39.726 38.477	$\frac{1.00}{1.00}$	26.94 25.82
1977	Ö	ASN	223	145.233	26.910	38.538	1.00	27.88
1978	CB	ASN	223	148.261	25.185	39.915	1.00	23.98
1979	CC	ASN	223	148.732	24.633	41.242	1.00	28.15
1980	OD1	ASN	223	143.586	25.271	42.281	1.00	30.20
1981	ND2	ASN	223	149.291	23.423	41.219	1.00	25.49
1982 1983	H 1HD2	ASN ASN	223 223	147.118 149.595	22.925 23.063	39.901 42.082	$\frac{1.00}{1.00}$	25.00 25.00
1984	2HD2	ASN	223	149.377	22.938	40.382	1.00	25.00
1985	N	VAL	224	146.224	25.036	37.346	1.00	24.73
1988	CA	VAL	224	145.743	25.667	36.124	1.00	27.15
1987	C	VAL	224	144.263	26.026	36.304	1.00	28.87
1988	O	VAL	224	143.852	27.150	36.019	1.00	29.97
1989 1990	CB CG1	VAL	224	145.914	24.742	34.900	1.00	31.17
1990 1991	CG1 CG2	VAL VAL	224 224	145.359 147.382	25.404 24.400	33.651 34.707	$\frac{1.00}{1.00}$	30.27 30.28
1992	H	VAL	224	146.488	24.096	37.329	1.00	25.00
1993	N	LEU	225	143.486	25.089	36.843	1.00	25.78
1994	CA	LEU	225	142.057	25.303	37.069	1.00	25.90
1995	C	LEU	225	141.792	26.380	38.125	1.00	26.51
1996	O	LEU	225	140.900	27.214	37.956	1.00	26.55
1997 1998	CB CG	LEU	225 225	141.386 141.398	23.991 22.848	37.463 38.441	1.00	20.07
1330	CU	LEU	223	141.370	22.040	30. <del>44</del> 1	1.00	21.81

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
1999	CD1	LEU	225	140.664	21.638	36.991	1.00	10.56
2000	CD2	LEU	225	140.780	23.295	35.126	1.00	19.02
2001 2002	H N	LEU LEU	225 226	143.883 142.566	24.229 26.369	37.092 39.207	$\frac{1.00}{1.00}$	25.00 22.12
2002	CA	LEU	226	142.400	27.387	40.261	1.00	27.39
2003	C	LEU	228	142.724	28.775	39.743	1.00	28.78
2005	O	LEU	226	141.967	29.720	39.969	1.00	34.36
2006	CB	LEU	226	143.282	27.033	41.468	1.00	25.60
2007	CG	LEU	226	143.170	27.984	42.6665	1.00	26.39
2008	CD1	LEU	226	141.731	28.037	43.183	1.00	20.17
2009 2010	CD2 H	LEU LEU	226 226	144.110 143.249	27.532 25.673	43.763 39.298	$\frac{1.00}{1.00}$	26.04 25.00
2010	N	ARG	227	143.842	28.904	39.036	1.00	28.64
2012	CA	ARG	227	144.270	30.183	38.473	1.00	30.51
2013	C	ARG	227	143.186	30.688	37.508	1.00	29.19
2014	O	ARG	227	142.770	31.849	37.567	1.00	25.86
2015	CB	ARG	227	145.607	29.989	37.742	1.00	30.51
2016	CG	ARG	227	146.171	31.215	37.037	1.00	32.00
2017 2018	CD NE	ARG ARG	227 227	140.883 147.414	32.162 33.314	37.981 37.256	$\frac{1.00}{1.00}$	35.49 34.46
2019	CZ	ARG	227	147.799	34.464	37.822	1.00	32.62
2020	NH1	ARG	227	147.727	34.611	39.136	1.00	33.39
2021	NH2	ARG	227	148.214	35.460	37.066	1.00	35.87
2022	H	ARG	227	144.402	28.114	38.884	1.00	25.00
2023	HE	ARG	227	147.480	33.245	36.292	1.00	25.00
2024	1HH1	ARG	227	147.381	33.871	39.712	1.00	25.00
2025	2HH1	ARG	227	148.020	35.470	39.564	$\frac{1.00}{1.00}$	25.00
2026 2027	1HH2 2HH2	ARG ARG	227 227	148.236 148.505	35.359 36.315	36.073 37.491	1.00	25.00 25.00
2027	N	PHE	228	142.723	29.788	36.645	1.00	28.49
2029	CA	PHE	228	141.678	30.063	35.656	1.00	30.24
2030	C	PHE	228	140.411	30.575	36.369	1.00	29.56
2031	O	PHE	228	139.909	31.662	36.068	1.00	31.17
2032	CB	PHE	228	141.394	28.752	34.892	1.00	32.31
2033	CG CD1	PHE	228 228	140.441 139.889	28.879 30.103	33.721 33.348	1.00	30.15
2034 2035	CD1 CD2	PHE PHE	228	140.084	27.740	32.999	1.00 $1.00$	29.90 24.53
2036	CE1	PHE	228	138.994	30.186	32.277	1.00	28.21
2037	CE2	PHE	228	139.193	27.811	31.930	1.00	23.61
2038	CZ	PHE	228	138.646	29.036	31.568	1.00	29.31
2039	H	PHE	228	143.108	28.886	36.672	1.00	25.00
2040	N	ALA	229	139.935	29.809	37.344	1.00	25.42
2041 2042	CA C	ALA ALA	229 229	138.737 138.808	30.156 31.533	38.099 38.764	$\frac{1.00}{1.00}$	25.11 29.20
2042	o	ALA	229	137.847	32.310	38.703	1.00	26.99
2044	CB	ALA	229	138.456	29.085	39.135	1.00	20.89
2045	H	ALA	229	140.406	28.982	37.565	1.00	25.00
2046	N	LYS	230	139.944	31.838	39.389	1.00	9.21
2047	CA	LYS	230	140.127	33.121	40.068	1.00	29.75
2048 2049	C O	LYS LYS	230 230	140.100 139.405	34.306 35.298	39.109 39.350	$\frac{1.00}{1.00}$	30.11 30.69
2050	СВ	LYS	230	141.434	33.136	40.866	1.00	30.09
2051	CG	LYS	230	141.422	32.247	42.100	1.00	29.79
2052	CD	LYS	230	142.686	32.430	42.923	1.00	24.40
2053	CE	LYS	230	142.595	31.664	44.227	1.00	29.55
2054	NZ	LYS	230	143.790	31.883	45.079	1.00	36.37
2055	H 1117	LYS	230	140.676	31.183	39.394	1.00	25.00
2056 2057	1HZ 2HZ	LYS LYS	230 230	143.881 143.689	32.896 31.356	45.296 45.965	$\frac{1.00}{1.00}$	25.00 25.00
2058	33HZ	LYS	230	144.640	31.560	44.573	1.00	25.00
2059	N	LEU	231	140.852	34.201	38.016	1.00	31.56
2060	CA	LEU	231	140.911	35.275	37.032	1.00	30.17
2061	C	LEU	231	139.549	35.506	36.394	1.00	30.20
2062	O	LEU	231	139.085	36.645	36.299	1.00	26.72
2063 2084	CB CG	LEU LEU	231 231	141.941 143.408	34.959 34.790	35.941 36.340	$\frac{1.00}{1.00}$	29.21 26.11
2084	CD1	LEU	231	143.408	34.631	35.077	1.00	20.11
2066	CD2	LEU	231	143.890	35.990	37.139	1.00	22.48
2067	Н	LEU	231	141.374	33.380	37.869	1.00	25.00
2068	N	ASP	232	138.898	34.417	35.995	1.00	29.28
2069	CA	ASP	232	137.593	34.490	35.351	1.00	28.41
2070	С	ASP	232	136.549	35.149	36.247	1.00	25.56
2071 2072	O CB	ASP ASP	232 232	135.820 137.131	36.044 33.097	35.813 34.932	$\frac{1.00}{1.00}$	25.21 26.05
2012	CD	11.31	202	101.101	55.057	54.752	1.00	20.05

TABLE 11-continued

		In the	e Absence of			ле буни		
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
2073	CG	ASP	232	136.143	33.143	33.793	1.00	33.93
2074	OD1	ASP	232	136.587 134.927	33.188	32.627	1.00	32.45
2075 2076	OD2 H	ASP ASP	232 232	134.927	33.157 33.539	34.060 36.141	$\frac{1.00}{1.00}$	29.97 25.00
2077	N	PHE	232	136.510	34.730	37.507	1.00	24.45
2078	CA	PHE	233	135.569	35.286	38.466	1.00	22.77
2079	С	PHE	233	135.788	36.788	38.603	1.00	28.10
2080	O	PHE	233	134.835	37.568	38.516	1.00	31.99
2081	CB	PHE	233	135.732	34.605	39.831	1.00	19.36
2082	CG	PHE	233	134.714	35.035	40.854	1.00	22.51
2083 2084	CD1 CD2	PHE PHE	233 233	1344.85 133.604	736.241 34.237	41.540 41.127	$\frac{1.00}{1.00}$	22.18 23.90
2085	CE1	PHE	233	133.908	36.646	42.481	1.00	26.05
2086	CE2	PHE	233	132.650	34.631	42.085	1.00	23.08
2087	CZ	PHE	233	132.801	35.839	42.743	1.00	24.51
2088	H	PHE	233	137.126	34.022	37.802	1.00	25.00
2089	N	ASN	234	137.044	37.189	38.801	1.00	28.39
2090	CA	ASN	234	137.393	38.602	38.965	1.00	28.37
2091	C O	ASN	234 234	137.079	39.446	37.743	1.00	27.25
2092 2093	СВ	ASN ASN	234	136.608 138.867	40.575 38.762	37.868 39.342	$\frac{1.00}{1.00}$	32.70 27.32
2094	CG	ASN	234	139.152	38.353	40.778	1.00	30.63
2095	OD1	ASN	234	138.242	38.219	41.595	1.00	29.34
2096	ND2	ASN	234	140.426	38.166	41.092	1.00	37.28
2097	H	ASN	234	137.753	36.513	38.851	1.00	25.00
2098	1HD2	ASN	234	140.628	37.901	42.015	1.00	25.00
2099	2HD2	ASN	234	141.114	38.292	40.408	1.00	25.00
2100 2101	N CA	LEU LEU	235 235	137.339 137.059	38.902 39.616	36.561 35.321	1.00 1.00	28.66 29.93
2102	C	LEU	235	135.551	39.830	35.167	1.00	30.84
2103	Ō	LEU	235	135.105	40.949	34.905	1.00	31.89
2104	CB	LEU	235	137.625	38.852	34.119	1.00	29.05
2105	CG	LEU	235	137.476	39.509	32.742	1.00	30.21
2108	CD1	LEU	235	138.045	40.922	32.769	1.00	29.02
2107 2108	CD2 H	LEU LEU	235 235	138.173 137.721	38.667 38.001	31.684 36.525	$\frac{1.00}{1.00}$	31.45 25.00
2108	N	LEU	236	134.766	38.769	35.352	1.00	29.90
2110	CA	LEU	238	133.311	38.875	35.245	1.00	28.94
2111	C	LEU	238	132.774	39.874	36.263	1.00	28.31
2112	O	LEU	238	131.833	40.623	35.979	1.00	29.04
2113	CB	LEU	236	132.632	37.518	35.463	1.00	26.96
2114	CG CD1	LEU LEU	236	132.722	36.463 35.299	34.359 34.694	1.00	32.49 28.63
2115 2116	CD1 CD2	LEU	236 236	131.797 132.326	35.299	33.026	$\frac{1.00}{1.00}$	30.07
2117	H	LEU	236	135.173	37.900	35.561	1.00	25.00
2118	N	GLN	237	133.362	39.870	37.454	1.00	25.46
2119	CA	GLN	237	132.953	40.777	38.521	1.00	25.58
2120	С	GLN	237	133.059	42.231	38.082	1.00	27.92
2121	O	GLN	2337	132.201	43.054	38.387 39.769	1.00	29.65
2122 2123	CB CG	GLN GLN	237 237	133.807 133.342	40.549 41.314	40.993	$\frac{1.00}{1.00}$	20.55 23.60
2124	CD	GLN	237	134.216	41.046	42.197	1.00	31.25
2125	OE1	GLN	237	135.435	41.196	42.134	1.00	31.93
2126	NE2	GLN	237	133.602	40.634	43.298	1.00	28.03
2127	H	GLN	237	134.094	39.237	37.625	1.00	25.00
2128	1HE2	GLN	237	134.156	40.446	44.081	1.00	25.00
2129 2130	2HE2 N	GLN MET	237 238	132.635 134.096	40.519 42.537	43.287 37.286	$\frac{1.00}{1.00}$	25.00 28.91
2131	CA	MET	238	134.288	43.888	36.776	1.00	33.08
2132	C	MET	238	133.084	44.282	35.924	1.00	33.53
2133	O	MET	238	132.562	45.391	36.049	1.00	37.31
2134	CB	MET	238	135.573	43.976	35.954	1.00	32.86
2135	CG	MET	238	136.836	43.837	36.782	1.00	39.06
2136	SD	MET	238	138.318	43.815	35.763	1.00	43.74
2137 2138	CE H	MET MET	238 238	139.508 134.751	43.186 41.842	36.929 37.055	$\frac{1.00}{1.00}$	46.28 25.00
2138	n N	LEU	238 239	134.751	43.356	35.087	1.00	31.84
2140	CA	LEU	239	131.465	43.599	34.233	1.00	30.89
2141	C	LEU	239	130.219	43.801	35.097	1.00	28.36
2142	O	LEU	239	129.450	44.742	34.885	1.00	32.90
2143	CB	LEU	239	131.255	42.427	33.271	1.00	29.61
2144	CG CD1	LEU	239	129.969	42.432	32.436	1.00	29.85
2145 2146	CD1 CD2	LEU LEU	239 239	129.929 129.870	43.636 41.138	31.499 31.649	$\frac{1.00}{1.00}$	25.22 26.65
2140	CD2	LEU	237	122.070	71.130	J1.047	1.00	20.00

TABLE 11-continued

		In the	Absence o	f Bound St	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
2147	H	LEU	239	133.084	42.491	35.043	1.00	25.00
2148	N	HIS	240	130.042	42.942	36.095	1.00	25.85
2149	CA	HIS	240 240	128.891	43.042 44.406	36.990	1.00	28.32
2150 2151	C O	HIS HIS	240	128.885 127.824	44.974	37.671 37.940	$\frac{1.00}{1.00}$	30.20 30.10
2152	СВ	HIS	240	128.925	41.926	38.036	1.00	25.55
2153	CG	HIS	240	128.881	40.545	37.448	1.00	24.20
2154	ND1	HIS	240	129.449	39.455	38.058	1.00	25.29
2155	CD2	HIS	240	128.358	40.103	36.283	1.00	21.44
2156	CE1	HIS	240	129.289	38.394	37.302	1.00	27.58
2157 2158	NE2 H	HIS HIS	240 240	128.627 130.713	38.750 42.244	36.209 36.233	1.00 1.00	21.74 25.00
2159	HD11	HIS	240	129.918	39.455	38.924	1.00	25.00
2160	HE2	HIS	240	128.362	38.151	35.470	1.00	25.00
2161	N	LYS	241	130.079	44.926	37.935	1.00	34.60
2162	CA	LYS	241	130.239	46.230	38.563	1.00	32.03
2163	С	LYS	241	129.855	47.374	37.613	1.00	31.99
2164	O	LYS	241	129.280	48.374	38.045	1.00	29.66
2165 2166	CB CG	LYS LYS	241 241	131.675 131.984	46.391 45.614	39.072 40.356	$\frac{1.00}{1.00}$	32.10 34.40
2167	CD	LYS	241	133.447	45.807	40.758	1.00	41.19
2168	CE	LYS	241	133.701	45.417	42.208	1.00	49.51
2169	NZ	LYS	241	135.044	45.881	42.698	1.00	56.06
2110	H	LYS	241	130.875	44.403	37.703	1.00	25.00
2171	1HZ	LYS	241	135.111	46.916	42.625	1.00	25.00
2172	2HZ	LYS	241	135.179	45.610	43.698	1.00	25.00
2173 2174	3HZ	LYS	241	135.803	45.446	42.135	1.00	25.00
2174	N CA	GLN GLN	242 242	130.121 129.799	47.201 48.211	36.316 35.303	$\frac{1.00}{1.00}$	34.13 38.39
2176	C	GLN	242	128.288	48.278	35.161	1.00	39.98
2177	ŏ	GLN	242	127.702	49.353	34.990	1.00	45.19
2178	CB	GLN	242	130.376	47.827	33.942	1.00	42.52
2179	CG	GLN	242	131.883	47.683	33.920	1.00	60.88
2180	CD	GLN	242	132.417	47.252	32.574	1.00	69.61
2181	OE1 NE2	GLN	242 242	131.663	47.052 47.102	31.620	1.00	75.13 78.40
2182 2183	H	GLN GLN	242	133.730 130.490	46.358	32.488 35.996	1.00 1.00	25.00
2184	1HE2	GLN	242	134.0772	46.820	31.621	1.00	25.00
2185	2HE2	GLN	242	134.282	47.272	33.272	1.00	25.00
2186	N	GLU	243	127.674	47.105	35.219	1.00	33.89
2187	CA	GLU	243	126.233	46.975	35.107	1.00	28.66
2188	С	GLU	243	125.568	47.591	36.325	1.00	28.27
2189 2190	O CB	GLU	243 243	124.635	48.381	36.193 34.982	$\frac{1.00}{1.00}$	32.26 25.09
2191	CG	GLU GLU	243	125.857 126.416	45.505 44.820	33.741	1.00	24.03
2192	CD	GLU	243	126.182	43.329	33.738	1.00	26.58
2193	OE1	GLU	243	125.633	42.810	34.726	1.00	24.63
2194	OE2	GLU	243	126.556	42.665	32.750	1.00	29.69
2195	H	GLU	243	128.248	46.333	35.341	1.00	25.00
2196	N	LEU	244	126.078	47.267	37.508	1.00	24.94
2197 2198	CA C	LEU LEU	244 244	125.522 125.635	47.798 49.324	38.745 38.766	$\frac{1.00}{1.00}$	30.48 36.45
2199	ō	LEU	244	124.700	50.021	39.163	1.00	35.07
2200	СB	LEU	244	126.233	47.185	39.957	1.00	28.96
2201	CG	LEU	244	125.765	47.658	41.339	1.00	29.10
2202	OD1	LEU	244	124.249	47.527	41.484	1.00	25.23
2203	CD2	LEU	244	126.464	48.859	42.423	1.00	27.57
2204 2205	H N	LEU ALA	244 245	126.855 126.778	48.662 49.832	37.535 38.318	$\frac{1.00}{1.00}$	25.00 37.15
2206	CA	ALA	245	127.023	51.268	38.270	1.00	38.62
2207	C	ALA	245	126.030	51.937	37.325	1.00	39.57
2208	O	ALA	245	125.352	52.902	37.692	1.00	43.21
2209	CB	ALA	245	128.452	51.539	37.802	1.00	35.23
2210	H	ALA	248	127.477	49.215	38.032	1.00	25.00
2211	N C4	GLN	248	125.920	51.380	36.123	1.00	35.82
2212 2213	CA C	GLN GLN	248 248	125.025 123.577	51.888 51.989	35.086 35.566	$\frac{1.00}{1.00}$	40.78 43.61
2214	o	GLN	248	122.907	53.016	35.404	1.00	43.37
2215	СВ	GLN	248	125.088	50.963	33.872	1.00	42.45
2216	CG	GLN	248	124.151	51.350	32.733	1.00	59.73
2217	CD	GLN	248	124.148	50.338	31.610	1.00	65.77
2218	OE1	GLN	248	125.149	49.663	31.357	1.00	67.95
2219 2220	NE2 H	GLN GLN	248 248	123.009 126.450	50.223 50.576	30.920 35.964	$\frac{1.00}{1.00}$	66.48 25.00
2220		CLIT	240	220.70U	50.570	22.207	1.00	25.00

TABLE 11-continued

	Struct	In the	e Absence o			ene Syntha		
Atom Type	Atom	Residue	Residue #	X	Y	Z	occ	B-factor
2221	1HE2	GLN	248	123.038	49.559	30.198	1.00	25.00
2222	2HE2	GLN	248	122.232	50.763	31.132	1.00	25.00
2223 2224	N C4	VAL VAL	247 247	123.115	50.916	36.185	1.00	39.07
2224	CA C	VAL VAL	247	121.762 121.538	50.830 51.732	36.692 37.908	$\frac{1.00}{1.00}$	37.02 40.33
2226	o	VAL	247	120.435	52.248	38.106	1.00	39.92
2227	СВ	VAL	247	121.387	49.341	36.948	1.00	35.95
2228	CG1	VAL	247	120.417	49.201	38.091	1.00	37.82
2229	CG2	VAL	247	120.794	48.754	35.686	1.00	32.90
2230	H	VAL	247	123.730	50.170	36.340	1.00	25.00
2231	N	SER	248	122.579	51.926	38.715	1.00	44.51
2232	CA	SER	248	122.483	52.798	39.887	1.00	46.80
2233 2234	C O	SER SER	248 248	122.250 121.454	54.234 54.976	39.410 39.997	1.00 1.00	47.29
2235	СВ	SER	248	123.759	52.727	40.726	1.00	46.67 44.77
2238	CC	SER	243	123.859	51.479	41.381	1.00	45.74
2237	H	SER	248	123.423	51.465	38.531	1.00	25.00
2238	HG	SER	248	123.876	50.777	40.722	1.00	25.00
2239	N	ARG	249	122.938	54.615	38.334	1.00	44.35
2240	CA	ARG	249	122.789	55.943	37.750	1.00	48.22
2241	С	ARG	249	121.354	56.097	37.256	1.00	47.00
2242	O	ARG	249	120.710	57.119	37.504	1.00	47.12
2243 2244	CB CG	ARG	249 249	123.785	56.147	36.604 37.075	$\frac{1.00}{1.00}$	52.69
2244	CD	ARG ARG	249	125.165 126.154	56.590 56.712	35.924	1.00	66.38 73.20
2246	NE	ARG	249	126.919	55.484	35.712	1.00	75.40
2247	CZ	ARG	249	126.922	54.778	334.584	1.00	77.33
2248	NH1	ARG	249	126.194	55.165	33.542	1.00	74.66
2249	NH2	ARG	249	127.669	53.686	34.493	1.00	83.65
2250	H	ARG	249	123.579	53.986	37.936	1.00	25.00
2251	HE	ARG	249	127.471	55.153	36.453	1.00	25.00
2252	1HH1	ARG	249	125.633	55.990	33.598	$\frac{1.00}{1.00}$	25.00
2253 2254	2HH1 1HH2	ARG ARG	249 249	126.203 128.229	54.625 53.396	32.700 35.269	1.00	25.00 25.00
2255	2HH2	ARG	249	123.229	53.153	33.648	1.00	25.00
2256	N	TRP	250	120.848	55.053	36.603	1.00	46.33
2257	CA	TRP	250	119.480	55.024	36.092	1.00	43.84
2258	C	TRP	250	118.488	55.311	37.230	1.00	46.38
2259	O	TRP	250	117.566	56.118	37.075	1.00	44.72
2260	CB	TRRP	250	119.201	53.652	35.456	1.00	38.48
2261	CG	TRP	250	117.747	53.324	35.232	1.00	37.37
2262 2263	CD1 CD2	TRP TRP	250 250	116.986 116.891	53.661 52.569	34.150 36.105	1.00 $1.00$	33.31 36.85
2264	NE1	TRP	250	115.713	53.164	34.293	1.00	34.01
2265	CE2	TRP	250	115.626	52.490	35.483	1.00	36.82
2266	CE3	TRP	250	117.070	51.952	37.352	1.00	34.91
2267	CZ2	TRP	250	114.543	51.816	36.065	1.00	39.04
2268	CZ3	TRP	250	115.992	51.281	37.932	1.00	39.64
2269	CH2	TRP	250	114.748	51.220	37.286	1.00	39.49
2270	H	TRP	250 250	121.423	54.272	36.448	1.00	25.00
2271 2272	HE1 N	TRP TRP	250 251	114.984 118.718	53.281 54.683	33.650 38.382	$\frac{1.00}{1.00}$	25.00 46.65
2272	CA	TRP	251	117.859	54.851	39.551	1.00	54.20
2274	C	TRP	251	117.864	56.279	40.089	1.00	59.43
2275	O	TRP	251	116.814	56.823	40.445	1.00	62.13
2276	CB	TRP	251	118.284	53.889	40.657	1.00	51.89
2277	CG	TRP	251	117.358	53.872	41.836	1.00	58.17
2278	CD1	TRP	251	117.596	54.402	43.071	1.00	60.91
2279	CD2	TRP	251	116.069	53.248	41.908	1.00	60.72 64.22
2280 2281	NE1 CE2	TRP TRP	251 251	116.541 115.589	54.136 53.429	43.912 43.225	$\frac{1.00}{1.00}$	60.87
2282	CE3	TRP	251	115.274	52.546	40.989	1.00	56.90
2283	CZ2	TRP	251	114.351	52.934	43.648	1.00	58.08
2284	CZ3	TRP	251	114.042	52.054	41.410	1.00	53.15
2285	CH2	TRP	251	113.594	52.252	42.729	1.00	53.35
2286	H	TRP	251	119.485	54.076	38.442	1.00	25.00
2287	HE1	TRP	251	116.481	54.409	44.851	1.00	25.00
2288	N	LYS	252	119.049	56.875	40.167	1.00	64.71
2289	CA	LYS	252	119.191	58.241	40.661	1.00	68.07 65.20
2290 2291	C O	LYS LYS	252 252	118.440 117.831	59.227 60.169	39.779 40.283	$\frac{1.00}{1.00}$	65.20 65.28
2291	СВ	LYS	252	120.668	58.621	40.755	1.00	75.64
2293	CG	LYS	252	121.400	57.815	41.803	1.00	85.57
2294	CD	LYS	252	122.890	57.818	41.588	1.00	92.92

TABLE 11-continued

		In the	Absence o			ene synth	iase	
Atom Type	Atom	Residue	Residue #	X	Y	z	OCC	B-factor
2295	CE	LYS	252	123.526	56.776	42.486	1.00	95.43
2296	NZ	LYS	252	124.902	56.526	42.101	1.00	94.64
2297	H	LYS	252	119.849	56.379	39.887	1.00	25.00
22998 2299	1HZ 2HZ	LYS	252	125.056	56.256	41.101	$\frac{1.00}{1.00}$	25.00 25.00
2300	3HZ	LYS LYS	252 252	125.376 125.417	57.437 55.867	42.219 42.716	1.00	25.00
2301	N	ASP	253	118.453	58.983	38.469	1.00	63.40
2302	CA	ASP	253	117.762	59.846	37.515	1.00	63.30
2303	C	ASP	253	116.265	59.872	37.796	1.00	63.95
2304	O	ASP	253	115.635	60.925	37.729	1.00	68.15
2305	CB	ASP	253	118.003	59.376	36.077	1.00	68.18
2306	CG	ASP	253	119.467	59.453	35.664	1.00	76.46
2307	OD1	ASP	253	120.293	60.015	36.419	1.00	78.84
2308	OD2	ASP	253	119.793	58.943	34.570	1.00	79.35
2309	H	ASP	253	118.948	58.203	38.137	1.00	25.00
2310	N CA	LEU	254 254	115.697	58.710	38.105	1.00	66.44
2311 2312	CA C	LEU LEU	254 254	114.271 113.947	58.611 59.482	38.409 39.616	$\frac{1.00}{1.00}$	65.57 67.65
2312	o	LEU	254	112.815	59.462	39.784	1.00	68.72
2314	СВ	LEU	254	113.885	57.162	38.698	1.00	61.14
2315	CG	LEU	254	114.124	56.166	37.564	1.00	57.75
2316	CD1	LEU	254	113.718	54.785	38.021	1.00	58.43
2317	CD2	LEU	254	113.343	56.571	36.328	1.00	55.32
2318	H	LEU	254	116.252	57.902	38.125	1.00	25.00
2319	N	ASP	255	114.947	59.661	40.475	1.00	73.64
2320	CA	ASP	255	114.842	60.490	41.670	1.00	78.31
2321	C	ASP	255	113.664	60.123	42.574	1.00	79.28
2322	O	ASP	255	113.079	60.985	43.230	1.00	81.76
2323 2324	CB CG	ASP ASP	255 255	114.777 115.238	61.971	41.261 42.364	1.00	83.27 87.17
2324	OD1	ASP ASP	255 255	115.238	62.915 62.443	42.304	$\frac{1.00}{1.00}$	86.98
2325	OD2	ASP	255 255	115.121	64.144	42.165	1.00	87.49
2327	H H	ASP	255	115.805	59.226	40.294	1.00	25.00
2328	N	PHE	256	113.357	58.834	42.659	1.00	80.00
2329	CA	PHE	256	112.254	58.378	43.500	1.00	84.32
2330	C	PHE	256	112.504	58.649	44.976	1.00	89.15
2331	O	PHE	256	111.562	58.742	45.759	1.00	87.59
2332	CB	PHE	256	111.987	56.887	43.290	1.00	81.00
2333	CG	PHE	256	111.352	56.566	41.972	1.00	76.86
2334	CD1	PHE	256	110.671	57.544	41.251	1.00	76.59
2335	CD2	PHE	256	111.431	55.283	41.449	1.00	73.54
2336 2337	CE1	PHE PHE	256	110.080	57.247	40.030 40.229	1.00	76.03
2338	CE2 CZ	PHE	256 256	110.844 110.167	54.976 55.958	39.518	$\frac{1.00}{1.00}$	71.61 74.95
2339	H	PHE	256	113.874	58.189	42.139	1.00	25.00
2340	N	VAL	257	113.774	58.796	45.344	1.00	97.79
2341	CA	VAL	257	114.160	59.053	46.730	1.00	104.36
2342	C	VAL	257	113.428	60.269	47.303	1.00	105.53
2343	O	VAL	257	112.952	60.239	48.439	1.00	106.91
2344	CB	VAL	257	115.692	59.270	46.854	1.00	107.20
2345	CG1	VAL	257	116.092	59.432	48.316	1.00	107.16
2346	CG2	VAL	257	116.445	58.101	46.220	1.00	106.33
2347	H	VAL	257	114.464	58.732	44.658	1.00	25.00
2348 2349	N CA	THR THR	258 258	113.332 112.660	61.329 62.550	46.5506 46.935	$\frac{1.00}{1.00}$	105.84 104.41
2350	C	THR	258	111.183	62.608	46.531	1.00	104.41
2351	Ö	THR	258	110.339	63.047	47.314	1.00	104.07
2352	CB	THR	258	113.392	63.805	48.408	1.00	104.30
2353	OG1	THR	258	113.620	63.674	45.000	1.00	104.41
2354	CG2	THR	258	114.729	63.982	47.117	1.00	105.42
2355	H	THR	258	113.711	61.308	45.603	1.00	25.00
2356	HG1	THR	258	114.068	64.458	44.669	1.00	25.00
2357	N	THR	259	110.872	62.161	45.317	1.00	103.41
2358	CA	THR	259	109.497	62.182	44.825	1.00	100.89
2359	С	THR	259	108.599	61.133	45.482	1.00	101.97
2360	O	THR	259	107.414	61.375	45.707	1.00	103.53
2361 2362	CB OG1	THR THR	259 259	109.445 110.219	62.022	43.289 42.894	1.00	97.37 95.69
2362	CG2	THR	259 259	110.219	60.883 63.267	42.894 42.602	1.00 $1.00$	95.69 95.50
2364	H	THR	259 259	111.579	61.811	44.728	1.00	25.00
2365	HG1	THR	259	109.870	60.086	43.290	1.00	25.00
2366	N	LEU	260	109.164	59.969	45.783	1.00	102.18
2367	CA	LEU	260	108.415	58.884	48.412	1.00	103.22
2368	С	LEU	260	109.112	58.431	47.696	1.00	106.86

TABLE 11-continued

		In the	e Absence of	f Bound St	ıbstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
2369	O	LEU	260	109.742	57.372	47.732	1.00	107.73
2370	CB	LEU	260	108.282	57.700	45.445	1.00	98.81
2371	CG	LEU	260		57.928	44.119	1.00	94.81
2372	CD1	LEU	260	107.620	56.670	43.269	1.00	89.99
2373	CD2	LEU	260 260		58.320	44.380	1.00 1.00	92.83
2374 2375	H N	LEU PRO	260 261	110.112 108.963	59.844	45.595 48.784	1.00	25.00 110.17
2376	CA	PRO	261	109.580		50.077	1.00	110.17
2377	C	PRO	261		57.726	50.855	1.00	114.04
2378	Ö	PRO	261		57.813	52.073	1.00	117.31
2379	СВ	PRO	261	109.441	60.207	50.836	1.00	113.40
2380	CG	PRO	261	108.124	60.715	50.347	1.00	112.70
2381	CD	PRO	261	108.223	60.477	48.856	1.00	111.17
2382	N	TYR	262	108.599		50.163	1.00	113.40
2383	CA	TYR	262	108.012	55.479	50.822	1.00	112.08
2384	C	TYR	262	108.608	54.178	50.284	1.00	112.01
2385	O	TYR	262	108.125	53.086	50.582	1.00	110.30
2386	CB	TYR	262	106.477		50.702	1.00	109.31
2387 2388	CG CD1	TYR TYR	262 262	105.931 105.777	55.286	49.303 48.440	1.00 1.00	104.93 103.18
2389	CD1	TYR	262	105.555	54.021	48.846	1.00	102.84
2390	CE1	TYR	262	105.262		47.159	1.00	101.09
2391	CE2	TYR	262	105.040		47.565	1.00	100.35
2392	CZ	TYR	262		54.938	46.727	1.00	100.07
2393	OH	TYR	262		54.781	45.459	1.00	97.25
2394	H	TYR	262	103.735	56.615	49.199	1.00	25.00
2395	HH	TYR	262	104.276	53.862	45.236	1.00	25.00
2396	N	ALA	263		54.310	49.497	1.00	113.11
2397	CA	ALA	263	110.360		48.913	1.00	114.09
2398	C	ALA	263	111.856		49.146	1.00	114.75
2399	O	ALA	263	112.375	54.457	49.055	1.00	115.90
2400	СВ	ALA	263 263	110.064		47.428	1.00 1.00	112.96
2401 2402	H N	ALA ARG	264	110.034 112.543	52.252	49.305 49.467	1.00	25.00 113.69
2403	CA	ARG	264	113.979		49.726	1.00	114.09
2404	C	ARG	264	114.847	51.948	48.526	1.00	109.03
2405	Ö	ARG	264	114.394		47.594	1.00	109.80
2406	СВ	ARG	264		51.448	50.945	1.00	116.51
2407	CG	ARG	264	113.434		51.235	1.00	118.71
2408	CD	ARG	264	113.486	49.190	50.151	1.00	120.73
2409	NE	ARG	264	112.543	48.105	50.418	1.00	118.02
2410	CZ	ARG	264	111.607	47.696	49.565	1.00	115.79
2411	NH1	ARG	264	111.479	48.277	48.380	1.00	114.07
2412	NH2	ARG	264	110.788	46.709	49.904	1.00	113.33
2413	H	ARG	264	112.079	51.393	49.495	1.00	25.00
2414 2415	HE 111111	ARG	264	112.602	47.651	51.284 48.120	1.00	25.00 25.00
2415 2416	1HH1 2HH1	ARG ARG	264 264	112.083 110.766	49.029 47.967	47.751	1.00 $1.00$	25.00
2417	1HH2	ARG	264	110.766	46.283	50.807	1.00	25.00
2418	2HH2	ARG	264	110.000	46.408	49.273	1.00	25.00
2419	N	ASP	265	116.089	52.421	48.550	1.00	103.18
2420	CA	ASP	265	117.045	52.152	47.485	1.00	97.60
2421	С	ASP	265	117.480	50.688	47.579	1.00	92.70
2422	O	ASP	265	118.533	50.371	48.138	1.00	96.57
2423	CB	ASP	265	118.255	53.086	47.619	1.00	100.16
2424	CG	ASP	265	1199.317	52.826	46.565	1.00	106.00
2425	OD1	ASP	265	118.992	52.887	45.363	1.00	107.25
2426	OD2	ASP	265	120.479	52.556	46.940	1.00	109.23
2427	H	ASP	265	116.373	52.965	49.310	1.00	25.00
2428	N	ARG	266	116.654	49.799	47.041	1.00	82.19
2429 2430	CA	ARG ARG	266 266	116.942	48.372	47.073 45.775	$\frac{1.00}{1.00}$	73.27 63.23
2430	C O	ARG	266	117.613 117.711	47.910 46.712	45.773	1.00	66.31
2432	СВ	ARG	266	115.640	47.585	47.320	1.00	77.54
2433	CG	ARG	266	115.801	46.402	48.274	1.00	83.80
2434	CD	ARG	266	114.480	45.672	48.520	1.00	86.40
2435	NE	ARG	266	114.015	44.911	47.358	1.00	87.97
2436	CZ	ARG	266	114.383	43.661	47.077	1.00	87.83
2437	NH1	ARG	266	115.229	43.014	47.869	1.00	86.43
2438	NH2	ARG	266	113.895	43.049	46.005	1.00	80.67
2439	H	ARG	266	115.816	50.112	46.631	1.00	25.00
2440	HE	ARG	266	113.386	45.348	46.747	1.00	25.00
2441	1HH1	ARG	265	115.600	43.463	48.681	1.00	25.00
2442	2HH1	ARG	266	115.502	42.079	47.647	1.00	25.00

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
2443	1HH2	ARG	266	113.246	43.525	45.410	1.00	25.00
2444	2HH2	ARG	266	114.170	42.111	45.794	1.00	25.30
2445 2446	N CA	VAL VAL	267 267	118.130 118.778	43.859 48.560	45.000 43.722	$\frac{1.00}{1.00}$	54.00 46.85
2447	C	VAL	267	119.835	47.496	43.784	1.00	44.14
2448	Ö	VAL	267	119.311	46.609	42.935	1.00	45.14
2449	CB	VAL	267	119.361	49.827	43.076	1.00	45.45
2450	CG1	VAL	267	113.991	49.499	41.733	1.00	42.99
2451	CG2	VAL	267	118.273	50.848	42.892	1.00	51.05
2452 2453	H N	VAL	267 268	118.072 120.738	49.785 47.576	45.299 44.781	1.00 1.00	25.00
2455 2454	CA	VAL VAL	268	121.813	46.597	44.781	1.00	40.56 38.21
2455	C	VAL	268	121.242	45.185	45.125	1.00	35.40
2456	O	VAL	268	121.708	44.220	44.513	1.00	29.56
2457	CB	VAL	268	122.785	46.976	46.046	1.00	40.34
2458	CG1	VAL	268	123.983	46.055	46.040	1.00	41.41
2459	CG2	VAL	268	123.239	48.414	45.880 45.428	1.00	43.02
2460 2461	H N	VAL GLU	268 269	120.663 120.202	48.303 45.081	45.952	$\frac{1.00}{1.00}$	25.00 33.49
2462	CA	GLU	2669	119.553	43.796	46.220	1.00	31.75
2463	C	GLU	269	118.910	43.278	44.936	1.00	31.29
2464	O	GLU	269	119.023	42.095	44.607	1.00	34.74
2465	CB	GLU	269	118.477	43.940	47.300	1.00	30.42
2466	CG	GLU	269	118.998	44.124	48.719	1.00	41.96
2467 2468	CD OE1	GLU GLU	269 26 <b>P</b>	119.777 119.387	45.418 46.465	48.921 48.356	$\frac{1.00}{1.00}$	53.70 52.14
2469	OE2	GLU	269	120.785	45.385	49.658	1.00	60.44
2470	H	GLU	269	119.849	45.888	46.369	1.00	25.00
2471	N	CYS	270	118.258	44.179	44.204	1.00	25.87
2472	CA	CYS	270	117.603	43.829	42.948	1.00	31.04
2473	С	CYS	270	118.628	43.330	41.944	1.00	31.66
2474 2475	O CB	CYS CYS	270 270	118.352 116.841	42.406 45.029	41.170 42.380	1.00 $1.00$	34.14 31.00
2476	SG	CYS	270	115.468	45.566	43.429	1.00	41.23
2477	Н	CYS	270	118.204	45.102	44.523	1.00	25.00
2478	N	TYR	271	119.817	43.927	41.968	1.00	30.55
2479	CA	TYR	271	120.875	43.506	41.065	1.00	29.41
2480	С	TYR	271	121.365	42.109	41.459	1.00	29.51
2481 2482	O CB	TYR	271	121.662	41.284	40.592	1.00	30.16
2483	CG	TYR TYR	271 271	122.048 123.125	44.496 44.061	41.048 40.077	1.00 $1.00$	27.85 27.35
2484	CD1	TYR	271	123.006	44.327	38.714	1.00	25.72
2485	CD2	TYR	271	124.198	43.279	40.502	1.00	20.62
2486	CE1	TYR	271	123.923	43.813	37.798	1.00	22.98
2487	CE2	TYR	271	125.116	42.762	39.593	1.00	27.42
2488 2489	CZ	TYR	271	124.970 125.859	43.030 42.497	38.245	1.00	21.79
2489 2490	OH H	TYR TYR	271 271	119.985	44.662	37.343 42.594	$\frac{1.00}{1.00}$	23.43 25.00
2491	HH	TYR	271	126.485	41.987	37.843	1.00	25.00
2492	N	PHE	272	121.453	41.845	42.760	1.00	26.47
2493	CA	PHE	272	121.892	40.535	43.220	1.00	28.71
2494	C	PHE	272	120.957	39.486	42.633	1.00	31.32
2495	O	PHE	272	121.408	38.470	42.102	1.90	31.50
2496 2497	CB CG	PHE PHE	272 272	121.881 122.165	40.442 39.058	44.747 45.264	$\frac{1.00}{1.00}$	32.54 34.15
2498	CD1	PHE	272	123.471	38.577	45.323	1.00	32.37
2499	CD2	PHE	272	121.120	38.211	45.638	1.00	34.09
2500	CE11	PHE	272	123.732	37.271	45.739	1 00	35.66
2501	CE2	PHE	272	121.369	36.902	46.055	1.00	34.18
2502	CZ	PHE	272	122.679	36.431	46.105	1.00	36.58
2503 2504	H N	PHE TRP	272 273	121.228 119.656	42.541 39.744	43.413 42.712	$\frac{1.00}{1.00}$	25.00 29.73
2505	CA	TRP	273	118.670	38.817	42.712	1.00	30.60
2506	C	TRP	273	118.924	38.551	40.685	1.00	30.33
2507	ŏ	TRP	273	118.971	37.396	40.250	1.00	32.10
2508	CB	TRP	273	117.255	39.357	42.365	1.00	28.17
2509	CG	TRP	273	116.707	39.092	43.721	1.00	33.41
2510	CD1	TRP	273	117.241	39.478	44.915	1.00	37.71
2511 2512	CD2 NE1	TRP TRP	273 273	115.506 116.445	38.381 39.053	44.029 45.950	1.00 $1.00$	41.82 39.11
2513	CE2	TRP	273	115.372	38.378	45.435	1.00	43.90
2514	CE3	TRP	273	114.528	37.747	43.253	1.00	47.13
2515	CU	TRP	273	114.296	37.764	46.083	1.00	49.35
2516	CZ3	TRP	273	113.458	37.136	43.898	1.00	53.38

TABLE 11-continued

		In the	e Absence of			оно Бунина		
Atom Type	Atom	Residue	Residue #	X	Y	Z	occ	B-factor
2517	CH2	TRP	273	113.352	37.150	45.300	1.00	53.17
2518 2519	H HE1	TRP TRP	273 273	119.359	40.569	43.153	$\frac{1.00}{1.00}$	25.00 25.00
2520	N N	ALA	274	116.622 119.117	39.201 39.617	46.903 39.915	1.00	26.20
2521	CA	ALA	274	119.371	39.472	38.489	1.00	25.12
2522	C	ALA	274	120.638	38.657	38.263	1.00	27.03
2523	0	ALA	274	120.686	37.816	37.366	1.00	29.08
2524	CB	ALA	274	119.491	40.832	37.831	1.00	22.68
2525	H	ALA	274	119.0884		40.313	1.00	25.00
2526	N	LEU	275	121.648	38.886	39.104	1.00	26.74
2527	CA	LEU	275	122.922	38.175	39.011	1.00	23.65
2528 2529	C O	LEU LEU	275 275	122.727 123.432	36.689	39.329 38.798	1.00 $1.00$	20.19 18.69
2530	СВ	LEU	275	123.432	35.825 38.802	39.963	1.00	22.77
2531	CG	LEU	275	125.377	38.280	39.867	1.00	22.23
2532	CD1	LEU	275	125.859	38.352	38.427	1.00	20.45
2533	CD2	LEU	275	126.274	39.097	40.779	1.00	24.10
2534	H	LEU	275	121.532	39.552	39.808	1.00	25.00
2535	N	GLY	276	121.765	36.406	40.204	1.00	19.17
2536	CA	GLY	276	121.453	35.035	40.561	1.00	19.61
2537	С	GLY	276	120.811	34.299	39.392	1.00	25.48
2538	O	GLY	276	121.060	33.108	39.199	1.00	28.59
2539 2540	H N	GLY VAL	276 277	121.266 120.000	37.137 35.006	40.623 38.603	$\frac{1.00}{1.00}$	25.00 20.84
2541	CA	VAL VAL	277	119.323	34.415	37.440	1.00	19.19
2542	C	VAL	277	120.304	34.028	36.319	1.00	19.17
2543	ŏ	VAL	277	120.086	33.048	35.606	1.00	21.35
2544	CB	VAL	277	118.201	35.355	36.904	1.00	21.19
2545	CG1	VAL	277	117.560	34.777	35.650	1.00	15.33
2546	CG2	VAL	277	117.138	35.550	37.976	1.00	11.13
2547	H	VAL	277	119.858	35.954	38.810	1.00	25.00
2548	N	TYR	273	121.345	34.834	36.137	1.00	21.16
2549 2550	CA C	TYR TYR	278 278	122.401 123.583	34.587 35.532	35.150 35.351	1.00 1.00	24.77 29.35
2551	o	TYR	278	123.405	36.738	35.531	1.00	27.69
2552	СВ	TYR	278	121.910	34.611	33.687	1.00	25.96
2553	CG	TYR	278	120.741	35.517	33.341	1.00	26.89
2554	CD1	TYR	278	120.580	36.768	33.937	1.00	25.88
2555	CD2	TYR	278	119.800	35.116	32.388	1.00	27.14
2556	CE1	TYR	278	119.508	37.595	33.594	1.00	29.19
2557	CE2	TYR	278	118.729	35.934	32.037	1.00	31.49
2558 2559	CZ OH	TYR	278	118.587	37.171	32.643	1.00	31.78
2560	Н	TYR TYR	278 278	117.522 121.419	37.976 35.652	32.301 36.684	1.00 $1.00$	31.66 25.00
2561	HH	TYR	278	117.022	37.548	31.600	1.00	25.00
2562	N	PHE	279	124.789	34.968	35.332	1.00	31.85
2563	CA	PHE	279	126.017	35.732	35.549	1.00	28.28
2564	C	PHE	279	126.910	35.844	34.318	1.00	28.86
2565	O	PHE	279	127.855	36.636	34.310	1.00	28.24
2566	CB	PHE	279	126.829	35.087	36.678	1.00	24.38
2567	CG CD1	PHE	279	127.334	33.707	36.344	1.00	23.04
2568 2569	CD1 CD2	PHE	279 279	128.563 126.557	33.535 32.582	35.706 36.616	$\frac{1.00}{1.00}$	16.44 22.61
2570	CE1	PHE PHE	279	120.337	32.265	35.339	1.00	22.82
2571	CE2	PHE	279	126.989	31.309	36.254	1.00	22.81
2572	CZ	PHE	279	128.214	31.149	35.613	1.00	20.63
2573	H	PHE	279	124.845	34.006	35.181	1.00	25.00
2574	N	GLU	280	126.653	35.010	33.315	1.00	25.83
2575	CA	GLU	280	127.450	34.995	32.093	1.00	25.96
2576	C	GLU	280	127.464	36.347	31.384	1.00	32.09
2577	O	GLU	280	126.461	37.067	31.378	1.00	33.29
2578	CB	GLU GLU	280 280	126.947	33.909 32.479	31.140	1.00	29.64 31.98
2579 2580	CG CD	GLU	280	127.116 125.873	31.921	31.652 32.338	1.00 $1.00$	41.40
2581	OE1	GLU	280	125.089	32.696	32.938	1.00	33.42
2582	OE2	GLU	280	125.681	30.688	32.273	1.00	43.76
2583	H	GLU	280	125.894	34.421	33.402	1.00	25.00
2584	N	PRO	281	128.593	36.687	30.735	1.00	34.75
2585	CA	PRO	281	128.736	37.961	30.018	1.00	32.48
2586	С	PRO	281	127.718	38.182	28.899	1.00	30.11
2587	O	PRO	281	127.273	39.309	28.675	1.00	33.55
2588 2589	CB CG	PRO PRO	281 281	130.177 130.447	37.901 36.426	229.492 29.366	$\frac{1.00}{1.00}$	33.45 34.83
2590	CD	PRO	281	129.824	35.882	30.625	1.00	32.94
_0,0					22.002	23.020	2.00	

TABLE 11-continued

	Struc	tural Coordinat In the	es of Tobac Absence o			ene Synth	ase	
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
2591	N	GLN	282	127.319	37.106	28.226	1.00	30.38
2592	CA	GLN	282	126.350	37.195	27.134	1.00	31.74
2593 2594	C O	GLN	282 282	124.980	37.704	27.600	$\frac{1.00}{1.00}$	33.53
2595	СВ	GLN GLN	282	124.194 126.183	38.202 35.829	26.792 26.462	1.00	34.52 35.52
2596	CG	GLN	282	125.442	34.819	27.323	1.00	47.53
2597	CD	GLN	282	125.543	33.400	26.807	1.00	54.68
2598	CE1	GLN	282	126.378	32.624	27.273	1.00	58.66
2599	NE2	GLN	282	124.675	33.040	25.866	1.00	55.48
2600	Н	GLN	282	127.685	36.233	28.468	1.00	25.00
2601	1HE2	GLLN	282	124.740	32.122	25.538	1.00	25.00
2602 2603	2HE2 N	GLN TYR	282 283	124.018 124.698	33.690 37.577	25.551 28.896	$\frac{1.00}{1.00}$	25.00 27.81
2604	CA	TYR	283	123.417	38.015	29.447	1.00	26.64
2605	C	TYR	283	123.470	39.392	30.099	1.00	29.80
2606	O	TYR	283	122.615	39.732	30.922	1.00	29.85
2607	CB	TYR	283	122.885	36.982	30.444	1.00	24.82
2608	CG	TYR	283	122.670	35.614	29.840	1.00	26.77
2609	CD1	TYR	283	121.786	35.432	28.775	1.00	28.94
2610 2611	CD2 CE1	TYR TYR	283 283	123.373 121.610	34.506 34.174	30.313 28.192	$\frac{1.00}{1.00}$	26.51 32.76
2612	CE2	TYR	283	123.205	33.247	29.740	1.00	28.52
2613	CZ	TYR	283	122.324	33.086	28.680	1.00	30.97
2614	OH	TYR	283	122.164	31.845	28.104	1.00	25.13
2615	H	TYR	283	125.361	37.198	29.509	1.00	25.00
2616	HH	TYR	283	122.723	31.208	28.556	1.00	25.00
2617	N	SER	284	124.449	40.196	29.697	1.00	30.66
2618	CA	SER	284 284	124.620 123.375	41.539	30.239	$\frac{1.00}{1.00}$	33.15 32.37
2619 2620	C O	SER SER	284	123.373	42.412 42.999	30.040 30.999	1.00	32.25
2621	СВ	SER	284	125.848	42.201	29.609	1.00	32.60
2622	OG	SER	284	126.037	43.511	30.110	1.00	38.65
2623	H	SER	284	125.090	39.876	29.027	1.00	25.00
2624	HG	SER	284	126.140	43.520	31.058	1.00	25.00
2625	N	GLN	285	122.882	42.477	28.805	1.00	35.28
2626	CA	GLN	285	121.693	43.273	28.505	1.00	36.59
2627 2628	C O	GLN GLN	285 285	120.489 119.713	42.735 43.504	29.284 29.856	$\frac{1.00}{1.00}$	33.06 33.43
2629	СВ	GLN	285	121.399	43.255	27.002	1.00	36.57
2630	CG	GLN	285	120.138	44.020	26.611	1.00	48.54
2631	CD	GLN	285	119.829	43.943	25.123	1.00	54.77
2632	OE1	GLN	285	120.079	42.927	24.470	1.00	55.77
2633	NE2	GLN	285	119.280	45.024	24.581	1.00	56.33
2634	H 1HE2	GLN	285 285	123.323 119.084	41.984 44.982	28.086 23.621	1.00	25.00 25.00
2635 2636	1HE2 2HE2	GLN GLN	285	119.004	45.802	25.141	$\frac{1.00}{1.00}$	25.00
2637	N	ALA	286	120.364	41.410	29.324	1.00	32.38
2638	CA	ALA	286	119.272	40.750	30.032	1.00	28.79
2639	C	ALA	286	119.254	41.117	31.512	1.00	26.70
2640	O	ALA	286	118.200	41.438	32.060	1.00	31.71
2641 2642	CB H	ALA ALA	286 286	119.370 121.023	39.244 40.859	29.859 28.861	$\frac{1.00}{1.00}$	30.35 25.00
2643	N	ARG	287	120.422	41.097	32.152	1.00	26.48
2644	CA	ARG	287	120.517	41.442	33.568	1.00	27.31
2645	С	ARG	2871	20.056	42.870	33.826	1.00	27.08
2646	O	ARG	2871	19.290	43.118	34.760	1.00	28.78
2647	CB	ARG	2871	21.946	41.266	34.096	1.00	28.17
2648	CG	ARG	2871	22.240	39.891	34.652	1.00	25.50
2649 2650	CD NE	ARG ARG	2871 2871	23.566 24.703	39.859 40.191	35.396 34.535	1.00 1.00	24.46 20.40
2651	CZ	ARG	2871	25.252	39.373	33.641	1.00	23.12
2652	NH1	ARG	2871	24.781	38.146	33.473	1.00	23.32
2653	NH2	ARG	2871	26.268	39.793	32.897	1.00	22.53
2654	H	ARG	2871	21.232	40.840	31.670	1.00	25.00
2655	HE	ARG	2971	25.093	41.059	34.627	1.00	25.00
2656 2657	1HH1	ARG	2871	24.002	37.833	34.014	1.00	25.00
2657 2658	2HH1 1HH2	ARG ARG	2871 2871	25.192 26.623	37.542 40.719	32.796 33.014	$\frac{1.00}{1.00}$	25.00 25.00
2659	2HH2	ARG	2871	26.677	39.179	32.225	1.00	25.00
2660	N	VAL	2881	20.512	43.802	32.992	1.00	30.28
2661	CA	VAL	2881	20.144	45.208	33.148	1.00	29.94
2662	C	VAL	2881	18.628	45.388	33.043	1.00	28.55
2663	O	VAL	2881	18.018	46.044	33.890	1.00	35.55
2664	СВ	VAL	2881	20.874	46.106	32.120	1.00	35.29

TABLE 11-continued

Atom Type	Atom	Residue	Residue #	X	Y	Z	occ	B-factor
2665	CG1	VAL	2881	20.536	47.572	32.363	1.00	30.30
2666	CG2	VAL	2881	22.378	45.896	32.221	1.00	31.86
2667	H	VAL	2881	21.107	43.535	32.256	1.00	25.00
2668	N	MET	2891	18.018	44.775	32.031	1.00	27.91
2669	CA	MET	2891	16.567	44.856	31.854	1.00	27.36
2670	C	MET	2891	15.857	44.248	33.066	1.00	27.49
2671	O	MET	2891	14.938	44.845	33.627	1.00	29.43
2672	CB	MET	2891	16.136	44.129	30.572	1.00	28.18
2673	CG	MET	2891	16.578	44.819	29.282	1.00	28.82
2674	SD	MET	2891	16.207	43.882	27.770	1.00	38.33
2675	CE	MET	2891	14.526	44.384	27.438	1.00	39.91
2676	H	MET	2891	18.554	44.257	31.391	1.00	25.00
2677	N	LEU	2901	16.335	43.089	33.511	1.00	25.99
2678	CA	LEU	2901	15.743	42.408	34.654	1.00	25.33
2679	С	LEU	2901	15.805	43.222	35.949	1.00	26.44
2680	O	LEU	2901	14.815	43.289	36.687	1.00	30.04
2681	CB	LEU	2901	16.393	41.035	34.843	1.00	25.26
2682	CG	LEU	2901	15.880	40.125	35.964	1.00	25.88
2683	CD1	LEU	2901	14.357	40.043	35.951	1.00	19.53
2684	CD2	LEU	2901	16.499	38.741	35.796	1.00	18.49
2685	H	LEU	2901	17.098	42.681	33.057	1.00	25.00
2686	N	VAL	2911	16.947	43.857	36.210	1.00	27.54
2687	CA	VAL	2911	17.124	44.667	37.421	1.00	28.37
2688	C	VAL	2911	16.101	45.799	37.502	1.00	27.52
2689	O	VAL	2911	15.487	46.023	38.5550	1.00	27.61
2690	CB	VAL	2911	18.544	45.289	37.507	1.00	28.94
2691	CG1	VAL	2911	18.706	46.054	38.803	1.00	25.65
2692	CG2	VAL	2911	19.592	44.214	37.431	1.00	36.38
2693	H	VAL	2911	17.687	43.782	35.573	1.00	25.00
2694	N	LYS	2921	15.911	46.502	36.392	1.00	27.46
2695	CA	LYS	2921	14.968	47.611	36.345	1.00	28.57
2696	С	LYS	2921	13.546	47.158	36.677	1.00	30.77
2697	O	LYS	292	112.834	47.824	37.433	1.00	31.18
2698	CB	LYS	292	115.029	48.285	34.976	1.00	29.35
2699	CG	LYS	292	116.391	48.890	34.676	1.00	29.57
2700	CD	LYS	292	116.463	49.431	33.261	1.00	34.35
2701	CE	LYS	292	117.810	50.079	32.999	1.00	37.92
2702	NZ	LYS	292	117.909	50.619	31.619	1.00	40.64
2703	H	LYS	292	116.413	46.260	35.581	1.00	25.00
2704	1HZ	LYS	292	117.780	49.846	30.936	1.00	25.00
2705	2HZ	LYS	292	117.139	51.334	31.476	1.00	25.00
2706	3HZ	LYS	292	118.644	51.052	31.483	1.00	25.00
2707	N	THR	293	113.143	48.010	36.137	1.00	34.24
2708	CA	THR	293	111.817	45.483	36.395	1.00	27.43
2709	С	THR	293	111.657	45.123 45.493	37.872	1.00	31.58
2710	O	THR	293	110.655		38.491	1.00	28.71
2711	CB	THR	293	111.561	44.214	35.534	1.00	25.43
2712	OG1	THR	293	111.354	44.616	34.175	1.00	30.49
2713	CG2	THR	293	110.348	43.433	36.029	1.00	22.44
2714	H HG1	THR	293 293	113.756	45.525 45.180	35.533 34.129	1.00 $1.00$	25.00
2715	HG1	THR ILE	293 294	110.577		34.129 38.440		25.00 30.35
2716	N C4		294 294	112.647	44.439 44.064		1.00	
2717	CA	ILE	294 294	112.596		39.853 40.725	1.00	30.45
2718	С	ILE ILE	294 294	112.481	45.317	40.725	1.00	28.69 30.57
2719	O CB		294 294	111.709	45.348	41.685	1.00	
2720	CB CG1	ILE		113.837	43.230	40.272 39.399	1.00	29.95
2721	CG1	ILE	294	113.948	41.977 42.818	39.399 41.734	1.00	24.34
2722 2723	CG2 CD1	ILE ILE	294 294	113.733	42.818	41.734 39.687	1.00	18.35 30.84
2723	H H	ILE	294 294	115.165	41.133	39.687 37.906	1.00 $1.00$	25.00
				113.432				
2725	N C4	SER	295	113.219	46.361	40.359	1.00	32.61
2726	CA	SER	295 295	113.196	47.623	41.097	1.00	37.07 35.65
2727	С	SER	295 205	111.820	48.276	41.002	1.00	35.65
2728	O	SER	295	111.227	48.673	42.011	1.00	33.09
2729	CB	SER	295	114.246	48.584	40.533	1.00	35.83
2730	OG H	SER	295	115.543	48.024	40.608	1.00	42.38
2731	H	SER	295	113.804	46.290	39.575	1.00	25.00
2732	HG	SER	295	115.756	47.823	41.523	1.00	25.00
2733	N	MET	296	111.306	48.342	39.779	1.00	34.54
2734	CA	MET	296	110.017	48.951	39.506	1.00	35.02
				108.864	48.263	40.230	1.00	36.33
2735	С	MET	296					
2735 2736	O	MET	296	108.080	48.919	40.923	1.00	35.61
2735								

TABLE 11-continued

		In the	e Absence o	f Bound St	ıbstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
2739	SD	MET	296	109.993	51.708	38.067	1.00	51.57
2740	CE	MET	296	108.888	52.271	39.359	1.00	53.40
2741	H	MET	296	111.819	47.966	39.036	1.00	25.00
2742	N	ILE	297	108.780	46.941	40.103	1.00	37.42
2743	CA	ILE	297	107.709	46.185	40.745	1.00	34.54
2744	C	ILE	297	107.813	46.267	42.267	1.00	36.45
2745	O	ILE	297	106.817	46.101	42.976	1.00	35.56
2746	CB	ILE	297	107.675	44.709	40.265	1.00	36.67
2747	CG1	ILE	297	106.342	44.059	40.648	1.00	30.90
2748	CG2	ILE	297	108.858	43.925	40.821	1.00	34.15
2749	CD1	ILE	297	105.135	44.654	39.935	1.00	28.47
2750	H	ILE	297	109.454	46.468	39.576	1.00	25.00
2751	N	SER	298	109.015	46.559	42.761	1.00	37.95
2752	CA	SER	298 298	109.250	46.696	44.195	1.00	40.22
2753 2754	C O	SER SER	298 298	108.531 107.934	47.961 47.979	44.673 45.753	$\frac{1.00}{1.00}$	38.96 36.43
2755	СВ	SER	298 298	1107.934	46.791	44.481	1.00	43.72
2756	OG	SER	298 298	111.009	46.764	46.873	1.00	62.08
2757	H	SER	298	109.775	46.674	42.153	1.00	25.00
2758	HG	SER	298	110.573	47.504	46.304	1.00	25.00
2759	N	ILE	299	108.582	49.011	43.857	1.00	39.57
2760	CA	ILE	299	107.912	50.271	44.175	1.00	40.91
2761	C	ILE	299	106.412	49.996	44.293	1.00	40.75
2762	ŏ	ILE	299	105.771	50.378	45.276	1.00	40.26
2763	СВ	ILE	299	108.128	51.329	43.060	1.00	37.27
2764	CG1	ILE	299	109.614	51.653	42.908	1.00	37.90
2765	CG2	ILE	299	1077.345	52.592	43.370	1.00	42.54
2766	CD1	ILE	299	110.260	52.146	44.173	1.00	39.79
2767	H	ILE	299	109.091	48.942	43.021	1.00	25.00
2768	N	VAL	300	105.876	49.290	43.301	1.00	34.47
2769	CA	VAL	300	104.462	48.949	43.267	1.00	33.70
2770	C	VAL	300	104.050	48.145	44.497	1.00	38.81
2771	O	VAL	300	103.020	48.431	45.116	1.00	39.02
2772	CB	VAL	300	104.116	48.166	41.990	1.00	35.45
2773	CG1	VAL	300	102.629	47.848	41.951	1.00	37.16
2774	CG2	VAL	300	104.522	48.970	40.762	1.00	29.01
2775	H	VAL	300	106.459	48.994	42.572	1.00	25.00
2776	N	ASP	301	104.866	47.162	44.865	1.00	39.28
2777	CA	ASP	301	104.585	46.327	46.030	1.00	39.76
2778	С	ASP	301	104.477	47.200	47.281	1.00	43.93
2779	O	ASP	301	103.588	47.002	48.113	1.00	43.98
2780	CB	ASP	301	105.684	45.270 44.299	46.205	1.00	41.93
2781 2782	CG OD1	ASP ASP	301 301	105.401 104.219	44.299	47.348 47.633	$\frac{1.00}{1.00}$	47.18 52.50
2783	OD1 OD2	ASP	301	104.219	43.817	47.033	1.00	53.58
2784	H	ASP	301	105.672	46.984	44.338	1.00	25.00
2785	N	ASP	302	105.373	48.175	47.401	1.00	48.10
2786	CA	ASP	302	105.371	49.088	48.541	1.00	51.62
2787	C	ASP	302	104.090	49.918	48.560	1.00	50.46
2788	Ö	ASP	302	103.480	50.114	49.615	1.00	51.17
2789	CB	ASP	302	106.587	50.017	48.487	1.00	55.75
2790	CG	ASP	302	107.904	49.271	48.619	1.00	62.15
2791	OD1	ASP	302	107.922	48.165	49.207	1.00	63.08
2792	OD2	ASP	302	108.928	49.798	48.133	1.00	68.25
2793	H	ASP	302	106.056	48.281	46.704	1.00	25.00
2794	N	THR	303	103.684	50.388	47.383	1.00	50.52
2795	CA	THR	303	102.479	51.192	47.230	1.00	50.05
2796	C	THR	303	101.260	50.472	47.808	1.00	51.84
2797	O	THR	303	100.563	51.013	48.668	1.00	58.07
2798	CB	THR	303	102.222	51.512	45.745	1.00	50.09
2799	OG1	THR	303	103.377	52.153	45.190	1.00	45.79
2800	CG2	THR	303	101.015	52.425	45.593	1.00	50.31
2801	H	THR	303	104.219	50.190	46.581	1.00	25.00
2802	HG1	THR	303	103.480	52.938	45.719	1.00	25.00
2803	N C4	PHE	304	101.025	49.246	47.352	1.00	50.29
2804	CA	PHE	304	99.893 99.997	48.450	47.817	1.00	53.29
2805	C O	PHE PHE	304 304	98.981	48.024	49.275 49.940	1.00	59.68 62.17
2806 2807	CB	PHE	304 304	98.981	47.832 47.182	46.971	$\frac{1.00}{1.00}$	43.62
2808	CG CG	PHE	304	99.744	47.182	45.654	1.00	37.53
2809	CD1	PHE	304	99.063	47.857	44.555	1.00	32.17
2810	CD2	PHE	304	97.711	47.113	45.506	1.00	35.92
2811	CE1	PHE	304	99.158	48.029	43.326	1.00	36.26
2812	CE2	PHE	304	97.079	47.280	44.283	1.00	29.26

TABLE 11-continued

		In the	e Absence o			ene Synth		
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
2813	CZ	PHE	304	97.802	47.739	43.189	1.00	34.73
2814	H	PHE	304	101.636	48.860	46.683	1.00	25.00
2815 2816	N CA	ASP ASP	305 305	101.223 101.450	47.873 47.405	49.765 51.129	$\frac{1.00}{1.00}$	69.36 78.46
2817	C	ASP	305	101.436	48.406	52.279	1.00	80.92
2818	Ö	ASP	305	100.774	48.064	53.329	1.00	79.13
2819	CB	ASP	305	102.798	46.675	51.210	1.00	84.84
2820	CG	ASP	305	102.851	45.663	52.345	1.00	90.63
2821	OD1	ASP	305	102.142	44.635	52.265	1.00	90.70
2822	OD2	ASP	305	103.610	45.891	53.312	1.00	92.23
2823	H	ASP	305	101.996	48.072	49.197	1.00	25.00
2824 2825	N CA	ALA ALA	306 306	101.818 101.752	49.631 50.595	52.104 53.201	$\frac{1.00}{1.00}$	84.70 89.65
2825	C	ALA	306	101.752	52.057	52.874	1.00	91.11
2827	Ö	ALA	306	101.606	52.916	53.745	1.00	93.41
2828	CB	ALA	306	103.027	50.500	54.040	1.00	89.50
2829	H	ALA	306	102.229	49.879	51.249	1.00	25.00
2830	N	TYR	307	101.022	52.359	51.655	1.00	90.94
2831	CA	TYR	307	100.743	53.752	51.329	1.00	92.48
2832	С	TYR	307	99.374	54.011	50.701	1.00	90.37
2833	O	TYR	307	98.599	54.824	51.207 50.453	1.00	91.55
2834 2835	CB CG	TYR TYR	307 307	101.858 102.031	54.336 55.837	50.453	$\frac{1.00}{1.00}$	98.27 106.75
2836	CD1	TYR	307	102.031	56.732	49.825	1.00	100.73
2837	CD2	TYR	307	102.918	56.364	51.552	1.00	107.24
2838	CE1	TYR	307	101.447	58.113	49.973	1.00	107.24
2839	CE2	TYR	307	103.072	57.744	51.708	1.00	106.57
2840	CZ	TYR	307	102.332	58.611	50.915	1.00	106.75
2841	OH	TYR	307	102.477	59.971	51.060	1.00	103.97
2842	H	TYR	307	100.876	51.657	50.988	1.00	25.00
2843 2844	HH N	TYR GLY	307 308	101.900 99.079	60.427 53.325	50.443 49.603	1.00	25.00 85.79
2845	CA	GLY	308	97.808	53.523	48.930	$\frac{1.00}{1.00}$	82.54
2846	C	GLY	308	96.583	53.120	49.730	1.00	81.67
2847	ŏ	GLY	308	96.589	52.105	50.428	1.00	80.43
2848	H	GLY	308	99.710	52.666	49.258	1.00	25.00
2849	N	THR	309	95.531	53.928	49.637	1.00	82.42
2850	CA	THR	309	94.282	53.649	50.338	1.00	82.64
2851	C	THR	309	93.397	52.796	49.433	1.00	83.27
2852	O CB	THR	309 309	93.592	52.771	48.215	1.00	87.28
2853 2854	OG1	THR THR	309	93.519 93.166	54.946 55.648	50.696 49.495	$\frac{1.00}{1.00}$	80.48 74.90
2855	CG2	THR	309	94.371	55.844	51.583	1.00	75.37
2856	H	THR	309	95.600	54.729	49.085	1.00	25.00
2857	HG1	THR	309	92.598	55.106	48.954	1.00	25.00
2858	N	VAL	310	92.383	52.172	50.025	1.00	80.18
2859	CA	VAL	310	91.447	51.304	49.309	1.00	75.06
2860	C	VAL	310	91.067	51.822	47.919	1.00	74.05
2861 2862	O CB	VAL VAL	310 310	91.209 93.149	51.115 51.103	46.921 50.127	$\frac{1.00}{1.00}$	73.13 79.21
2863	CG1	VAL	310	89.284	50.020	49.494	1.00	80.28
2864	CG2	VAL	310	90.478	50.760	51.575	1.00	78.07
2865	H	VAL	310	92.275	52.293	50.987	1.00	25.00
2866	N	LYS	311	90.622	53.072	47.859	1.00	73.34
2867	CA	LYS	311	90.210	53.682	46.600	1.00	71.58
2868	C	LYS	311	91.366	53.946	45.639	1.00	67.72
2869	O	LYS	311	91.269 89.433	53.642	44.448	1.00	65.13
2870 2871	CB CG	LYS LYS	311 311	89.433 87.977	54.977 54.774	46.866 47.306	$\frac{1.00}{1.00}$	79.85 89.57
2872	CD	LYS	311	87.842	53.976	48.607	1.00	98.38
2873	CE	LYS	311	88.473	54.694	49.795	1.00	102.39
2874	NZ	LYS	311	87.808	55.997	50.082	1.00	107.12
2875	H	LYS	311	90.572	53.594	48.679	1.00	25.00
2876	1HZ	LYS	311	87.884	56.617	49.250	1.00	25.00
2877	2HZ	LYS	311	86.804	55.833	50.299	1.00	25.00
2878	3HZ	LYS	311	88.268	56.452	50.896	1.00	25.00
2879 2880	N CA	GLU GLU	312 312	92.461 93.634	54.495 54.805	46.162 45.346	1.00	64.41 61.39
2880	CA	GLU	312	93.034	54.805 53.556	45.346 44.667	$\frac{1.00}{1.00}$	61.26
2882	Ö	GLU	312	94.169	53.585	43.483	1.00	59.48
2883	CB	GLU	312	94.724	55.465	46.194	1.00	63.39
2884	CG	GLU	312	94.348	56.830	46.748	1.00	70.79
2885	CD	GLU	312	95.456	57.456	47.578	1.00	75.21
2886	OE1	GLU	312	95.879	56.840	48.579	1.00	77.19

TABLE 11-continued

		In the	Absence of	Bound St	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
2887	OE2	GLU	312	95.903	58.570	47.233	1.00	79.86
2888	H	GLU	312	92.482	54.687	47.112	1.00	25.00
2889	N	LEU	313	94.257	52.459	45.418	1.00	57.55
2890	CA	LEU	313	94.765	51.198	44.891	1.00	52.69
2891	C	LEU	313	93.885	50.678	43.762	1.00	52.82
2892	O	LEU	313	94.391	50.281	42.713	1.00	51.15
2893	CB	LEU	313	94.883	50.158	46.005	1.00	48.63
2894	CG	LEU	313	95.886	50.519	47.102	1.00	46.47
2895	CD1	LEU	313	95.941	49.416	48.140	1.00	48.38
2896	CD2	LEU	313	97.259	50.748	46.495	1.00	48.00
2897	H	LEU	313	93.952	52.495	46.340	1.00	25.00
2898	N	GLU	314	92.569	50.724	43.957	1.00	52.92
2899	CA	GLU	314	91.634	50.269	42.933	1.00	54.92
2900	C	GLU	314	91.840	51.087	41.651	1.00	52.05
2901	O	GLU	314	91.801	50.548	40.541	1.00	49.73
2902	CB	GLU	314	90.189	50.400	43.431	1.00	58.82
2903	CG	GLU	314	89.137	49.809	42.488	1.00	68.28
2904	CD	GLU	314	89.281	48.303	42.292	1.00	74.40
2905	OE1	GLU	314	89.097	47.550	43.275	1.00	76.74
2906	OE2	GLU	314	89.568	47.871	41.152	1.00	74.09
2907	H	GLU	314	92.221	51.057	44.810	1.00	25.00
2908	N	ALA	315	92.088	52.383	41.813	1.00	51.92
2909	CA	ALA	315	92.323	53.270	40.678	1.00	54.57
2910	C	ALA	315	93.649	52.920	39.993	1.00	51.51
2911	O	ALA	315	93.762	52.977	38.763	1.00	49.98
2912	CB	ALA	315	92.335	54.722	41.142	1.00	52.19
2913	H	ALA	315	92.103	52.753	42.723	1.00	25.00
2914	N	TYR	316	04.540	52.542	40.796	1.00	49.90
2915	CA	TYR	316	95.960	52.177	40.289	1.00	46.00
2916	C	TYR	316	95.911	50.864	39.506	1.00	42.05
2917	O	TYR	316	96.503	50.756	38.424	1.00	36.96
2918	CB	TYR	316	96.954	52.070	41.445	1.00	48.32
2919	CG	TYR	316	98.405	52.154	41.029	1.00	52.17
2920	CD1	TYR	316	98.975	53.371	40.657	1.00	53.66
2921	CD2	TYR	316	99.218	51.023	41.033	1.00	58.41
2922	CE1	TYR	316	100.320	53.461	40.303	1.00	56.28
2923	CE2	TYR	316	100.566	51.101	40.681	1.00	63.22
2924	CZ	TYR	316	101.110	52.323	40.319	1.00	58.59
2925	OH	TYR	316	102.442	52.405	39.986	1.00	51.77
2926	H	TYR	316	94.483	52.514	41.763	1.00	25.00
2927	HH	TYR	316	102.631	53.311	39.758	1.00	25.00
2928	N	THR	317	95.186	49.881	40.040	1.00	38.98
2929	CA	THR	317	95.044	48.574	39.396	1.00	40.08
2930	C	THR	317	94.391	48.732	38.025	1.00	41.19
2931	O	THR	317	94.755	48.046	37.065	1.00	40.64
2932	CB	THR	317	94.189	47.619	40.245	1.00	39.21
2933	OG1	THR	317	94.658	47.632	41.598	1.00	40.99
2934	CG2	THR	317	94.277	46.198	39.698	1.00	41.75
2935	H	THR	317	94.740	50.022	40.900	1.00	25.00
2936	HG1	THR	317	94.120	47.056	42.134	1.00	25.00
2937	N	ASP	318	93.423	49.641	37.945	1.00	46.22
2938	CA	ASP	318	92.719	49.920	36.700	1.00	44.99
2939	C	ASP	318	93.631	50.595	35.693	1.00	38.74
2940	O	ASP	318	93.695	50.183	34.536	1.00	39.12
2941	CB	ASP	318	91.497	50.799	36.959	1.00	55.80
2942	CG	ASP	318	90.215	50.006	36.977	1.00	64.67
2943	OD1	ASP	318	89.924	49.364	38.010	1.00	73.33
2944	OD2	ASP	318	89.507	50.014	35.948	1.00	72.06
2945	H	ASP	318	93.171	50.138	38.755	1.00	25.00
2940	N	ALA	319	94.340	51.628	36.135	1.00	37.74
2947	CA	ALA	319	95.258	52.347	35.260	1.00	40.21
2948	C	ALA	319	96.245	51.360	34.644	1.00	42.98
2949	O	ALA	319	96.528	51.422	33.446	1.00	41.24
2950	CB	ALA	319	95.995	53.426	36.039	1.00	42.19
2951	H	ALA	319	94.240	51.923	37.067	1.00	25.00
2952	N	ILE	320	96.727	50.423	35.462	1.00	41.94
2953	CA	ILE	320	97.670	49.403	35.005	1.00	40.80
2954	C	ILE	320	97.064	48.488	33.934	1.00	37.20
2955	O	ILE	320	97.711	48.200	32.923	1.00	31.26
2956	CB	ILE	320	98.198	48.549	36.191	1.00	40.22
2957	CG1	ILE	320	99.093	49.404	37.091	1.00	38.13
2958	CG2	ILE	320	98.973	47.332	35.680	1.00	36.19
2959	CE1	ILE	320	100.340	49.924	36.392	1.00	35.64
2960	H	ILE	320	96.447	50.429	36.402	1.00	25.00

TABLE 11-continued

Structural	Coordinates	of Tobacco	5-Epi-Aristolo	ochene Synthase
	In the A	bsence of E	Bound Substrat	ie .

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
2961	N	GLN	321	95.830	48.037	34.149	1.00	37.70
2962	CA	GLN	321	95.167	47.161	33.180	1.00	43.79
2963	С	GLN	321	94.959	47.367	31.839	1.00	44.60
2964	O	GLN	321	95.104	47.254	30.777	1.00	43.29
2965	СВ	GLN	321	93.818	46.662	33.713	1.00	45.39
2966	CG	GLN	321	93.879	45.997	35.079	1.00	50.49
2967	CD	GLN	321	94.981	44.963	35.182	1.00	52.12
2968 2969	OE1 NE2	GLN GLN	321 321	95.097 95.801	44.073 45.078	34.341 36.220	1.00 $1.00$	53.96 49.92
2970	H	GLN	321	95.360	48.301	34.967	1.00	25.00
2971	1HE2	GLN	321	96.511	44.405	36.295	1.00	25.00
2972	2HE2	GLN	321	95.664	45.810	36.855	1.00	25.00
2973	N	ARG	322	94.595	49.148	31.894	1.00	48.28
2974	CA	ARG	322	94.376	49.935	30.683	1.00	48.66
2975	C	ARG	322	95.697	50.181	29.976	1.00	48.74
2976	O	ARG	322	95.756	50.167	28.745	1.00	52.54
2977	CB	ARG	322	93.701	51.272	31.003	1.00	53.96
2978	CG	ARG	322	92.175	51.230	31.029	1.00	62.97
2979	CD	ARG	322	91.642	50.367	32.164	1.00	68.65
29880	NE	ARG	322	90.183	50.288	32.167	1.00	71.23
2981 2982	CZ NH1	ARG ARG	322 322	89.377 89.878	51.274 52.432	32.546 32.959	$\frac{1.00}{1.00}$	73.12 75.57
2982	NH2	ARG	322	88.064	51.101	32.512	1.00	75.20
2984	H	ARG	322	94.477	49.570	32.771	1.00	25.00
2985	HE	ARG	322	83.771	49.449	31.873	1.00	25.00
2986	1HH1	ARG	322	90.868	52.574	32.983	1.00	25.00
2987	2HH1	ARG	322	89.263	53.172	33.235	1.00	25.00
2988	1HH2	ARG	322	87.684	50.229	32.203	1.00	25.00
2989	2HH2	ARG	322	87.455	51.842	32.793	1.00	25.00
2990	N	TRP	323	96.740	50.434	30.765	1.00	48.78
2991	CA	TRP	323	98.086	50.674	30.248	1.00	50.53
2992	C	TRP	323	98.038	51.721	29.139	1.00	51.62
2993	0	TRP	323	98.388	51.442	27.984	1.00	46.60
2994	CB	TRP	323	98.678	49.361	29.719	1.00	45.66
2995 2998	CG CD1	TRP TRP	323 323	100.171 100.972	49.334 49.778	29.673 28.660	$\frac{1.00}{1.00}$	43.84 41.07
2997	CD2	TRP	323	101.045	48.811	30.677	1.00	42.00
2998	NE1	TRP	323	102.292	49.559	28.969	1.00	41.37
2999	CE2	TRP	323	102.367	48.967	30.202	1.00	42.95
3000	CE3	TRP	323	100.841	48.222	31.932	1.00	43.56
3001	CZ2	TRP	323	103.483	48.555	30.939	1.00	41.64
3002	CZ3	TRP	323	101.952	47.812	32.666	1.00	46.00
3003	CH2	TRP	323	103.256	47.982	32.164	1.00	42.99
3004	H	TRP	323	96.604	50.455	31.735	1.00	25.90
3005	HE1	TRP	323	103.052	49.790	28.396	1.00	25.00
3006	N	ASP	324	97.624	52.931	29.503	1.00	59.55
3007	CA C	ASP	324 324	97.500	54.015	28.539	1.00	65.59
3008 3009	Ö	ASP ASP	324	98.480 99.591	55.143 55.176	28.844 28.316	$\frac{1.00}{1.00}$	64.51 68.35
3010	СВ	ASP	324	96.056	54.541	28.552	1.00	70.35
3011	CG	ASP	324	95.713	55.365	27.320	1.00	74.75
3012	OD1	ASP	324	96.439	56.333	27.008	1.00	77.92
3013	OD2	ASP	324	94.698	55.043	26.668	1.00	77.68
3014	H	ASP	324	97.422	53.067	30.447	1.00	25.00
3015	N	ILE	325	98.025	56.067	29.685	1.00	59.63
3016	CA	ILE	325	98.765	57.248	30.131	1.00	62.22
3017	C	ILE	325	97.699	58.281	30.472	1.00	62.36
3018	O	ILE	325	97.807	58.988	31.467	1.00	57.92
3019	CB CC1	ILE	325	99.752	57.824	29.066	1.00	58.28
3020 3021	CG1 CG2	ILE ILE	325 325	100.656 99.004	58.874	29.713 27.882	1.00 $1.00$	56.91 53.60
3021	CD1	ILE	325	101.760	58.424 59.364	28.812	1.00	65.51
3023	Н	ILE	325	97.138	55.950	30.059	1.00	25.00
3024	N	ASN	326	96.622	58.287	29.687	1.00	64.69
3025	CA	ASN	326	95.504	59.203	29.902	1.00	68.20
3026	C	ASN	326	94.857	58.860	31.238	1.00	71.30
3027	Ō	ASN	326	94.171	59.684	31.846	1.00	76.27
3028	CB	ASN	326	94.462	59.058	28.787	1.00	67.58
3029	CG	ASN	326	95.041	59.297	27.406	1.00	69.04
3030	OD1	ASN	326	96.055	59.975	27.225	1.00	68.18
3031	ND2	ASN	326	94.410	58.716	26.395	1.00	67.11
3032	H	ASN	326	96.599	57.685	28.918	1.00	25.00
3033	1HD2	ASN	326	94.783	58.869	25.501	1.00	25.00
3034	2HD2	ASN	326	93.618	58.173	26.568	1.00	25.00

TABLE 11-continued

		In the	e Absence o	f Bound St	ıbstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3035	N	GLU	327	95.095	57.631	31.691	1.00	70.98
3036	CA	GLU	327	94.553	57.144	32.952	1.00	69.92
3037 3038	C O	GLU GLU	327 327	95.259 94.751	57.798 57.777	34.138 35.260	$\frac{1.00}{1.00}$	67.58 68.23
3039	СВ	GLU	327	94.709	55.622	33.034	1.00	69.72
3040	CG	GLU	327	94.147	54.858	31.838	1.00	66.84
3041	CD	GLU	327	92.650	55.040	31.659	1.00	69.61
3042	OE1	GLU	327	91.899	54.845	32.639	1.00	69.86
3043	OE2	GLU	327	92.225	55.369	30.530	1.00	66.87
3044	H	GLU	327 328	95.646	57.037	31.159 33.872	1.00	25.00
3045 3048	N CA	ILE ILE	328 328	96.411 97.212	58.407 59.065	34.901	1.00 1.00	64.07 63.90
3047	C	ILE	328	96.425	60.142	35.657	1.00	70.44
3048	O	ILE	328	96.624	60.338	36.857	1.00	69.65
3049	CB	ILE	328	98.508	59.669	34.286	1.00	56.23
3050	CG1	ILE	328	99.578	59.844	35.359	1.00	56.00
3051	CG2	ILE	328	98.223	61.007	33.618	1.00	53.09
3052 3053	CD1 H	ILE ILE	328 328	100.948 96.744	60.162 58.430	34.799 32.958	$\frac{1.00}{1.00}$	58.67 25.00
3054	N	ASP	329	95.487	60.780	34.961	1.00	75.68
3055	CA	ASP	329	94.659	61.844	35.531	1.00	79.88
3056	С	ASP	329	93.764	61.377	36.677	1.00	79.44
3057	O	ASP	329	93.303	62.188	37.483	1.00	79.71
3058	CB	ASP	329	93.796	62.476	34.435	1.00	85.36
3059	CG	ASP	329	94.608	62.912	33.226	1.00	90.96
3060	OD1	ASP	329 329	95.719	63.461	33.409 32.090	1.00	93.02 93.14
3061 3062	OD2 H	ASP ASP	329 329	94.133 95.344	62.699 60.527	34.026	$\frac{1.00}{1.00}$	25.00
3063	N	ARG	330	93.501	60.074	36.730	1.00	76.85
3064	CA	ARG	330	92.658	59.501	37.775	1.00	75.38
3065	C	ARG	330	93.488	59.059	38.981	1.00	71.57
3066	O	ARG	330	92.935	58.631	39.998	1.00	69.70
3067	CB	ARG	330	91.881	58.300	37.227	1.00	75.33
3068	CG	ARG	330	91.177	58.562	35.905	1.00	78.99
3069	CD	ARG	330 330	90.383 39.861	57.350 57.517	35.454 34.100	1.00 1.00	80.32
3070 3071	NE CZ	ARG ARG	330	88.851	56.816	33.592	1.00	86.31 88.51
3072	NH1	ARG	330	88.239	55.894	34.325	1.00	91.01
3073	NH2	ARG	330	88.458	57.030	32.344	1.00	89.50
3074	H	ARG	330	93.891	59.477	36.060	1.00	25.00
3075	HE	ARG	330	90.281	58.188	33.523	1.00	25.00
3076	1HH1	ARG	330	88.533	55.723	35.265	1.00	25.00
3077	2HH1	ARG	330	87.475 88.917	55.375	33.942	1.00	25.00
3078 3079	1HH2 2HH2	ARG ARG	330 330	87.692	57.720 56.508	31.786 31.966	1.00 1.00	25.00 25.00
3080	N	LEU	331	94.809	59.174	38.867	1.00	67.63
3081	CA	LEU	331	95.723	58.761	39.930	1.00	62.95
3082	С	LEU	331	96.290	59.919	40.735	1.00	60.42
3083	O	LEU	331	96.590	60.974	40.186	1.00	58.28
3084	CB	LEU	331	96.906	57.985	39.338	1.00	58.68
3085 3086	CG CD1	LEU	331	96.664	56.739	38.486 37.941	1.00	54.70
3080	CD1 CD2	LEU LEU	331	97.992	56.255 55.654		1.00	46.64 48.82
3088	H	LEU	331 331		55.654 59.574	39.304 38.061	1.00 $1.00$	48.82 25.00
3089	N	PRO	332	96.426	59.743	42.058	1.00	57.70
3090	CA	PRO	332	96.981	60.814	42.886	1.00	58.87
3091	С	PRO	332	98.455	60.977	42.521	1.00	61.32
3092	O	PRO	332	99.132	59.997	42.207	1.00	63.75
3093	CB	PRO	332	96.800	60.278	44.307	1.00	58.32
3094	CG CD	PRO	332 332	96.819 95.978	58.793	44.122	$\frac{1.00}{1.00}$	60.80 59.09
3095 3096	N N	PRO ASP	333	98.944	58.616 62.210	42.892 42.585	1.00	68.56
3097	CA	ASP	333	100.324	62.554	42.237	1.00	71.37
3098	С	ASP	333	101.432	61.520	42.437	1.00	68.42
3099	O	ASP	333	102.188	61.247	41.504	1.00	65.66
3100	CB	ASP	333	100.715	63.879	42.891	1.00	79.84
3101	CG OD1	ASP	333	99.967	65.057	42.298	1.00	86.67
3102 3103	OD1 OD2	ASP ASP	333 333	100.442 98.897	65.608 65.418	41.283 42.834	1.00 $1.00$	91.38 90.77
3103	H	ASP	333	98.343	62.928	42.866	1.00	25.00
3105	N	TYR	334	101.538	60.936	43.627	1.00	65.59
3106	CA	TYR	334	1022.588	59.953	43.861	1.00	63.25
3107	C	TYR	334	102.455	58.740	42.938	1.00	60.61
3108	O	TYR	334	103.452	58.243	42.411	1.00	63.00

TABLE 11-continued

	In the Absence of Bound Substrate							
Atom Type	Atom	Residue	Residue #	X	Y	z	OCC	B-factor
3109	CB	TYR	334	102.664	59.545	45.341	1.00	65.74
3110	CG	TYR	334	101.539	58.674	45.852	1.00	68.46
3111 3112	CD1 CD2	TYR	334 334	100.343 101.690	59.232 57.289	46.303 45.929	1.00 $1.00$	69.57 68.64
3112	CE1	TYR TYR	334	99.326	58.432	46.824	1.00	69.54
3114	CE2	TYR	334	100.682	56.482	46.446	1.00	69.64
3115	CZ	TYR	334	99.504	57.058	46.892	1.00	70.47
3116	OH	TYR	334	98.515	56.257	47.413	1.00	68.60
3117	H	TYR	334	100.910	61.170	44.337	1.00	25.00
3118	HH	TYR	334	97.786	56.812	47.691	1.00	25.00
3119 3120	N CA	MET MET	335 335	101.220 100.977	58.311 57.174	42.691 41.809	1.00 1.00	51.84 46.91
3121	C	MET	335	101.236	57.558	40.356	1.00	46.00
3122	Ö	MET	335	101.540	56.701	39.525	1.00	49.65
3123	CB	MET	335	99.552	56.646	41.969	1.00	41.83
3124	CG	MET	335	99.268	56.054	43.333	1.00	37.39
3125	SD	MET	335	97.625	55.322	43.450	1.00	44.89
3126	CE	MET	335	97.914	54.042	44.666	1.00	45.77
3127 3128	H N	MET LYS	335 336	100.460 101.122	58.772 58.848	43.094 40.052	1.00 $1.00$	25.00 47.61
3129	CA	LYS	336	101.122	59.340	38.699	1.00	48.27
3130	C	LYS	336	102.836	59.143	38.325	1.00	46.89
3131	O	LYS	336	103.161	58.829	37.177	1.00	49.33
3132	CB	LYS	336	101.000	60.824	38.588	1.00	51.58
3133	CG	LYS	336	99.517	61.132	38.743	1.00	54.94
3134	CD	LYS	336	99.233	62.600	38.446	1.00	60.67
3135	CE	LYS	336	97.739	62.882	38.431	1.00	63.81
3136 3137	NZ H	LYS LYS	336 336	97.404 100.869	64.287 59.485	38.083 40.750	1.00 $1.00$	68.16 25.00
3138	1HZ	LYS	336	97.836	64.931	38.776	1.00	25.00
3139	2HZ	LYS	336	97.770	64.503	37.134	1.00	25.00
3140	3HZ	LYS	338	96.371	64.409	38.092	1.00	25.00
3141	N	ILE	337		59.321	39.303	1.00	44.58
3142	CA	ILE	337	105.164	59.162	39.089	1.00	47.99
3143	С	ILE	337	105.469	57.701	38.782	1.00	50.21 52.97
3144 3145	O CB	ILE ILE	3337 337	106.153 105.957	57.400 59.595	37.800 40.336	1.00 1.00	52.97
3146	CG1	ILE	337	105.533	61.001	40.770	1.00	54.20
3147	CG2	ILE	337	107.455	59.569	40.034	1.00	49.66
3148	CD1	ILE	337	106.048	61.406	42.131	1.00	53.35
3149	H	ILE	337	103.390	59.566	40.195	1.00	25.00
3150	N	SER	338	104.951	56.802	39.618	1.00	46.40
3151	CA C	SER	338 338	105.161	55.364 54.905	39.458 38.098	1.00 1.00	41.92
3152 3153	o	SER SER	338	104.640 105.385	54.347	37.286	1.00	39.82 36.78
3154	СВ	SER	338	104.423	54.598	40.560	1.00	37.35
3155	OG	SER	338	104.502	55.268	41.805	1.00	52.45
3156	H	SER	338	104.411	57.109	40.381	1.00	25.00
3157	HG	SER	338	105.419	55.324	42.084	1.00	25.00
3158	N	TYR	339	103.363	55.183	37.848	1.00	39.53
3159 3160	CA C	TYR TYR	339 339		54.804 55.247	36.608 35.362	$\frac{1.00}{1.00}$	40.68 39.79
3161	Ö	TYR	339	103.719		34.458	1.00	39.78
3162	СВ	TYR	339	101.272		36.586	1.00	39.96
3163	CG	TYR	339	100.388	54.833	35.480	1.00	44.71
3164	CD1	TYR	339	99.948	53.507	35.494	1.00	40.18
3165	CD2	TYR	339	99.992	55.646	34.416	1.00	42.64
3166 3167	CE1	TYR TYR	339 339	99.136 99.180	53.004 55.151	34.475 33.393	$\frac{1.00}{1.00}$	40.02 46.52
3168	CE2 CZ	TYR	339		53.830	33.431	1.00	40.74
3169	OH	TYR	339	97.968	53.3422	32.417	1.00	44.28
3170	H	TYR	339	102.843	55.668	38.522	1.00	25.00
3171	HH	TYR	339	97.819	54.042	31.792	1.00	25.00
3172	N	LYS	340	103.864		35.324	1.00	42.19
3173	CA	LYS	340	134.599	57.032	34.179	1.00	42.37
3174 3175	C O	LYS	340 340	105.930 106.264	56.306 55.885	33.992 32.882	1.00	40.71 41.93
3175 3176	СВ	LYS LYS	340 340	106.264	55.885 58.541	34.306	1.00 $1.00$	41.93 48.48
3177	CG	LYS	340	105.461	59.136	33.063	1.00	61.36
3178	CD	LYS	340	105.412	60.647	33.041	1.00	76.14
3179	CE	LYS	340	105.947	61.164	31.713	1.00	85.02
3180	NZ	LYS	340	105.783	62.636	31.566	1.00	94.84
3181	H	LYS	340	103.661	57.115	36.074	1.00	25.00
3182	1HZ	LYS	340	104.774	62.881	31.624	1.00	25.00

TABLE 11-continued

		In the	Absence o			ene Synth		
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3183	2HZ	LYS	340	106.302	63.119	32.327	1.00	25.00
3184	3HZ	LYS	340	106.160	62.937	30.645	1.00	25.00
3185 3186	N CA	ALA ALA	341 341	106.668 107.963	56.134 55.455	35.084 35.052	$\frac{1.00}{1.00}$	40.84 36.82
3187	C	ALA	341	107.837	54.053	34.469	1.00	35.10
3188	Ö	ALA	341	108.657	53.635	33.650	1.00	34.92
3189	СВ	ALA	341	108.548	55.388	36.451	1.00	37.09
3190	H	ALA	341	106.336	56.478	35.942	1.00	25.00
3191	N	ILE	342	106.796	53.338	34.884	1.00	32.81
3192	CA	ILE	342	106.547	51.983	34.409	1.00	31.78
3193	С	ILE	342	106.357	51.982	32.891	1.00	38.13
3194	O	ILE	342	107.061	51.269	32.163	1.00	37.84
3195 3196	CB CG1	ILE ILE	342 342	105.306 105.585	51.377 51.219	35.109 36.608	1.00 1.00	27.44 30.45
3197	CG2	ILE	342	103.363	50.031	34.499	1.00	29.34
3198	CD1	ILE	342	104.399	50.759	37.420	1.00	30.40
3199	Н	ILE	342	106.179	53.740	35.532	1.00	25.00
3200	N	LEU	343	105.447	52.827	32.414	1.00	44.76
3201	CA	LEU	343	105.168	52.920	30.984	1.00	42.79
3202	С	LEU	343	106.428	53.290	30.214	1.00	40.23
3203	O	LEU	343	106.706	52.724	29.153	1.00	39.11
3204	СВ	LEU	343	104.061	53.943	30.715	1.00	42.96
3205	CG	LEU	343	102.731	53.704	31.436	1.00	46.89
3206 3207	CD1	LEU	343	101.704	54.723	30.978 31.166	1.00	51.34 44.17
3207	CD2 H	LEU LEU	343 343	102.233 104.954	52.302 53.403	33.039	$\frac{1.00}{1.00}$	25.00
3208	N	ASP	343 344	104.934	54.218	30.770	1.00	40.93
3210	CA	ASP	344	108.442	54.660	30.144	1.00	43.89
3211	C	ASP	344	109.443	53.515	30.053	1.00	43.08
3212	O	ASP	344	110.049	53.299	29.001	1.00	38.31
3213	CB	ASP	344	109.056	55.831	30.921	1.00	50.27
3214	CG	ASP	344	108.259	57.124	30.775	1.00	58.66
3215	OD1	ASP	344	107.376	57.206	29.891	1.00	59.02
3216	OD2	ASP	344	108.525	58.070	31.549	1.00	62.70
3217 3218	H N	ASP LEU	344 345	106.928 109.585	54.612 52.764	31.623 31.144	$\frac{1.00}{1.00}$	25.00 40.54
3219	CA	LEU	345	110.511	51.633	31.196	1.00	36.66
3220	C	LEU	345	110.256	50.661	30.048	1.00	36.17
3221	O	LEU	345	111.188	50.256	29.343	1.00	35.58
3222	CB	LEU	345	110.393	50.903	32.540	1.00	38.27
3223	CG	LEU	345	111.284	40.672	32.755	1.00	35.02
3224	CD1	LEU	345	112.750	50.043	32.587	1.00	28.76
3225	CD2	LEU	346	111.030	49.087	34.132	1.00	30.95
3226	H	LEU	345 346	139.050	52.975 50.304	31.934 29.844	1.00 $1.00$	25.00 35.43
3227 3228	N CA	TYR TYR	346 346	108.992 108.650	49.389	28.768	1.00	32.38
3229	C	TYR	346	108.030	49.969	27.388	1.00	34.86
3230	ŏ	TYR	346	109.183	49.228	26.446	1.00	36.74
3231	CB	TYR	346	107.227	48.870	28.927	1.00	33.82
3232	CG	TYR	346	107.173	47.798	29.980	1.00	31.79
3233	CD1	TYR	346	107.531	46.487	29.675	1.00	34.43
3234	CD2	TYR	346	106.856	48.107	31.302	1.00	34.30
3235	CE1	TYR	346	107.585	45.507	30.659	1.00	32.57
3236 3237	CE2 CZ	TYR TYR	346 346	106.906 107.275	47.137 45.839	32.296 31.965	$\frac{1.00}{1.00}$	34.14 34.31
3238	OH	TYR	346	107.351	44.878	32.938	1.00	32.03
3239	Н	TYR	346	108.288	50.665	30.428	1.00	25.00
3240	НН	TYR	346	107.610	44.036	32.562	1.00	25.00
3241	N	LYS	347	108.861	51.295	27.276	1.00	44.24
3242	CA	LYS	347	109.143	51.955	26.004	1.00	44.41
3243	C	LYS	347	110.630	51.792	25.716	1.00	43.81
3244	0	LYS	347	111.030	51.558	24.572	1.00	42.39
3245	CB	LYS	347	108.762	53.437	26.060	1.00	51.50
3246	CG	LYS	347	107.268	53.672	25.945	1.00	55.25
3247 3248	CD CE	LYS LYS	347 347	106.759 105.251	53.062 52.978	24.650 24.608	$\frac{1.00}{1.00}$	59.99 60.17
3248 3249	NZ	LYS	347 347	103.231	52.152	23.446	1.00	53.42
3250	H	LYS	347	108.627	51.840	28.057	1.00	25.00
3251	1HZ	LYS	347	105.241	51.196	23.537	1.00	25.00
3252	2HZ	LYS	347	103.803	52.090	23.409	1.00	25.00
3253	3HZ	LYS	347	105.190	52.590	22.569	1.00	25.00
3254	N	ASP	348	111.439	51.874	26.771	1.00	45.04
3255	CA	ASP	348	112.884	51.712	26.654	1.00	47.19
3256	С	ASP	348	113.178	50.289	26.211	1.00	44.53

TABLE 11-continued

		In the	e Absence of			ene Synth		
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3257	0	ASP	348	113.992	50.074	25.316	1.00	46.94
3258 3259	CB CG	ASP ASP	348 348	113.582	51.981 53.430	27.991	$\frac{1.00}{1.00}$	55.77 63.79
3239 3260	OD1	ASP ASP	348	113.469 113.017	54.288	28.441 27.648	1.00	66.77
3261	OD2	ASP	348	113.846	53.710	29.600	1.00	65.20
3262	Н	ASP	348	111.048	52.057	27.652	1.00	25.00
3263	N	TYR	349	112.507	49.321	26.835	1.00	39.87
3264	CA	TYR	349	112.692	47.913	26.491	1.00	40.93
3265	C	TYR	349	112.412	47.704	25.008	1.00	41.26
3266	O	TYR	349	113.189	47.051	24.302	1.00	40.44
3267 3268	CB CG	TYR TYR	349 349	111.752 112.115	47.015 46.841	27.310 28.773	$\frac{1.00}{1.00}$	35.88 33.98
3269	CD1	TYR	349	113.396	47.144	29.250	1.00	29.99
3270	CD2	TYR	349	111.172	46.360	29.680	1.00	27.01
3271	CE1	TYR	349	113.723	45.971	30.596	1.00	27.43
3272	CE2	TYR	349	111.485	46.182	31.021	1.00	32.24
3273	CZ	TYR	349	112.759	46.491	31.476	1.00	34.71
3274	OH	TYR	349	113.045	46.346	32.813	1.00	31.76
3275	H	TYR	349 349	111.880	49.567	27.549	1.00	25.00
3276 3277	HH N	TYR GLU	349 350	112.295 111.302	45.961 48.269	33.270 24.541	$\frac{1.00}{1.00}$	25.00 44.18
3278	CA	GLU	350	110.911	58.156	23.140	1.00	47.18
3279	C	GLU	350	111.972	48.767	22.235	1.00	45.46
3280	O	GLU	350	112.337	48.175	21.221	1.00	45.14
3281	CB	GLU	350	109.557	48.828	22.903	1.00	48.54
3282	CG	GLU	350	108.396	48.141	23.609	1.00	52.79
3283	CD	GLLU	350	107.076	48.883	23.473	1.00	59.87
3284 3285	OE1 OE2	GLU GLU	350 350	107.070 106.037	50.040 48.304	23.000 23.852	$\frac{1.00}{1.00}$	66.09 65.68
3286	H	GLU	350	110.728	48.772	25.158	1.00	25.00
3287	N	LYS	351	112.497	49.923	22.636	1.00	48.01
3288	CA	LYS	351	113.530	50.618	21.871	1.00	51.81
3289	C	LYS	351	114.794	49.756	21.788	1.00	52.11
3290	O	LYS	351	115.311	49.505	20.696	1.00	48.04
3291	CB	LYS	351	113.861	51.973	22.518	1.00	53.86
3292 3293	CG CD	LYS LYS	351 351	114.151 115.235	53.095 52.708	21.520 20.517	1.00 1.00	60.98 68.26
3293	CE	LYS	351	115.255	53.551	19.253	1.00	75.67
3295	NZ	LYS	351	115.951	52.975	18.132	1.00	74.32
3296	H	LYS	351	112.180	50.322	23.471	1.00	25.00
3297	1HZ	LYS	351	116.950	52.914	18.410	1.00	25.00
3298	2HZ	LYS	351	115.590	52.025	17.907	1.00	25.00
3299	3HZ	LYS	351	115.855	53.584 49.297	17.293 22.944	1.00	25.00
3300 3301	N CA	GLU GLU	352 352	115.275 116.474	48.461	23.031	$\frac{1.00}{1.00}$	56.12 54.04
3302	C	GLU	352	116.409	47.241	22.120	1.00	52.28
3303	O	GLU	352	117.410	46.851	21.514	1.00	52.78
3304	CB	GLU	352	116.688	47.971	24.466	1.00	58.92
3305	CG	GLU	352	117.135	49.023	25.460	1.00	67.31
3306	CD OF1	GLU GLU	352 352	117.386	48.438	26.842	1.00	71.98
3307 3308	OE1 OE2	GLU	352 352	118.383 116.582	47.694 48.718	27.004 27.760	$\frac{1.00}{1.00}$	69.15 65.74
3309	H	GLU	352	114.800	49.527	23.762	1.00	25.00
3310	N	LEU	353	115.235	46.624	22.052	1.00	49.88
3311	CA	LEU	353	115.053	45.435	21.233	1.00	51.47
3312	C	LEU	353	114.701	45.732	19.772	1.00	55.82
3313	O	LEU	353	114.606	44.809	18.955	1.00	56.53
3314 3315	CB CG	LEU LEU	353 353	114.009	44.511	21.876 23.297	1.00	44.77 40.31
3316	CD1	LEU	353 353	114.320 113.151	44.017 43.224	23.855	$\frac{1.00}{1.00}$	35.62
3317	CD2	LEU	353	115.586	43.172	23.302	1.00	33.71
3318	H	LEU	353	114.478	46.975	22.571	1.00	25.00
3319	N	SER	354	114.538	47.012	19.437	1.00	62.13
3320	CA	SER	354	114.202	47.423	18.071	1.00	66.31
33221	С	SER	354	115.245	46.970	17.058	1.00	64.64
3322	O CB	SER	354 354	114.904 114.043	46.378	16.035	1.00	66.26
3323 3324	CB OG	SER SER	354 354	114.043	48.945 49.406	17.978 18.763	$\frac{1.00}{1.00}$	69.09 80.83
3324	Н	SER	354 354	114.623	47.710	20.121	1.00	25.00
3326	HG	SER	354	112.888	50.360	18.684	1.00	25.00
3327	N	SER	355	116.516	47.223	17.359	1.00	65.77
3328	CA	SER	355	117.616	46.850	16.472	1.00	67.77
3329	С	SER	355 355	117.631	45.364	16.110	1.00	68.81
3330	О	SER	355	118.082	44.990	15.028	1.00	69.36

TABLE 11-continued

In the Absence of Bound Substrate								
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3331	CB	SER	355	118.956	47.245	17.099	1.00	66.08
3332	OG	SER	355	119.067	46.741	18.419	1.00	68.44
3333	H	SER	355	116.729	47.670	18.209	1.00	25.00
3334 3335	HG N	SER ALA	355 356	119.043 117.150	45.780 44.525	18.416 17.024	$\frac{1.00}{1.00}$	25.00 69.39
3336	CA	ALA	356	117.115	43.082	16.802	1.00	68.66
3337	C	ALA	356	115.741	42.585	16.347	1.00	69.25
3338	O	ALA	356	115.561	41.395	16.084	1.00	71.52
3339	CB	ALA	356	117.549	42.347	18.067	1.00	64.46
3340	H	ALA	356	116.806	44.884	17.867	1.00	25.00
3341 3342	N CA	GLY	357	114.773	43.493	16.270	1.00	67.77
3342 3343	CA	GLY GLY	357 357	113.432 112.754	43.118 42.202	15.854 16.856	$\frac{1.00}{1.00}$	62.16 58.02
3344	Ö	GLY	357	111.969	41.327	16.481	1.00	58.07
3345	H	GLY	357	114.965	44.428	16.477	1.00	25.00
3346	N	ARG	358	113.039	42.416	18.138	1.00	53.28
3347	CA	ARG	358	112.461	41.601	19.204	1.00	50.96
3348	C	ARG	358	111.488	42.359	20.106	1.00	50.55
3349	O CB	ARG	358 358	110.885	41.774	20.999	1.00	51.85
3350 3351	СG	ARG ARG	358	113.568 114.360	40.953 39.872	20.047 19.314	$\frac{1.00}{1.00}$	44.69 43.66
3352	CD	ARG	358	115.389	39.206	20.217	1.00	43.11
3353	NE	ARG	358	114.768	38.503	21.338	1.00	42.40
3354	CZ	ARG	358	114.997	38.783	22.618	1.00	43.84
3355	NH1	ARG	358	115.836	39.754	22.951	1.00	49.95
3356	NH2	ARG	358	114.389	38.089	23.571	1.00	45.03
3357	H	ARG	358	113.664	43.136	18.368	1.00	25.00
3358 3359	HE 1HH1	ARG ARG	358 358	114.142 116.301	37.776 40.282	21.138 22.241	$\frac{1.00}{1.00}$	25.00 25.00
3360	2HH1	ARG	358	116.006	39.958	23.915	1.00	25.00
3361	1HH2	ARG	358	113.755	37.355	23.327	1.00	25.00
3362	2HH2	ARG	358	114.562	38.301	24.533	1.00	25.00
3363	N	SER	359	111.270	43.639	19.826	1.00	50.59
3364	CA	SER	359	110.363	44.464	20.625	1.00	47.98
3365	С	SER SER	359 359	108.948	43.888	20.767	1.00	48.46
3368 3367	O CB	SER	359 359	108.247 110.315	44.177 45.879	21.737 20.050	$\frac{1.00}{1.00}$	46.16 51.38
3368	OG	SER	359	110.450	45.839	18.639	1.00	63.31
3369	H	SER	359	111.730	44.045	19.067	1.00	25.00
3370	HG	SER	359	111.323	45.514	18.419	1.00	25.00
3371	N	HIS	360	108.559	43.029	19.8829	1.00	46.52
3372	CA	HIS	360	107.234	42.401	19.837	1.00	47.05
3373 3374	C O	HIS HIS	360 360	106.998 105.893	41.398 40.871	20.974 21.124	$\frac{1.00}{1.00}$	48.80 46.79
3375	СВ	HIS	360	105.893	41.713	18.492	1.00	47.13
3376	CG	HIS	360	108.026	40.724	18.100	1.00	47.23
3377	ND1	HIS	360	107.885	39.365	18.289	1.00	50.30
3378	CD2	HIS	360	109.242	40.899	17.532	1.00	49.82
3379	CE1	HIS	380	103.969	38.746	17.855	1.00	47.18
3380 3381	NE2 H	HIS HIS	360 360	109.808 109.183	39.655 42.837	17.391 19.111	$\frac{1.00}{1.00}$	46.40 25.00
3382	HD1	HIS	360	107.098	38.919	18.675	1.00	25.00
3383	HE2	HIS	360	110.702	39.494	16.992	1.00	25.00
3384	N	ILE	361	108.042	41.098	21.741	1.00	47.17
3385	CA	ILE	361	107.922	40.152	22.845	1.00	40.37
3386	C	ILE	361	107.657	40.850	24.175	1.00	35.93
3387	O	ILE	361	107.118 109.187	40.240	25.103	1.00	41.86
3388 3389	CB CG1	ILE ILE	361 361	110.392	39.277 40.152	22.987 23.346	$\frac{1.00}{1.00}$	44.49 39.20
3390	CG2	ILE	361	109.421	38.477	21.707	1.00	38.28
3391	CD1	ILE	361	111.680	39.405	23.464	1.00	49.02
3392	H	ILE	361	108.913	41.517	21.574	1.00	25.00
3393	N	VAL	362	108.007	42.131	24.256	1.00	29.83
3394	CA	VAL	362	107.818	42.911	25.478	1.00	28.00
3395 3396	C O	VAL VAL	362 362	106.396 106.209	42.815 42.788	26.041 27.262	$\frac{1.00}{1.00}$	32.52 34.43
3397	СВ	VAL	362	108.203	44.401	25.259	1.00	30.93
3398	CG1	VAL	362	107.851	45.233	26.484	1.00	28.70
3399	CG2	VAL	362	109.699	44.524	24.952	1.00	22.99
3400	H	VAL	362	108.395	42.568	23.472	1.00	25.00
3401	N	CYS	363	105.405	42.701	25.160	1.00	29.77
3402	CA	CYS	363	104.011	42.610	25.592	1.00	31.29
3403 3404	C O	CYS CYS	363 363	103.757 102.942	41.470 41.610	26.581 27.499	$\frac{1.00}{1.00}$	29.63 25.93
2101	-		2 32	30 <u>20</u> , 1 <u>0</u>		2	2.00	

TABLE 11-continued

Structural	Coordinates of Tobacco 5-Epi-Aristolochene Synthase						
In the Absence of Bound Substrate							

In the Absence of Bound Substrate								
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3405	CB	CYS	363	103.066	42.504	24.387	1.00	33.73
3406	SG	CYS	363	103.387	41.126	23.270	1.00	40.61
3407	Н	CYS	363	105.612	42.677	24.204	1.00	25.00
3408	N	HIS	364	104.480	40.362	26.421	1.00	26.78
3409	CA	HIS	364	104.332	39.216	27.315	1.00	24.36
3410	С	HIS	364	104.685	39.599	28.754	1.00	31.26
3411	O	HIS	364	104.029	39.163	29.703	1.00	32.48
3412 3413	CB CG	HIS HIS	364 364	105.220 104.826	38.064 37.486	26.855 25.531	$\frac{1.00}{1.00}$	26.25 29.86
3414	ND1	HIS	364	103.731	36.663	25.372	1.00	39.99
3415	CD2	HIS	364	105.398	37.588	24.308	1.00	32.37
3416	CE1	HIS	364	103.646	36.282	24.110	1.00	36.28
3417	NE2	HIS	364	104.648	36.829	23.444	1.00	32.87
3418	Н	HIS	364	105.139	40.332	25.695	1.00	25.00
3419	HD1	HIS	364	103.113	38.392	26.086	1.00	25.00
3420	HE2	HIS	364	104.819	36.718	22.488	1.00	25.00
3421	N	ALA	365	105.711	40.431	28.911	1.00	28.85
3422	CA	ALA	365	106.134	40.880	30.232	1.00	30.11
3423	C	ALA	365	105.075	41.813	30.826	1.00	31.00
3424	O	ALA	365	104.727	41.708	32.005	1.00	31.25
3425 3426	CB H	ALA ALA	365 365	107.482 106.191	41.585 40.762	30.140 28.123	$\frac{1.00}{1.00}$	31.82 25.00
3427	N	ILE	366	100.191	42.708	29.992	1.00	33.54
3428	CA	ILE	366	103.512	43.653	30.417	1.00	33.21
3429	C	ILE	366	102.287	42.889	30.922	1.00	29.56
3430	O	ILE	366	101.743	43.199	31.987	1.00	31.04
3431	CB	ILE	366	103.036	44.582	29.255	1.00	33.17
3432	CG1	ILE	386	104.264	45.455	28.823	1.00	30.87
3433	CG2	ILE	366	101.908	45.453	29.674	1.00	28.96
3434	CD1	ILE	366	103.987	46.289	27.599	1.00	35.02
3435	H	ILE	366	104.869	42.731	29.067	1.00	25.00
3436	N	GLU	367	101.874	41.875	30.167	1.00	28.47
3437	CA	GLU	367	100.726	41.081	30.548	1.00	30.13
3438	С	GLU	367	100.945	40.439	31.929	1.00	30.58
3439 3440	O CB	GLU GLU	367 367	100.029	40.407 39.966	32.754 29.507	$\frac{1.00}{1.00}$	31.55 38.78
3441	CG	GLU	367	100.461 100.228	40.472	28.074	1.00	52.31
3442	CD	GLU	367	99.180	41.585	27.970	1.00	62.83
3443	OE1	GLU	367	98.144	41.525	28.675	1.00	55.58
3444	OE2	GLU	367	99.395	42.523	27.168	1.00	63.90
3445	Н	GLU	367	102.351	41.673	29.337	1.00	25.00
3446	N	ARG	368	102.167	39.985	32.196	1.00	28.25
3447	CA	ARG	368	102.479	39.385	33.487	1.00	21.98
3448	С	ARG	368	102.462	40.420	34.607	1.00	22.36
3449	O	ARG	368	102.080	40.108	35.738	1.00	22.36
3450	CB	ARG	368	103.821	38.661	33.440	1.00	23.96
3451	CG	ARG ARG	368	103.796 102.812	37.364	32.642	1.00	17.80
3452 3453	CD NE	ARG	368 368	102.812	36.352 35.034	33.224 32.626	$\frac{1.00}{1.00}$	19.62 19.60
3454	CZ	ARG	368	102.516	33.897	33.113	1.00	20.41
3455	NH1	ARG	368	101.773	33.898	34.211	1.00	26.21
3458	NH2	ARG	368	102.843	32.743	32.548	1.00	22.02
3457	H	ARG	368	102.868	40.040	31.510	1.00	25.00
3458	HE	ARG	368	103.526	34.993	31.804	1.00	25.00
3459	1HH1	ARG	368	101.580	34.754	34.685	1.00	25.00
3460	2HH1	ARG	368	101.410	33.036	34.566	1.00	25.00
3461	1HH2	ARG	368	103.454	32.730	31.755	1.00	25.00
3462	2HH2	ARG	368	102.476	31.888	32.904	1.00	25.00
3463	N	MET	369	102.849	41.654	34.293	1.00	23.19
3464	CA	MET	369	102.845	42.716	35.295	1.00	20.55
3465 3466	C O	MET MET	369 369	101.410 101.085	43.060 43.248	35.657 36.833	$\frac{1.00}{1.00}$	20.66 24.28
3467	СВ	MET	369	103.565	43.966	34.789	1.00	24.43
3468	CG	MET	369	103.575	45.097	35.806	1.00	27.10
3469	SD	MET	369	104.503	46.538	35.283	1.00	33.91
3470	CE	MET	369	105.378	46.942	36.804	1.00	36.11
3471	H	MET	369	103.151	41.852	33.380	1.00	25.00
3472	N	LYS	370	100.550	43.142	34.645	1.00	27.52
3473	CA	LYS	370	99.135	43.441	34.865	1.00	27.03
3474	C	LYS	370	98.572	42.392	35.817	1.00	26.80
3475	O	LYS	370	97.854	42.720	36.766	1.00	31.01
3476	CB	LYS	370	98.361	43.415	33.545	1.00	28.62
3477	CG	LYS	370	98.699	44.546	32.591	1.00	26.77
3478	CD	LYS	370	97.881	44.437	31.318	1.00	32.86

TABLE 11-continued

		In the	Absence of					
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3479	CE	LYS	370	98.174	45.591	30.371	1.00	40.98
3480	NZ	LYS	370	97.397	45.502	29.099	1.00	45.53
3481	H	LYS	370	100.870	43.001	33.729	1.00	25.00
3482	1HZ	LYS	370	96.379 97.630	45.511	29.313	1.00	25.00
3483 3484	2HZ	LYS	370 370	97.630	46.314 44.619	28.492 28.607	1.00	25.00
	3HZ	LYS GLU	3771	98.959		35.581	1.00 1.00	25.00
3485	N				41.139			26.63
3485 3485	N	GLU GLU	3771	98.959 98.959	41.139	35.581 35.581	1.00	26.63
3486 3486	N CA	GLU	3771 371	98.541	41.139 40.006	36.398	1.00 1.00	26.63 23.18
3487	C	GLU	371	98.981	40.191	37.854	1.00	31.56
3488	o	GLU	371	98.180	40.023	38.782	1.00	33.30
3489	СВ	GLU	371	99.125	38.719	35.815	1.00	24.88
3490	CG	GLU	371	98.779	37.449	36.569	1.00	25.01
3491	CD	GLU	371	99.346	36.212	35.892	1.00	38.13
3492	OE1	GLU	371	100.588	36.080	35.815	1.00	34.30
3493	OE2	GLU	371	98.549	35.373	35.425	1.00	44.97
3494	H	GLU	371	99.542	40.972	34.811	1.00	25.00
3495	N	VAL	372	100.243	40.567	38.056	1.00	29.02
3496	CA	VAL	372	100.765	40.789	39.406	1.00	26.07
3497	C	VAL	372	99.952	41.869	40.126	1.00	28.22
3498	ŏ	VAL	372	99.582	41.705	41.293	1.00	27.36
3499	СВ	VAL	372	102.261	41.216	39.388	1.00	27.23
3500	CG1	VAL	372	102.738	41.520	40.801	1.00	19.82
3501	CG2	VAL	372	103.124	40.119	38.770	1.00	23.40
3502	H	VAL	372	100.836	40.688	37.283	1.00	25.00
3503	N	VAL	373	99.657	42.964	39.426	1.00	31.06
3504	CA	VAL	373	98.893	44.063	40.018	1.00	32.80
3505	C	VAL	373	97.453		40.378	1.00	33.82
3506	0	VAL	373	96.952	44.054	41.441	1.00	30.74
3507	СВ	VAL	373	98.908	45.314	39.118	1.00	34.89
3508	CG1	VAL	373	98.134		39.775	1.00	32.71
3509	CG2	VAL	373	100.345	45.741	38.855	1.00	34.17
3510	H	VAL	373	99.960	43.0288	38.495	1.00	25.00
3511	N	ARG	374	96.794	42.900	39.512	1.00	31.81
3512	CA	ARG	374	95.428	42.447	39.789	1.00	29.59
3513	C	ARG	374	95.422	41.667	41.093	1.00	31.35
3514	O	ARG	374	94.613	41.933	41.989	1.00	36.45
3515	CB	ARG	374	94.910	41.519	38.689	1.00	29.25
3516	CG	ARG	374	94.668	42.166	37.349	1.00	32.69
3517	CD	ARG	374	94.034	41.169	36.396	1.00	33.05
3518	NE	ARG	374	94.840	40.973	35.194	1.00	37.26
3519	CZ	ARG	374	95.399	39.817	34.846	1.00	38.15
3520	NH1	ARG	374	95.247	38.741	35.608	1.00	41.80
3521	NH2	ARG	374	96.114	39.738	33.733	1.00	40.61
3522	H	ARG	374	97.230	42.625	38.677	1.00	25.00
3523	HE	ARG	374	94.974	41.744	34.603	1.00	25.00
3524	1HH1	ARG	374	94.711	38.795	36.448	1.00	25.00
3525	2HH1	ARG	374	95.672	37.878	35.336	1.00	25.00
3526	1HH2	ARG	374	96.232	40.550	33.158	1.00	25.00
3527	2HH2	ARG	374	96.537	38.873	33.467	1.00	25.00
3528	N	ASN	375	96.351	40.721	41.202	1.00	31.47
3529	CA	ASN	375	96.458	39.883	42.388	1.00	28.79
3530	C	ASN	375	96.897	40.652	43.625	1.00	26.40
3531	O	ASN	375	96.561	40.266	44.746	1.00	27.83
3532	CB	ASN	375	97.359	33.683	42.112	1.00	35.49
3533	CG	ASN	375	96.744	37.720	41.111	1.00	32.04
3534	OD1	ASN	375	95.982	38.125	40.237	1.00	33.82
3535	ND2	ASN	375	97.075	36.442	41.231	1.00	34.73
3536	Н	ASN	375	96.970	40.566	40.454	1.00	25.00
3537	1HD2	ASN	375	96.671	35.322	40.590	1.00	25.00
3538	2HD2	ASN	375	97.686	36.184	41.941	1.00	25.00
3539	N	TYR	376	97.643	41.736	43.422	1.00	32.41
3540	CA	TYR	376	98.075	42.599	44.526	1.00	36.00
3541	С	TYR	376	96.803	43.220	45.101	1.00	36.51
3542	O	TYR	376	96.585	43.247	46.316	1.00	32.23
3543	CB	TYR	370	98.960	43.739	44.010	1.00	34.19
3544	CG	TYR	376	100.447	43.464	43.979	1.00	41.46
3545	CD1	TYR	376	100.993	42.339	44.601	1.00	40.12
3546	CD2	TYR	376	101.315	44.350	43.336	1.00	41.43
3547	CE1	TYR	376	102.365	42.104	44.580	1.00	38.75
3548	CE2	TYR	376	102.688	44.127	43.310	1.00	37.68
3549 3550	CZ	TYR	376 276	103.203	43.005	43.932	1.00	41.58
3550	OH	TYR	376	104.560	42.785	43.895	1.00	43.07

TABLE 11-continued

		In th	e Absence of	t Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3551	H	TYR	376	97.915	41.956	42.506	1.00	25.00
3552	HH	TYR	376	104.761	41.958	44.341	1.00	25.00
3553	N	ASN	377	95.965	43.713	44.194	1.00	37.58
3554 3555	CA C	ASN ASN	377 377	94.704 93.807	44.343 43.352	44.550 45.285	1.00 1.00	36.82 35.16
3556	o	ASN	377 377	93.276	43.658	46.353	1.00	35.66
3557	СВ	ASN	377	94.011	44.846	43.287	1.00	38.60
3558	CG	ASN	377	92.858	45.770	43.587	1.00	43.38
3559	OD1	ASN	377	92.949	46.628	44.462	1.00	38.97
3560	ND2	ASN	377	91.774	45.622	42.838	1.00	46.57
3561	H	ASN	377	96.210	43.651	43.245	1.00	25.00
3582	1HD2	ASN	377	91.023	46.218	43.029	1.00	25.00
3563	2HD2	ASN	377	91.765	44.936	42.143	1.00	25.00
3564	N	VAL	378	93.683	42.147	44.735	1.00	32.14
3565	CA	VAL	378	92.857	41.106	45.344	1.00	28.96
3566	С	VAL	378	93.339	40.801	46.766	1.00	33.76
3567	O	VAL	378	92.532	40.647	47.690	1.00	32.35
3568	CB CG1	VAL	378	92.858 92.051	39.818	44.490 45.169	1.00	30.20 28.32
3569 3570	CG2	VAL VAL	378 378	92.031	38.732 40.104	43.105	$\frac{1.00}{1.00}$	26.65
3571	H	VAL VAL	378	94.153	41.951	43.103	1.00	25.00
3572	N N	GLU	379	94.133	40.741	46.940	1.00	36.35
3573	CA	GLU	379	95.258	40.478	48.246	1.00	38.43
3574	C	GLU	379	94.875	41.598	49.209	1.00	36.80
3575	ŏ	GLU	379	94.579	41.352	50.383	1.00	37.49
3576	СВ	GLU	379	96.780	40.395	48.114	1.00	43.01
3577	CG	GLU	379	97.544	40.416	49.436	1.00	52.96
3578	CD	GLU	379	99.055	40.403	49.250	1.00	61.44
3579	OE1	GLU	379	99.526	40.225	48.107	1.00	70.64
3580	OE2	GLU	379	99.776	40.568	50.255	1.00	66.21
3581	H	GLU	379	95.247	40.882	46.167	1.00	25.00
3582	N	SER	380	94.894	42.827	48.700	1.00	37.49
3583	CA	SER	380	94.531	44.003	49.480	1.00	38.42
3584	С	SER	380	93.070	43.865	49.906	1.00	37.35
3585	O	SER	380	92.740	44.018	51.085	1.00	38.35
3586	CB	SER	380	94.721 94.344	45.264	48.634	1.00	37.87
3587 3588	OG H	SER SER	380 330	94.344	46.428 42.951	49.349 47.767	1.00 1.00	5i.23 25.00
3589	HG	SER	380	94.903	46.518	50.127	1.00	25.00
3590	N	THR	381	92.209	43.535	48.945	1.00	36.73
3591	CA	THR	381	90.785	43.349	49.198	1.00	31.81
3592	C	THR	381	90.574	42.288	50.278	1.00	33.52
3593	O	THR	381	89.846	42.514	51.245	1.00	35.95
3594	CB	THR	381	90.043	42.922	47.912	1.00	27.90
3595	OG1	THR	381	90.230	43.914	46.894	1.00	30.65
3596	CG2	THR	381	88.564	42.762	48.174	1.00	30.75
3597	H	THR	381	92.531	43.424	48.030	1.00	25.00
3598	HG1	THR	381	89.901	44.765	47.202	1.00	25.00
3599	N	TRP	382	91.246	41.146	50.137	1.00	31.98
3600	CA	TRP	382	91.124	40.059	51.106	1.00	34.10
3601 3602	C O	TRP TRP	382 382	91.498 90.840	40.511 40.145	52.513	$\frac{1.00}{1.00}$	37.61 37.71
3602 3603	CB	TRP	382 382	90.840	40.145 38.870	53.490 50.701	1.00	29.03
3604	CG CG	TRP	382 382	92.001	38.064	49.553	1.00	34.52
3605	CD1	TRP	382	90.298	38.272	48.872	1.00	32.13
3606	CD2	TRP	382	92.073	36.907	48.962	1.00	41.36
3607	NE1	TRP	382	90.141	37.315	47.897	1.00	33.27
3608	CE2	TRP	382	91.215	36.465	47.929	1.00	39.81
3609	CE3	TRP	382	93.262	36.198	49.205	1.00	42.10
3610	CZ2	TRP	382	91.507	35.344	47.138	1.00	41.24
3611	CZ3	TRP	382	93.552	35.082	48.417	1.00	37.35
3612	CH2	TRP	382	92.676	34.669	47.396	1.00	37.45
3613	H	TRP	382	91.841	41.035	49.370	1.00	25.00
3614	HE1	TRP	382	89.384	37.250	47.275	1.00	25.00
3615	N	PHE	383	92.551	41.317	52.601	1.00	42.79
3616	CA	PHE	383	93.040	41.836	53.875	1.00	44.89
3617	С	PHE	383	92.005	42.728	54.561	1.00	45.32
3618	O	PHE	383	91.714	42.557	55.748	1.00	44.05
3619	CB	PHE	383	94.346	42.611	53.657	1.00	45.88 46.70
3620 3621	CG CD1	PHE PHE	383 383	94.818 95.254	43.358 42.674	54.869 55.997	$\frac{1.00}{1.00}$	46.79 47.27
3622	CD1 CD2	PHE	383	93.234	44.751	54.893	1.00	50.35
3623	CE1	PHE	383	95.665	43.368	57.137	1.00	53.01
3624	CE2	PHHE	383	95.208	45.453	56.026	1.00	50.27
JU2T	UL2	LILL	505	22.200	10.700	50.020	1.00	50.27

TABLE 11-continued

		In the	Absence of	f Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3625	CZ	PHE	383	95.641	44.759	57.151	1.00	48.55
3626	H	PHE	383	93.023	41.569	51.778	1.00	25.00
3627	N	ILE	384	91.462	43.677	53.803	1.00	46.83
3628	CA	ILE	384	90.458	44.610	54.306	1.00	46.17
3629	C	ILE	384	89.185	43.894	54.774	1.00	47.27
3630	O	ILE	384	88.608	44.253	55.799	1.00	47.11
3631	CB	ILE	384	90.091	45.646	53.227	1.00	41.64
3632 3633	CG1 CG2	ILE ILE	384 384	91.337 89.031	46.434 46.597	52.817 53.750	$\frac{1.00}{1.00}$	42.54 46.00
3634	CD1	ILE	384	91.148	47.270	51.568	1.00	40.69
3635	Н	ILE	384	91.753	43.749	52.867	1.00	25.00
3636	N	GLU	385	88.756	42.884	54.022	1.00	45.25
3637	CA	GLU	385	87.554	42.123	54.360	1.00	43.73
3638	C	GLU	385	87.791	41.137	55.495	1.00	46.22
3639	O	GLU	385	36.842	40.636	56.097	1.00	51.43
3640	CB	GLU	385	87.051	41.346	53.142	1.00	42.68
3641	CG	GLU	385	86.657	42.211	51.956	1.00	46.85
3642	CD	GLU	385	86.265	41.397	50.730	1.00	50.00
3643	OE1	GLU	385	86.535	40.175	50.696	1.00	44.05
3644	OE2	GLU	385	85.689	41.988	49.791	1.00	52.88
3645	H	GLU	385	89.262	42.652	53.215	1.00	25.00
3648	N	GLY	386	89.055	40.846	55.777	1.00	45.48
3647	CA	GLY	386	89.371	39.893	56.824	1.00	41.52
3648	С	GLY	386	89.038	38.499	56.328	1.00	42.60
3649 3650	O H	GLY GLY	386 386	88.656 89.784	37.625 41.273	57.104 55.279	$\frac{1.00}{1.00}$	44.94 25.00
3651	N	TYR	387	89.190	38.297	55.023	1.00	42.34
3652	CA	TYR	387	88.897	37.020	54.382	1.00	43.55
3653	C	TYR	387	90.042	36.010	54.474	1.00	45.46
3654	ŏ	TYR	387	91.191	36.329	54.162	1.00	49.16
3655	CB	TYR	387	88.545	37.254	52.908	1.00	38.26
3656	CG	TYR	387	88.082	36.017	52.162	1.00	36.21
3657	CD1	TYR	387	87.152	35.142	52.727	1.00	36.57
3658	CD2	TYR	387	88.555	35.736	50.880	1.00	31.64
3659	CE1	TYR	387	86.704	34.015	52.035	1.00	32.07
3660	CE2	TYR	387	88.112	34.616	50.178	1.00	32.41
3661	CZ	TYR	387	87.187	33.759	50.763	1.00	34.67
36662	OH	TYR	387	86.749	32.648	50.082	1.00	38.16
3663	Н	TYR	387	89.541	39.025	54.474	1.00	25.00
3664 3665	HH N	TYR THR	387 388	87.147 89.706	32.629 34.787	49.211 54.872	$\frac{1.00}{1.00}$	25.00 45.36
3666	CA	THR	388	90.671	33.692	54.986	1.00	43.34
3667	C	THR	388	90.199	32.571	54.048	1.00	41.02
3668	ŏ	THR	388	89.474	31.660	54.459	1.00	45.75
3669	СВ	THR	388	90.748	33.161	56.444	1.00	42.74
3670	OG1	THR	388	91.169	34.220	57.314	1.00	43.28
3671	CG2	THR	388	91.741	32.008	56.561	1.00	39.83
3672	H	THR	388	88.782	34.613	55.129	1.00	25.00
3673	HG1	THR	388	91.243	33.885	58.206	1.00	25.00
3674	N	PRO	389	90.575	32.649	52.761	1.00	34.93
3675	CA	PRO	389	90.184	31.645	51.769	1.00	34.82
3676	С	PRO	389	90.846	30.293	51.974	1.00	39.00
3677	O	PRO	389	91.864	30.185	52.658	1.00	44.20
3678 3679	CB	PRO PRO	389 389	90.654 91.878	32.273	50.460	$\frac{1.00}{1.00}$	29.22 31.21
3680	CG CD	PRO	389 389	91.676	33.003 33.671	50.869 52.151	1.00	33.29
3681	N	PRO	390	90.243	29.231	51.425	1.00	39.35
3682	CA	PRO	390	90.830	27.896	51.566	1.00	38.80
3683	C	PRO	390	92.130	27.894	50.744	1.00	42.47
3684	O	PRO	390	92.264	28.683	49.801	1.00	41.59
3685	CB	PRO	390	89.756	26.991	50.960	1.00	35.31
3686	CG	PRO	390	89.094	27.876	49.944	1.00	39.05
3687	CD	PRO	390	88.968	29.177	50.690	1.00	35.38
3688	N	VAL	391	93.070	27.015	51.085	1.00	42.92
3689	CA	VAL	391	94.367	26.947	50.396	1.00	40.91
3690	C	VAL	391	94.310	27.035	48.869	1.00	40.48
3691	O	VAL	391	95.026	27.832	48.266	1.00	37.26
3692	CB	VAL	391	95.163	25.685	50.800	1.00	42.01
3693	CG1	VAL	391	96.542	25.698	50.149	1.00	37.12
3694 3695	CG2	VAL	391	95.298 92.886	25.616	52.307	1.00	37.77
3695 3696	H N	VAL SER	391 392	92.886	26.404 26.228	51.823 48.251	$\frac{1.00}{1.00}$	25.00 37.92
3697	CA	SER	392 392	93.433	26.223	46.799	1.00	36.67
3698	CA	SER	392	93.065	27.627	46.253	1.00	37.70
2070	-						2.50	0

TABLE 11-continued

Structural Coordinates of Tobacco 5-Epi-Ar	ristolochene Synthase	
In the Absence of Bound Su	bstrate	

		In the	e Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3699	O	SER	392	93.699	28.056	45.289	1.00	39.57
3700	CB	SER	392	92.167	25.301	46.399	1.00	41.29
3701	OG	SER	392	91.008	25.599	47.163	1.00	53.55
3702	H	SER	392	92.894	25.625	48.776	1.00	25.00
3703	HG	SER	392	90.720	26.495	46.965	1.00	25.00
3704	N	GLU	393	92.140	28.342	46.883	1.00	35.95
3705	CA	GLU	393	91.806	29.692	46.455	1.00	34.88
3706	C	GLU	393	92.951	30.643	46.783	1.00	31.14
3707	O	GLU	393	93.293	31.516	45.984	1.00	29.96
3708	CB	GLU	393	90.518	30.159	47.130	1.00	35.59
3709	CG	GLU	393	89.956	31.447	46.559	1.00	35.57
3710	CD OE1	GLU	393 393	88.746	31.951	47.318	1.00	39.64 40.88
3711 3712	OE2	GLU GLU	393 393	88.064 88.475	31.141 33.167	47.985 47.2242	$\frac{1.00}{1.00}$	40.88 40.96
3713	H H	GLU	393 393	91.694	27.967	47.2242	1.00	25.00
3713	N N	TYR	393 394	93.539	30.476	47.962	1.00	32.04
3715	CA	TYR	394	94.655	31.318	48.371	1.00	29.74
3716	C	TYR	394	95.743	31.287	47.302	1.00	31.70
3717	Ö	TYR	394	96.180	32.335	46.822	1.00	33.06
3718	СВ	TYR	394	95.238	30.844	49.706	1.00	33.01
3719	CG	TYR	394	96.548	31.520	50.059	1.00	42.08
3720	CD1	TYR	394	96.585	32.876	50.392	1.00	45.51
3721	CD2	TYR	394	97.752	30.815	50.021	1.00	35.80
3722	CE1	TYR	394	97.791	33.517	50.675	1.00	45.10
3723	CE2	TYR	394	98.963	31.448	50.299	1.00	36.76
3724	CZ	TYR	394	98.975	32.798	50.627	1.00	42.58
3725	OH	TYR	394	100.164	33.430	50.915	1.00	41.43
3726	H	TYR	394	93.225	29.778	48.567	1.00	25.00
3727	HH	TYR	394	99.991	34.360	51.082	1.00	25.00
3728	N	LEU	395	96.145	30.082	46.909	1.00	28.83
3729	CA	LEU	395	97.189	29.897	45.910	1.00	26.16
3730	С	LEU	395	96.865	30.472	44.541	1.00	29.12
3731	O	LEU	395	97.737	31.063	43.901	1.00	28.83
3732	CB	LEU	395	97.550	28.415	45.770	1.00	28.06
3733 3734	CG CD1	LEU LEU	395 395	98.263 98.511	27.754 26.290	46.951	1.00 1.00	27.93 28.97
3735	CD1	LEU	395 395	98.511	28.475	46.638 47.245	1.00	24.02
3736	H	LEU	395 395	95.720	29.293	47.243	1.00	25.00
3737	N	SER	396	95.620	30.324	44.093	1.00	29.97
3738	CA	SER	396	95.239	30.836	42.780	1.00	32.88
3739	C	SER	396	95.535	32.329	42.624	1.00	28.77
3740	Ö	SER	396	95.715	32.818	41.508	1.00	27.80
3741	СВ	SER	396	93.770	30.518	42.465	1.00	39.24
3742	OG	SER	396	92.896	30.992	43.472	1.00	46.99
3743	H	SER	396	94.943	29.836	44.642	1.00	25.00
3744	HG	SER	396	92.971	31.948	43.546	1.00	25.00
3745	N	ASN	397	95.597	33.046	43.745	1.00	25.18
3746	CA	ASN	397	95.907	34.472	43.723	1.00	29.15
3747	C	ASN	397	97.333	34.739	44.226	1.00	27.87
3748	O CB	ASN	397	98.106	35.459	43.588	1.00	28.50
3749	CB	ASN	397	94.909	35.264	44.577	1.00	29.41
3750	CG CD1	ASN	397	95.146	36.770	44.505	1.00	35.89
3751	OD1	ASN	397	94.831	37.404	43.502	1.00	37.46
3752	ND2	ASN	397 397	95.715	37.343	45.564	1.00	30.17
3753	H	ASN		95.421	32.602	44.603	1.00	25.00
3754	1HD2	ASN	397	95.872	38.310	45.510	1.00	25.00
3755 3756	2HD2	ASN	397 398	95.953 97.682	36.794 34.123	46.336 45.351	$\frac{1.00}{1.00}$	25.00 26.89
3757	N CA	ALA ALA	398	98.988	34.300	45.980	1.00	24.87
3758	C	ALA	398	100.205	33.854	45.178	1.00	28.89
3759	0	ALA	398	100.203	34.358	45.395	1.00	31.67
3760	СВ	ALA	398	98.992	33.646	47.337	1.00	24.15
3761	H	ALA	398	97.035	33.533	45.770	1.00	25.00
3762	N	LEU	399	100.039	32.910	44.262	1.00	27.33
3763	CA	LEU	399	101.181	32.464	43.474	1.00	29.45
3764	C	LEU	399	101.755	33.589	42.617	1.00	32.27
3765	O	LEU	399	102.967	33.807	42.603	1.00	34.30
3766	CB	LEU	399	100.823	31.254	42.611	1.00	25.44
3767	CG	LEU	399	100.621	29.949	43.390	1.00	24.86
3768	CD1	LEU	399	100.172	28.853	42.451	1.00	20.68
3769	CD2	LEU	399	101.900	29.549	44.104	1.00	22.68
3770	H	LEU	399	99.159	32.500	44.121	1.00	25.00
3771	N	ALA	400	100.887	34.336	41.943	1.00	29.07
3772	CA	ALA	400	101.343	35.434	41.094	1.00	31.03

TABLE 11-continued

		In the	Absence of	f Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3773	С	ALA	400	101.939	36.601	41.882	1.00	29.34
3774	O	ALA	400	102.813	37.303	41.373	1.00	26.86
3775	СВ	ALA	400 400	100.215	35.925	40.192 42.022	1.00	32.40
3776 3777	H N	ALA THR	400	99.932 101.500	34.143 36.796	43.125	$\frac{1.00}{1.00}$	25.00 27.66
3778	CA	THR	401	102.024	37.896	43.929	1.00	30.92
3779	С	THR	401	103.505	37.728	44.303	1.00	35.35
3789	O	THR	401	104.118	38.649	44.847	1.00	36.05
3781	CB	THR	401	101.170	38.174	45.194	1.00	27.88
3782	OG1	THR	401	101.106	37.007	46.021	1.00	27.11
3783 3784	CG2 H	THR THR	401 401	99.768 100.837	38.593 36.193	44.803 43.521	$\frac{1.00}{1.00}$	25.60 25.00
3785	HG1	THR	401	101.990	36.769	46.324	1.00	25.00
3786	N	THR	402	104.076	36.558	44.016	1.00	30.98
3787	CA	THR	402	105.492	36.310	44.295	1.00	28.76
3788	С	THR	402	106.317	37.040	43.240	1.00	28.41
3789	O	THR	402	107.509	37.248	43.422	1.00	30.86
3790	CB	THR	402	105.861	34.807	44.173	1.00	24.70
3791 3792	OG1 CG2	THR THR	402 402	105.656 105.039	24.372 33.954	42.320 45.117	$\frac{1.00}{1.00}$	22.39 21.26
3793	H	THR	402	103.554	35.829	43.612	1.00	25.00
3794	HG1	THR	402	105.851	33.431	42.770	1.00	25.00
3795	N	THR	403	105.656	37.373	42.130	1.00	28.49
3796	CA	THR	403	106.207	38.045	40.946	1.00	25.51
3797	C	THR	403	107.032	37.101	40.077	1.00	27.25
3798	O	THR	403	107.499	37.495	39.009	1.00	29.50
3799	CB	THR	403	107.060	39.323	41.246	1.00	29.54
3800 3801	OG1 CG2	THR THR	403 403	108.335 106.339	38.954 40.267	41.781 42.203	$\frac{1.00}{1.00}$	24.64 26.87
3802	H	THR	403	100.339	37.143	42.203	1.00	25.00
3803	HG1	THR	403	108.246	38.543	42.630	1.00	25.00
3804	N	TYR	404	107.120	35.833	40.474	1.00	25.89
3805	CA	TYR	404	107.914	34.880	39.728	1.00	22.27
3806	C	TYR	404	107.544	34.611	38.272	1.00	24.30
3807	O	TYR	404	108.439	34.511	37.434	1.00	26.21
3808 3809	CB CG	TYR TYR	404 404	108.062 109.278	33.551 33.544	40.509 41.419	$\frac{1.00}{1.00}$	30.29 30.18
3810	CD1	TYR	404	109.278	34.736	41.922	1.00	31.88
3811	CD2	TYR	404	109.925	32.352	41.755	1.00	28.35
3812	CE1	TYR	404	110.937	34.747	42.732	1.00	30.99
3813	CE2	TYR	404	111.065	32.353	42.569	1.00	29.01
3814	CZ	TYR	404	111.563	33.558	43.051	1.00	29.70
3815	OH	TYR	404	112.683	33.593	43.847	1.00	27.42
3816 3817	H HH	TYR TYR	404 404	106.644 113.022	35.551 32.697	41.285 43.955	$\frac{1.00}{1.00}$	25.00 25.00
3818	N	TYR	405	106.253	34.508	37.952	1.00	24.23
3819	CA	TYR	405	105.844	34.306	36.553	1.00	24.60
3820	С	TYR	405	106.361	35.507	35.766	1.00	23.35
3821	O	TYR	405	106.912	35.378	34.672	1.00	23.93
3822	CB	TYR	405	104.317	34.292	36.406	1.00	25.23
3823	CG CD1	TYR	405	103.593	33.163	37.099 36.548	1.00	24.35
3824 3825	CD2	TYR	405 405	103.561 102.894	31.879		1.00	23.45
3825 3826	CD2 CE1	TYR	405 4005	102.894	33.390	38.282 37.161	1.00 $1.00$	23.55 23.28
3827	CE2	TYR	405	102.179	32.374	38.901	1.00	27.86
3828	CZ	TYR	405	102.155	31.111	38.337	1.00	26.80
3829	OH	TYR	405	101.428	30.121	38.956	1.00	26.06
3830	H	TYR	405	105.577	34.557	38.658	1.00	25.00
3831 3832	HH N	TYR TYR	405 406	101.510 106.160	29.308 36.676	38.452 36.363	$\frac{1.00}{1.00}$	25.00 23.37
3833	CA	TYR	406	106.553	37.964	35.813	1.00	21.93
3834	C	TYR	406	108.072	38.084	35.621	1.00	25.55
3835	Ö	TYR	406	108.535	38.439	34.538	1.00	24.19
3836	CB	TYR	406	106.021	39.047	36.751	1.00	22.96
3837	CG	TYR	406	106.379	40.488	36.408	1.00	21.90
3838	CD1	TYR	406	106.352	40.926	35.093	1.00	20.58
3839 3840	CD2 CE1	TYR TYR	406 406	106.703 106.634	41.375 42.252	37.416 34.796	1.00 $1.00$	27.70 21.17
3840 3841	CE1	TYR	406 406	106.634	42.252	37.128	1.00	24.51
3842	CZ	TYR	406	106.947	43.131	35.820	1.00	23.02
3843	OH	TYR	406	107.207	44.449	35.541	1.00	28.99
3844	H	TYR	406	105.729	36.670	37.238	1.00	25.00
3845	HH	TYR	406	107.398	44.923	36.351	1.00	25.00
3846	N	LEU	407	108.844	37.750	36.652	1.00	25.38

TABLE 11-continued

		In the	Absence o	f Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3847	CA	LEU	407	110.303	37.826	36.574	1.00	23.31
3848	C	LEU	407	110.873	36.825	35.575	1.00	20.18
3849	O	LEU	407	111.803	37.147	34.836	1.00	22.15
3850	CB	LEU	407	110.940	37.620	37.952	1.00	20.82
3851	CG	LEU	407	110.514	38.602	39.048	1.00	25.79
3852	CD1	LEU	407	111.362	38.376	40.287	1.00	24.12
3853	CD2	LEU	407	110.635	40.045	38.562	1.00	15.72
3854 3855	H N	LEU ALA	407 408	108.434 110.299	37.445 35.625	37.475 35.539	1.00 $1.00$	25.00 14.35
3856	CA	ALA	408	110.233	34.591	34.609	1.00	17.38
3857	C	ALA	408	110.520	35.083	33.183	1.00	22.75
3858	ŏ	ALA	408	111.382	34.931	32.318	1.00	25.21
3859	CB	ALA	408	109.991	33.293	34.852	1.00	17.26
3860	H	ALA	408	109.560	35.429	36.149	1.00	25.00
3861	N	THR	409	109.362	35.695	32.949	1.00	23.86
3862	CA	THR	409	109.037	36.228	31.632	1.00	22.93
3863	С	THR	409	110.012	37.353	31.294	1.00	25.61
3864	O	THR	409	110.507	37.443	30.165	1.00	26.75
3865	CB	THR	409	107.598	36.776	31.589	1.00	26.26
3866	OG1	THR	409	106.689	35.765	32.042	1.00	26.48
3867 3868	CG2 H	THR THR	409 409	107.222 108.709	37.170 35.780	30.173 33.672	1.00 $1.00$	18.58 25.00
3869	н HG1	THR	409	106.709	35.516	32.932	1.00	25.00
3870	N	THR	410	110.316	38.185	32.287	1.00	26.70
3871	CA	THR	410	111.233	39.299	32.095	1.00	26.67
3872	C	THR	410	112.650	38.835	31.757	1.00	29.09
3873	O	THR	410	113.298	39.411	30.877	1.00	29.16
3874	CB	THR	410	111.281	40.208	33.333	1.00	28.30
3875	OG1	THR	410	109.962	40.684	33.626	1.00	30.05
3876	CG2	THR	410	112.189	41.404	33.082	1.00	28.81
3877	H	THR	410	109.905	38.064	33.169	1.00	25.00
3878	HG1	THR	410	109.991	41.259	34.393	1.00	25.00
3879	N	SER	411	113.105	37.760	32.399	1.00	24.71
3880	CA	SER	411	114.452	37.254	32.155	1.00	24.18
3881 3882	C O	SER SER	411 411	114.688 115.822	36.909 36.984	30.687 30.204	1.00 1.00	25.08 27.75
3883	СВ	SER	411	114.753	36.046	33.043	1.00	20.40
3884	OG	SER	411	114.010	34.914	32.644	1.00	21.48
3885	Н	SER	411	112.534	37.287	33.041	1.00	25.00
3886	HG	SER	411	114.205	34.667	31.738	1.00	25.00
3887	N	TYR	412	113.613	36.573	29.979	1.00	23.79
3888	CA	TYR	412	113.692	36.227	28.562	1.00	24.39
3889	C	TYR	412	113.8774	37.44	27.655	1.00	25.24
3890	O	TYR	412	114.437	37.326	26.570	1.00	27.76
3891	CB	TYR	412	112.419	35.511	28.116	1.00	25.37
3892	CG	TYR	412	112.273	34.072	28.539	1.00	29.38
3893 3894	CD1	TYR	412	113.043	33.531	29.569	1.00	29.32
3895	CD2 CE1	TYR TYR	412 412	111.338 112.878	33.248 32.199	27.910 29.964	$\frac{1.00}{1.00}$	24.28 26.36
3896	CE2	TYR	412	111.169	31.927	28.291	1.00	25.08
3897	CZ	TYR	412	111.937	31.408	29.318	1.00	29.45
3898	OH	TYR	412	111.750	30.099	29.693	1.00	27.61
3899	H	TYR	412	112.738	36.552	30.423	1.00	25.00
3900	HH	TYR	412	112.347	29.879	30.418	1.00	25.00
3901	N	LEU	413	113.396	38.604	28.100	1.00	25.11
3902	CA	LEU	413	113.467	39.832	27.304	1.00	27.32
3903	С	LEU	413	114.835	40.149	26.726	1.00	30.49
3904	O	LEU	413	114.957	40.434	25.533	1.00	30.50
3905	CB	LEU	413	112.959	41.039	28.103	1.00	23.58
3906	CG CD1	LEU	413	111.476	41.081	28.478	1.00	31.45
3907 3908	CD1 CD2	LEU LEU	413 413	111.179 110.613	42.362 40.996	29.242 27.231	$\frac{1.00}{1.00}$	31.49 30.34
3909	H	LEU	413	112.980	38.636	28.989	1.00	25.00
3910	N	GLY	414	115.859	40.098	27.573	1.00	28.96
3911	CA	GLY	414	117.203	40.404	27.129	1.00	27.47
3912	C	GLY	414	117.990	39.233	26.586	1.00	28.88
3913	Ö	GLY	414	119.186	39.362	26.340	1.00	34.59
3914	Н	GLY	414	115.698	39.833	28.496	1.00	25.00
3915	N	MET	415	117.353	38.079	26.436	1.00	29.79
3916	CA	MET	415	118.043	36.909	25.906	1.00	29.75
3917	С	MET	415	117.861	36.868	24.393	1.00	35.70
3918	O	MET	415	116.795	36.522	23.893	1.00	39.21
3919	CB	MET	415	117.515	35.630	26.554	1.00	22.67
3920	CG	MET	415	117.728	35.581	28.050	1.00	23.60

TABLE 11-continued

		In the	Absence o	f Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	z	OCC	B-factor
3921	SD	MET	415	117.062	34.095	28.794	1.00	32.91
3922	CE	MET	415	118.255	32.896	28.242	1.00	24.83
3923 3924	H N	MET LYS	415 416	116.398 118.933	38.012 37.181	26.652 23.677	$\frac{1.00}{1.00}$	25.00 40.25
3924	CA	LYS	416	118.933	37.233	22.218	1.00	43.20
3926	C	LYS	416	118.370	36.031	21.466	1.00	42.08
3927	O	LYS	416	118.037	36.143	20.289	1.00	44.24
3928	CB	LYS	416	120.362	37.539	21.735	1.00	48.69
3929	CG	LYS	416	120.916	38.828	22.333	1.00	60.37
3930	CD	LYS	416	122.427	38.949	22.191	1.00	70.42
3931 3932	CE NZ	LYS LYS	416 416	122.936 124.412	40.173 40.319	22.949 22.863	$\frac{1.00}{1.00}$	72.06 78.00
3933	N	LYS	416	119.749	37.416	24.164	1.00	25.00
3934	1HZ	LYS	416	124.870	39.475	23.262	1.00	25.00
3935	2HZ	LYS	416	124.687	40.423	21.865	1.00	25.00
3938	3HZ	LYS	416	124.709	41.163	23.393	1.00	25.00
3937	N	SER	417	118.239	34.893	22.138	1.00	39.46
3938	CA	SER	417	117.706	33.698	21.491	1.00	36.96
3939 3940	C O	SER SER	417 417	116.247 115.637	33.395 32.518	21.833 21.226	1.00 $1.00$	34.08 35.80
3941	СВ	SER	417	118.580	32.488	21.823	1.00	39.51
3942	OG	SER	417	119.907	32.675	21.358	1.00	45.86
3943	H	SER	417	118.485	34.856	23.077	1.00	25.00
3944	HG	SER	417	120.288	33.461	21.743	1.00	25.00
3945	N	ALA	418	115.688	34.106	22.806	1.00	30.28
3946	CA	ALA	418	114.303	33.879	23.208	1.00	38.39
3947	С	ALA	418	113.331	34.250	22.087	1.00	40.58
3948 3949	O CB	ALA ALA	418 418	113.145 113.981	35.427 34.659	21.779 24.484	1.00 $1.00$	42.57 32.32
3950	Н	ALA	418	116.190	34.823	23.238	1.00	25.00
3951	N	THR	419	112.750	33.232	21.457	1.00	41.43
3952	CA	THR	419	111.799	33.420	220.362	1.00	41.21
3953	C	THR	419	110.357	33.504	20.865	1.00	40.11
3954	O	THR	419	110.077	33.235	22.036	1.00	39.88
3955	CB	THR	419	111.892	32.264	19.338	1.00	39.89
3956	OG1 CG2	THR THR	419 419	111.666 113.261	31.016 32.238	20.005 18.672	1.00 $1.00$	50.18 37.82
3957 3958	H	THR	419	112.987	32.335	21.728	1.00	25.00
3959	HG1	THR	419	112.370	30.881	20.644	1.00	25.00
3960	N	GLU	420	109.443	33.848	19.963	1.00	39.64
3961	CA	GLU	420	108.027	33.958	20.292	1.00	39.75
3962	С	GLU	420	107.496	32.650	20.871	1.00	35.49
3963	O	GLU	420	106.718	32.652	21.828	1.00	35.76
3964	CB	GLU	420 420	107.222	34.321	19.041	1.00	46.65 56.30
3965 3966	CG CD	GLU GLU	420 420	106.741 105.668	35.765 36.081	18.980 20.015	1.00 $1.00$	65.12
3967	OE1	GLU	420	104.685	35.311	20.130	1.00	62.03
3968	OE2	GLU	420	105.804	37.112	20.707	1.00	72.02
3969	H	GLU	420	109.732	34.021	19.047	1.00	25.00
3970	N	GLN	421	107.938	31.537	20.291	1.00	32.71
3971	CA	GLN	421	107.520	30.206	20.722	1.00	36.14
3972	С	GLN	421	107.883	29.959	22.184	1.00	36.63
3973 3974	CB	GLN GLN	421 421	107.105	29.365	22.936 19.830	1.00 $1.00$	40.03
3975	CG	GLN	421	107.6[]2	29.086	18.398	1.00	53.07
3976	CD	GLN	421	107.819	30.390	17.638	1.00	62.17
3977	OE1	GLN	421	108.877	31.017	17.711	1.00	65.66
3978	NE2	GLN	421	106.788	30.815	16.917	1.00	68.97
3979	H	GLN	421	108.580	31.620	19.555	1.00	25.00
3980	1HE2	GLN	421	106.906	31.650	16.419	1.00	25.00
3981	2HE2	GLN	421	105.963	30.289	16.903	1.00	25.00
3982 3983	N CA	ASP ASP	422 422	109.052 109.521	30.448 30.292	22.589 23.960	1.00 $1.00$	34.98 32.53
3983 3984	CA	ASP	422	109.321	31.039	24.924	1.00	32.33 30.54
3985	Õ	ASP	422	108.272	30.525	25.992	1.00	33.81
3986	СВ	ASP	422	110.972	30.767	24.085	1.00	28.20
3987	CG	ASP	422	111.929	29.945	23.233	1.00	28.18
3988	OD1	ASP	422	111.755	28.710	23.148	1.00	33.24
3989	OD2	ASP	422	112.855	30.529	22.638	1.00	33.81
3990 3001	H	ASP	422	109.610	30.933	21.948	1.00	25.00
3991 3992	N CA	PHE PHE	423 423	108.166 107.261	32.229 33.022	24.523 25.348	1.00 $1.00$	29.33 28.15
3992 3993	CA	PHE	423	107.201	32.373	25.407	1.00	29.51
3994	ŏ	PHE	423	105.205	32.424	26.441	1.00	30.32

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
3995	CB	PHE	423	107.143	34.448	24.808	1.00	31.32
3996	CG	PHE	423	108.275	35.353	25.214	1.00	27.99
3997	CD1	PHE	423	108.227	36.045	26.421	1.00	26.74
3998	CD2	PHE	423	109.375	35.529	24.387	1.00	26.53
3999	CE1	PHE	423	109.255	36.900	26.794	1.00	21.72
4000	CE2	PHE	423	110.412	36.385	24.753	1.00	23.04
4001	CZ	PHE	423	110.350	37.070	25.958	1.00	23.46
4002	H	PHE	423	108.456	32.575	23.652	1.00	25.00
4000	N	PHE	424	105.450	31.771	24.297	1.00	30.30
4004	CA	GLU	424	104.140	31.107 29.911	24.233	1.00	36.62
4005 4006	C O	GLU GLU	424 424	104.128 103.131	29.651	25.172 25.343	1.00	34.48 34.44
4007	СВ	GLU	424	103.131	30.661	22.805	$\frac{1.00}{1.00}$	42.57
4007	CG	GLU	424	103.525	31.813	21.839	1.00	63.38
4009	CD	GLU	424	103.322	31.357	20.414	1.00	70.12
4010	OE1	GLU	424	103.895	30.329	19.988	1.00	72.19
4011	OE2	GLU	424	102.543	32.039	19.715	1.00	79.30
4012	H	GLU	424	106.023	31.773	23.503	1.00	25.00
4013	N	TRP	425	105.242	29.187	25.221	1.00	32.02
4014	CA	TRP	425	105.367	28.038	26.107	1.00	28.78
4015	С	TRP	425	105.262	28.531	27.553	1.00	33.19
4016	O	TRP	425	104.518	27.974	28.365	1.00	30.47
4017	CB	TRP	425	106.719	27.352	25.881	1.00	29.01
4018	CG	TRP	425	107.077	26.340	26.927	1.00	29.69
4019	CD1	TRP	425	106.621	25.058	27.019	1.00	28.88
4020	CD2	TRP	425	107.970	26.530	28.034	1.00	29.87
4021	NE1	TRP	425	107.171	24.437	28.116	1.00	29.58
4022	CE2	TRP	425	108.003	25.316	28.757	1.00	32.75
4023	CE3	TRP	425 425	108.742	27.609	28.488	1.00	30.70
4024 4025	CZ2 CZ3	TRP TRP	425 425	108.781 109.514	25.149 27.444	29.912 29.638	$\frac{1.00}{1.00}$	27.26 26.04
4026	CH2	TRP	425	109.514	26.222	30.335	1.00	27.12
4027	H	TRP	425	105.993	29.429	24.639	1.00	25.00
4028	HE1	TRP	425	106.983	23.517	28.395	1.00	25.00
4029	N	LEU	426	105.974	29.615	27.848	1.00	31.46
4030	CA	LEU	426	105.994	30.186	29.188	1.00	28.35
4031	C	LEU	426	104.627	30.692	29.650	1.00	31.98
4032	O	LEU	426	104.293	30.585	30.832	1.00	29.19
4033	CB	LEU	426	107.039	31.302	29.268	1.00	21.80
4034	CG	LEU	426	107.525	31.703	30.664	1.00	25.44
4035	CD1	LEU	426	108.240	30.535	31.331	1.00	20.60
4036	CD2	LEU	426	108.454	32.900	30.560	1.00	23.92
4037 4038	H N	LEU	426 427	106.512 103.824	30.035	27.141 28.720	1.00	25.00 33.95
4039	CA	SER SER	427	103.824	31.208 31.722	29.066	$\frac{1.00}{1.00}$	33.39
4040	C	SER	427	101.502	30.647	29.502	1.00	30.91
4041	ŏ	SER	427	100.515	30.951	30.170	1.00	31.38
4042	СB	SER	427	101.917	32.568	27.925	1.00	37.83
4043	OG	SER	427	101.970	31.892	26.683	1.00	46.06
4044	H	SER	427	104.124	31.247	27.790	1.00	25.00
4045	HG	SER	427	102.886	31.692	26.470	1.00	25.00
4046	N	LYS	428	101.780	29.392	29.151	1.00	30.99
4047	CA	LYS	428	100.914	28.271	29.518	1.00	30.38
4048	C	LYS	428	101.124	27.817	30.964	1.00	33.22
4049	O	LYS	428	100.505	26.845	31.410	1.00	34.36
4050	CB	LYS	428	101.166	27.070	28.601	1.00	33.08
4051	CG	LYS	428	100.690	27.213	27.166	1.00	40.80
4052 4053	CD CE	LYS LYS	428 428	100.885 100.314	25.888 25.910	26.433 25.002	$\frac{1.00}{1.00}$	48.04 54.20
4054	NZ	LYS	428	100.314	24.571	24.377	1.00	59.29
4055	H	LYS	428	102.587	29.209	28.628	1.00	25.00
4056	1HZ	LYS	428	99.919	23.866	24.937	1.00	25.00
4057	2HZ	LYS	428	100.037	24.613	23.418	1.00	25.00
4058	3HZ	LYS	428	101.440	24.300	24.323	1.00	25.00
4059	N	ASN	429	101.992	28.518	31.693	1.00	36.19
4060	CA	ASN	429	102.313	28.172	33.081	1.00	31.37
4061	C	ASN	429	102.855	26.740	33.172	1.00	29.89
4062	0	ASN	429	102.272	25.882	33.839	1.00	25.78
4063	CB	ASN	429	101.092	28.334	33.995	1.00	32.22
4064	CG OD1	ASN	429	100.814	29.782	34.358	1.00	36.78
4065	OD1	ASN	429	101.488	30.699	33.894	1.00	40.44
4066	ND2	ASN	429 429	99.826	29.991	35.215 31.299	1.00	41.36 25.00
4067 4068	H 1HD2	ASN ASN	429 429	102.436 99.643	29.294 30.918	31.299	$\frac{1.00}{1.00}$	25.00
-1000	1111/2	11311	727	//.UTJ	20.710	55.777	1.00	25.00

TABLE 11-continued

		In th	e Absence of	f Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
4069	2HD2	ASN	429	99.331	29.226	35.566	1.00	25.00
4070	N	PRO	430	103.997	26.472	32.508	1.00	29.30
4071	CA	PRO	430	104.649	25.157	32.492	1.00	25.30
4072	С	PRO	430	104.897	24.669	33.913	1.00	27.01
4073	O	PRO	430	105.218	25.463	34.801	1.00	27.38
4074	CB	PRO	430	105.975	25.447	31.799	1.00	24.23
4075 4076	CG CD	PRO PRO	430 430	105.664 104.835	26.610 27.468	30.932 31.820	1.00 1.00	30.06 30.25
4077	N N	LYS	431	104.833	23.358	34.108	1.00	25.65
4078	CA	LYS	431	105.020	22.774	35.426	1.00	25.93
4079	C	LYS	431	106.308	23.248	36.107	1.00	24.81
4080	O	LYS	431	106.297	23.570	37.292	1.00	24.38
4081	CB	LYS	431	105.000	21.252	35.325	1.00	26.70
4082	CG	LYS	431	104.584	20.547	36.604	1.00	40.34
4083	CD	LYS	431	104.361	19.068	36.330	1.00	52.65
4084	CE	LYS	431	103.775	18.345	37.531	1.00	60.92
4085	NZ	LYS	431	103.587	16.892	37.247	1.00	58.59
4086	H 1117	LYS	431	104.610	22.777	33.352	1.00	25.00
4087 4088	1HZ 2HZ	LYS LYS	431 431	102.939 104.506	16.775 16.462	36.442 37.018	1.00 1.00	25.00 25.00
4089	3HZ	LYS	431	103.189	16.424	38.086	1.00	25.00
4090	N	ILE	432	107.401	23.338	35.353	1.00	25.03
4091	CA	ILE	432	108.667	23.774	35.934	1.00	21.64
4092	C	ILE	432	108.561	25.188	36.505	1.00	24.42
4093	O	ILE	432	109.058	25.459	37.602	1.00	25.07
4094	CB	ILE	432	109.847	23.646	34.928	1.00	22.09
4095	CG1	ILE	432	111.179	23.898	35.647	1.00	20.75
4096	CG2	ILE	432	109.662	24.587	33.739	1.00	20.53
4097	CD1	ILE	432	112.403	23.511	34.838	1.00	17.10
4098	H	ILE	432	107.354	23.103	34.410	1.00	25.00
4099	N	LEU	433	107.868	26.070	35.788	1.00	25.32
4100	CA C	LEU	433 433	107.674	27.448	36.240	1.00	23.97
4101 4102	0	LEU LEU	433	106.758 107.051	27.446 28.078	37.464 38.483	1.00 1.00	27.11 28.11
4102	СВ	LEU	433	107.051	28.298	35.126	1.00	24.53
4104	CG	LEU	433	106.721	29.754	35.473	1.00	28.30
4105	CD1	LEU	433	107.968	30.488	35.945	1.00	23.75
4106	CD2	LEU	433	106.108	30.456	34.265	1.00	27.09
4107	H	LEU	433	107.456	25.788	34.948	1.00	25.00
4108	N	GLU	434	105.667	26.698	37.360	1.00	26.39
4109	CA	GLU	434	104.690	26.566	38.429	1.00	28.21
4110	С	GLU	434	105.393	26.139	39.723	1.00	25.73
4111	O	GLU	434	105.159	26.711	40.790	1.00	25.92
4112	CB	GLU	434	103.656	25.510	38.027	1.00	40.62
4113	CG	GLU GLU	434 434	102.371	25.510	38.835	1.00	59.16
4114 4115	CD OE1	GLU	434	101.447 101.135	26.650 26.803	38.457 37.255	1.00 1.00	69.11 76.28
4116	OE2	GLU	434	101.026	27.391	39.366	1.00	78.90
4117	H	GLU	434	105.516	26.206	36.531	1.00	25.00
4118	N	ALA	435	106.272	25.147	39.614	1.00	23.71
4119	CA	ALA	435	107.015	24.632	40.764	1.00	21.25
4120	С	ALA	435	107.915	25.704	41.377	1.00	21.57
4121	O	ALA	435	107.973	25.864	42.599	1.00	21.94
4122	CB	ALA	435	107.838	23.424	40.353	1.00	17.11
4123	H	ALA	435	106.427	24.748	38.737	1.00	25.00
4124	N	SER	436	108.603	26.448	40.519	1.00	20.07
4125	CA	SER	436	109.486	27.510	40.969	1.00	21.72
4126	С	SER	436	108.676	28.531	41.759	1.00	23.51
4127 4128	O CB	SER SER	436 436	109.095 110.147	28.979 28.179	42.832 39.765	1.00 1.00	25.11 21.38
4129	OG	SER	438	111.040	29.196	40.173	1.00	36.67
4130	Н	SER	436	108.521	26.272	39.556	1.00	25.00
4131	HG	SER	436	110.568	29.867	40.669	1.00	25.00
4132	N	VAL	437	107.501	28.876	41.235	1.00	23.94
4133	CA	VAL	437	106.622	29.846	41.880	1.00	18.07
4134	C	VAL	437	106.134	29.330	43.226	1.00	19.00
4135	O	VAL	437	106.179	30.040	44.227	1.00	24.40
4136	CB	VAL	437	105.410	30.192	40.990	1.00	22.63
4137	CG1	VAL	437	104.498	31.163	41.709	1.00	24.13
4138	CG2	VAL	437	105.879	30.794	39.677	1.00	13.85
4139	H	VAL	437	107.218	28.463	40.391	1.00	25.00
4140 4141	N C4	ILE	438	105.693	28.076	43.249	1.00	22.97
4141 4142	CA C	ILE ILE	438 438	105.204 106.279	27.443 27.478	44.472 45.566	1.00 1.00	25.17 27.56
4142		ILL	+30	100.279	21.410	+5.500	1.00	27.30

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
4143	O	ILE	438	105.996	27.831	46.718	1.00	25.37
4144	CB	ILE	438	104.776	25.975	44.200	1.00	28.36
4145	CG1	ILE	438	103.565	25.952	43.262	1.00	33.63
4146	CG2	ILE	438	104.457	25.255	45.505	1.00	31.72
4147	CD1	ILE	438	103.130	24.569	42.836	1.00	33.54
4148	H	ILE	438	105.697	27.559	42.417	1.00	25.00
4149	N	ILE	439	107.512	27.134	45.196	1.00	27.24
4150	CA	ILE	439	108.635	27.130	46.135	1.00	24.88
4151	С	ILE	439	108.769	28.502	46.787	1.00	20.20
4152	O	ILE	439	108.842	28.610	48.007	1.00	20.39
4153 4154	CB CC1	ILE ILE	439	109.961	26.739	45.429	1.00	23.09
4154	CG1 CG2	ILE	439 439	109.915 111.154	25.264 26.989	45.023 46.345	$\frac{1.00}{1.00}$	21.73 17.14
4156	CD1	ILE	439	111.134	24.859	44.043	1.00	22.40
4157	H	ILE	439	107.670	26.870	44.265	1.00	25.00
4158	N	CYS	440	108.763	29.552	45.974	1.00	21.23
4159	CA	CYS	440	103.873	30.901	46.508	1.00	23.58
4160	C	CYS	440	107.718	31.209	47.458	1.00	27.42
4161	Ö	CYS	440	107.933	31.707	48.563	1.00	29.41
4162	CB	CYS	440	108.897	31.928	45.376	1.00	26.26
4163	SG	CYS	440	109.015	33.625	45.934	1.00	17.39
4164	H	CYS	440	108.685	29.406	45.006	1.00	25.00
4165	N	ARG	441	106.502	30.869	47.038	1.00	30.12
4166	CA	ARG	441	105.295	31.118	47.825	1.00	28.33
4167	C	ARG	4441	105.280	30.448	49.197	1.00	28.78
4168	O	ARG	441	105.225	31.125	50.223	1.00	28.38
4169	CB	ARG	441	104.056	30.693	47.031	1.00	26.28
4170	CG	ARG	441	102.722	30.927	47.734	1.00	26.09
4171	CD	ARG	441	102.312	32.391	47.725	1.00	32.68
4172	NE	ARG	441	103.001	33.195	48.731	1.00	35.67
4173	CZ	ARG	441	103.243	34.499	48.611	1.00	31.42
4174	NH1	ARG	441 441	102.861	35.159	47.526	1.00	25.83
4175 4176	NH2 H	ARG ARG	441	103.851 106.413	35.153 30.429	49.591 46.165	1.00 $1.00$	32.87 25.00
4177	н НЕ	ARG	441	103.307	32.748	49.547	1.00	25.00
4178	1HH1	ARG	441	102.391	34.678	46.792	1.00	25.00
4179	2HH1	ARG	441	103.043	36.138	47.443	1.00	25.00
4180	1HH2	ARG	441	104.133	34.668	50.417	1.00	25.00
4181	2HH2	ARG	441	104.032	36.133	49.498	1.00	25.00
4182	N	VAL	442	105.329	29.120	49.220	1.00	28.03
4183	CA	VAL	442	105.289	28.392	50.484	1.00	29.35
4184	C	VAL	442	106.443	28.708	51.430	1.00	29.48
4185	O	VAL	442	106.248	28.754	52.644	1.00	30.67
4186	CB	VAL	442	105.171	26.864	50.272	1.00	28.34
4187	CG1	VAL	442	103.906	26.546	49.489	1.00	20.42
4188	CG2	VAL	442	106.394	26.322	49.562	1.00	28.17
4189	H	VAL	442	105.405	28.625	48.376	1.00	25.00
4190	N	ILE	443	107.635	28.941	50.885	1.00	32.67
4191 4192	CA	ILE	443 443	108.788	29.266 30.6722	51.722 52.283	1.00	32.60 34.90
4192	C O	ILE ILE	443	108.619 108.866	30.908	53.469	$\frac{1.00}{1.00}$	33.18
4194	СВ	ILE	443	110.134	29.150	50.955	1.00	34.36
4195	CG1	ILE	443	110.394		50.574	1.00	27.67
4196	CG2	ILE	443	111.290		51.822	1.00	28.68
4197	CD1	ILE	443	110.456		51.765	1.00	31.24
4198	H	ILE	443	107.746	28.891	49.910	1.00	25.00
4199	N	ASP	444	108.170	31.599	51.441	1.00	33.84
4200	CA	ASP	444	107.954	32.968	51.889	1.00	35.72
4201	C	ASP	444	106.935	32.959	53.023	1.00	38.14
4202	O	ASP	444	107.184	33.520	54.091	1.00	37.60
4203	CS	ASP	444		33.848	50.744	1.00	39.61
4204	CG	ASP	444	107.110	35.260	51.199	1.00	48.15
4205	OD11	ASP	444	105.972	35.485	51.667	1.00	51.59
4206	OD2	ASP	444	107.980	36.147	51.091	1.00	51.59
4207	H	ASP	444	107.985	31.364	50.507	1.00	25.00
4208 4209	N C4	ASP	445 445	105.812 104.749	32.279 32.199	52.803 53.803	1.00 $1.00$	40.33 39.63
4209	CA C	ASP ASP	445 445	104.749	32.199	55.124	1.00	39.63 36.43
4210	O	ASP ASP	445	103.221	32.060	56.195	1.00	38.68
4211	СВ	ASP	445	104.820		53.259	1.00	34.74
4212	CG	ASP	445	102.867	32.115	52.087	1.00	37.47
4214	OD1	ASP	445	103.173	33.301	51.818	1.00	32.84
4215	OD2	ASP	445	102.022	31.474	51.429	1.00	34.75
4216	H	ASP	445	105.694		51.945	1.00	25.00

TABLE 11-continued

		In the	Absence o	f Bound S	ubstrate			
Atom Typ	e Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
4217	N	THR	446	106.061	30.574	55.047	1.00	36.21
4218	CA	THR	446	106.586	29.933	56.247	1.00	34.20
4219	С	THR	446	107.469	30.902	57.034	1.00	34.56
4220	О	THR	446	107.396	30.968	58.259	1.00	36.69
4221	CB	THR	446	107.398	28.674	55.890	1.00	29.80
4222	OG1	THR	446	106.545	27.735	55.227	1.00	31.60
4223	CG2	THR	446	107.970	28.029	57.136	1.00	31.33
4224 4225	H HG1	THR THR	446 446	106.322 107.041	30.221 26.946	54.170 54.992	1.00 $1.00$	25.00 25.00
4226	N	ALA	447	108.266	31.687	56.318	1.00	39.84
4227	CA	ALA	447	109.172	32.644	56.941	1.00	42.07
4228	C	ALA	447	108.495	33.902	57.483	1.00	45.04
4229	O	ALA	447	108.675	34.259	58.647	1.00	48.71
4230	CB	ALA	447	110.2776	33.025	55.959	1.00	36.91
4231	H	ALA	447	108.237	31.624	55.337	1.00	25.00
4232	N	THR	448	107.708	34.565	56.644	1.00	50.59
4233	CA	THR	448	107.036	35.799	57.039	1.00	49.97
4234	C	THR	448	105.729	35.644	57.818	1.00	51.43
4235	O	THR	448	105.100	36.640	58.159	1.00	56.61
4236	CB	THR	448	106.790	36.715	55.811	1.00	47.11
4237 4238	OG1 CG2	THR THR	448 448	106.095 108.106	35.986 37.214	54.793 55.245	$\frac{1.00}{1.00}$	50.09 47.36
4239	H	THR	448	107.555	34.230	55.741	1.00	25.00
4240	HG1	THR	448	105.951	36.556	54.034	1.00	25.00
4241	N	TYR	449	105.352	34.415	58.157	1.00	51.91
4242	CA	TYR	449	104.103	34.182	58.881	1.00	53.63
4243	С	TYR	449	103.927	35.020	60.148	1.00	58.13
4244	O	TYR	449	102.939	35.745	60.282	1.00	57.88
4245	CB	TYR	449	103.926	32.696	59.218	1.00	50.75
4246	CG	TYR	449	102.674	32.406	60.025	1.00	55.96
4247	CD1	TYR	449	101.419	32.856	59.596	1.00	60.14
4248	CD2	TYR	449	102.740	31.716	61.235	1.00	56.00
4249	CE1	TYR	449	100.273	32.629	60.353	1.00	55.79
4250	CE2	TYR	449	101.605	31.483	62.000	1.00	57.85
4251 4252	CZ OH	$\mathrm{TYR}$ $\mathrm{TYR}$	449 449	100.375 99.250	31.943 31.724	61.554 62.316	$\frac{1.00}{1.00}$	60.06 61.67
4253	Н	TYR	449	105.912	33.653	57.905	1.00	25.00
4254	HH	TYR	449	98.487	32.104	61.878	1.00	25.00
4255	N	GLU	450	104.883	34.927	61.067	1.00	64.21
4256	CA	GLU	450	104.810	35.662	62.329	1.00	67.19
4257	C	GLU	450	104.604	37.167	62.173	1.00	68.02
4258	O	GLU	450	103.698	37.742	62.781	1.00	68.60
4259	CB	GLU	450	106.053	35.386	63.178	1.00	73.69
4260	CG	GLU	450	106.228	33.917	63.557	1.00	87.48
4261	CD	GLU	450	104.938	33.311	64.211	1.00	95.20
4262	OE1	GLU	450	104.311	34.009	65.000	1.00	100.27
4263	OE2	GLU	450	104.690	32.129	63.934	1.00	96.38
4264 4265	H	GLU	450 451	105.651 105.427	34.348 37.790	60.883	1.00	25.00
4266	N CA	VAL VAL	451	105.351	39.228	61.336 61.091	$\frac{1.00}{1.00}$	69.29 69.79
4267	C	VAL	451	104.011	39.634	60.479	1.00	71.86
4268	ŏ	VAL	451	103.383	40.594	60.925	1.00	73.87
4269	CB	VAL	451	106.482	39.692	60.149	1.00	68.77
4270	CG1	VAL	451	106.490	41.211	60.036	1.00	67.70
4271	CG2	VAL	451	107.825	39.180	60.647	1.00	74.55
4272	H	VAL	451	106.100	37.263	60.866	1.00	25.00
4273	N	GLU	452	103.572	38.893	59.467	1.00	73.34
4274	CA	GLU	452	102.311	39.191	58.798	1.00	74.52
4275	С	GLU	452	101.096	38.987	59.700	1.00	75.62
4276	O	GLU GLU	452	100.107	39.715	59.575	1.00	75.03
4277 4278	CB CG	GLU	452 452	102.176 103.194	38.385 38.774	57.502 56.427	$\frac{1.00}{1.00}$	73.01 77.67
4279	CD	GLU	452	103.194	38.007	55.118	1.00	81.85
428O	OE1	GLU	452	103.032	36.858	55.137	1.00	80.20
4281	OE2	GLU	452	102.337	38.558	54.061	1.00	82.09
4282	H	GLU	452	104.100	38.124	59.165	1.00	25.00
4283	N	LYS	453	101.173	38.038	60.631	1.00	78.11
4284	CA	LYS	453	100.050	37.799	61.538	1.00	81.43
4285	C	LYS	453	99.887	38.943	62.532	1.00	84.49
4286	O	LYS	453	98.768	39.393	62.783	1.00	88.82
4287	CB	LYS	453	100.176	36.468	62.280	1.00	79.17
4288	CG	LYS	453	98.907	36.134	63.054	1.00	78.09
4289	CD	LYS	453	98.928	34.759	63.674	1.00	79.02
4290	CE	LYS	453	97.583	34.462	64.319	1.00	81.56

TABLE 11-continued

	In the Absence of Bound Substrate							
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
4291	NZ	LYS	453	97.525	33.093	64.899	1.00	87.65
4292	H	LYS	453	101.981	37.486	60.701	1.00	25.00
4293	1HZ	LYS	453	98.261	32.991	665.627	1.00	25.00
4294	2HZ	LYS	453	96.590	32.937	65.327	1.00	25.00
4295	3HZ	LYS	453	97.682	32.390	64.148	1.00	25.00
4296	N	SER	454	100.996	39.424	63.088	1.00	84.86
4297	CA	SER	454	100.943	40.535	64.037	1.00	83.61
4298	C	SER	454	100.521	41.835	63.336	1.00	83.52
4299	O	SER	454	100.210	42.830	63.991	1.00	82.37
4300	CB	SER	454	102.286	40.703	64.755	1.00	81.64
4301	OG	SER	454	103.363	40.771	63.838	1.00	82.19
4302	H	SER	454	101.863	39.025	62.862	1.00	25.00
4303	HG	SER	454	103.411	39.963	63.320	1.00	25.00
4304	N	ARG	455	100.515	41.813	62.003	1.00	83.63
4305	CA	ARG	455	100.102	42.961	61.197	1.00	86.97
4306	C	ARRG	455	98.616	42.870	60.854	1.00	89.33
4307	O	ARG	455	98.073	43.748	60.183	1.00	89.55
4308	CB	ARG	455	100.892	43.025	59.891	1.00	86.51
4309	CG	ARG	455	102.319	43.493	60.014	1.00	89.64
4310	CD	ARG	455	102.926	43.581	58.632	1.00	98.07
4311	NE	ARG	455	104.296	44.078	58.647	1.00	109.66
4312	CZ	ARG	455	104.976	44.415	57.555	1.00	114.68
4313	NH1	ARG	455	104.411	44.309	56.357	1.00	117.44
4314	NH2	ARG	455	106.220	44.863	57.659	1.00	113.39
4315	H	ARG	455	100.816	41.007	61.542	1.00	25.00
4316	HE	ARG	455	104.745	44.172	59.512	1.00	25.00
4317	1HH1	ARG	455	103.471	43.978	56.270	1.00	25.00
4318	2HH1	ARG	455	104.920	44.568	55.536	1.00	25.00
4319	1HH2	ARG	455	106.646	44.949	58.558	1.00	25.00
4320	2HH2	ARG	455	108.724	45.120	56.834	1.00	25.00
4321	N	GLY	456	97.980	41.773	61.259	1.00	90.87
4322	CA	GLY	456	96.566	41.584	60.989	1.00	90.27
4323	C	GLY	456	96.256	40.876	59.681	1.00	91.84
4324	O	GLY	456	95.087	40.636	59.371	1.00	90.99
4325	H	GLY	456	98.464	41.081	61.752	1.00	25.00
4326	N	GLN	457	97.290	40.528	58.917	1.00	92.16
4327	CA	GLN	457	97.107	39.842	57.638	1.00	90.85
4328	C	GLN	457	96.662	38.395	57.846	1.00	89.74
4329	O	GLN	457	97.442	37.462	57.659	1.00	92.66
4330	CB	GLN	457	98.402	39.868	56.817	1.00	89.90
4331	CG	GLN	457	98.905	41.257	56.457	1.00	95.19
4332	CD	GLN	457	100.145	41.221	55.576	1.00	99.21
4333	OE1	GLN	457	100.325	40.303	54.775	1.00	102.27
4334	NE2	GLN	457	101.002	42.225	55.718	1.00	97.71
4335	H	GLN	457	98.199	40.716	59.228	1.00	25.00
4336	1HE2	GLN	457	101.798	42.200	55.151	1.00	25.00
4337	2HE2	GLN	457	100.809	42.931	56.364	1.00	25.00
4338	N	ILE	458	95.397	38.207	58.209	1.00	88.19
4339	CA	ILE	458	94.859	36.867	58.439	1.00	82.29
4340	C	ILE	458	94.715	36.043	57.159	1.00	75.34
4341	O	ILE	458	94.305	34.887	57.205	1.00	73.69
4342	CB	ILE	458	93.510	36.907	59.199	1.00	85.78
4343	CG1	ILE	458	92.566	37.933	58.562	1.00	86.17
43444	CG2	ILE	458	93.751	37.191	60.681	1.00	85.33
4345	CD1	ILE	458	91.240	38.069	59.277	1.00	89.41
4346	H	ILE	458	94.823	38.995	58.338	1.00	25.00
4347	N	ALA	459	95.077	36.632	56.025	1.00	68.80
4348	CA	ALA	459	95.007	35.938	54.747	1.00	62.63
4349	C	ALA	459	96.368	35.324	54.389	1.00	62.33
4350	O	ALA	459	96.664	35.093	53.216	1.00	62.33
4351	CB	ALA	459	94.549	36.893	53.653	1.00	62.28
4352	H	ALA	459	95.387	37.554	56.036	1.00	25.00
4353	N	THR	460	97.210	35.097	55.396	1.00	59.27
4354	CA	THR	460	98.531	34.513	55.170	1.00	57.11
4355	С	THR	460	98.424	33.034	54.826	1.00	53.62
4356	O	THR	460	97.587	32.319	55.383	1.00	53.81
4357	CB	THR	460	99.453	34.671	56.400	1.00	58.87
4358	OG1	THR	460	98.763	34.257	57.588	1.00	56.81
4359	CG2	THR	460	99.901	36.100	56.541	1.00	60.73
4360	H	THR	460	96.933	35.314	56.305	1.00	25.00
4361	HG1	THR	460	98.004	34.838	57.727	1.00	25.00
4362	N	GLY	461	99.298	32.574	53.937	1.00	44.28
4363	CA	GLY	461	99.289	31.184	53.526	1.00	37.76
4364	С	GLY	461	99.138	30.186	54.652	1.00	39.21

TABLE 11-continued

		In th	e Absence of	f Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
4365	О	GLY	461	98.265	29.318	54.599	1.00	37.71
4366	H	GLY	461	99.947	33.183	53.532	1.00	25.00
4367	N	ILE	462	99.965	30.320	55.684	1.00	41.07
4368	CA	ILE	462	99.915	29.405	56.821	1.00	43.03
4369	C O	ILE	462	98.567	29.461	57.539	1.00 1.00	43.30
4370 4371	СВ	ILE ILE	462 462	98.063 101.079	28.430 29.658	57.991 57.814	1.00	44.23 35.94
4372	CG1	ILE	462	102.418	29.419	57.116	1.00	34.23
4373	CG2	ILE	462	100.979	28.720	59.011	1.00	26.77
4374	CD1	ILE	462	102.625	27.987	56.675	1.00	31.75
4375	H	ILE	462	100.618	31.046	55.680	1.00	25.00
4376	N	GLU	463	97.972	30.652	57.602	1.00	47.45
4377	CA	GLU	463	96.673	30.842	58.253	1.00	48.26
4378	C	GLU	463	95.600	30.064	57.495	1.00	43.95
4379	O	GLU	463	94.876	29.250	58.077	1.00	45.29
4380	CB	GLU	463	96.307	32.329	58.291	1.00	55.50
4381	CG	GLU	463 463	95.120	32.664	59.182	1.00 $1.00$	60.77
4382 4383	CD OE1	GLU GLU	463 463	95.448 96.195	32.555 33.419	60.656 61.166	1.00	66.36 70.77
4384	OE2	GLU	463	94.957	31.605	61.303	1.00	66.97
4385	H	GLU	463	98.403	31.430	57.193	1.00	25.00
4386	N	CYS	464	95.534	30.295	56.186	1.00	38.49
4387	CA	CYS	464	94.575	29.616	55.322	1.00	40.30
4388	C	CYS	464	94.751	28.113	55.451	1.00	40.72
4389	O	CYS	464	93.778	27.364	55.550	1.00	43.89
4390	CB	CYS	464	94.798	30.010	53.860	1.00	33.19
4391	SG	CYS	464	94.721	31.780	53.533	1.00	41.13
4392	H	CYS	464	96.143	30.954	55.789	1.00	25.00
4393	N CA	CYS	465	96.007	27.682	55.464	1.00	42.83
4394 4395	CA	CYS CYS	465 465	96.337 95.791	26.271 25.654	55.557 56.852	1.00 $1.00$	45.44 48.27
4396	o	CYS	465	95.165	24.591	56.818	1.00	47.36
4397	СВ	CYS	465	97.850	26.074	55.487	1.00	39.55
4398	SG	CYS	465	98.332	24.350	55.349	1.00	40.18
4399	H	CYS	465	96.735	28.334	55.396	1.00	25.00
4400	N	MET	466	96.000	26.340	57.972	1.00	51.35
4401	CA	MET	466	95.531	25.854	59.267	1.00	55.36
4402	C	MET	466	94.019	25.699	59.312	1.00	56.30
4403	O	MET	466	93.512	24.647	59.698	1.00	55.90
4404	CB	MET	466	95.977	26.786	60.391	1.00	52.93
4405	CG SD	MET	466 466	97.464	26.797	60.618	1.00 1.00	49.91
4406 4407	CE	MET MET	466	97.890 97.679	27.866 29.434	61.980 61.228	1.00	56.57 45.63
4408	H	MET	466	96.483	27.193	57.936	1.00	25.00
4409	N	ARG	467	93.303	26.744	58.916	1.00	57.72
4410	CA	ARG	467	91.849	26.704	58.921	1.00	60.99
4411	C	ARG	467	91.271	25.700	57.935	1.00	57.48
4412	O	ARG	467	90.406	24.901	58.295	1.00	61.64
4413	CB	ARG	467	91.270	28.089	58.642	1.00	69.59
4414	CG	ARG	467	91.304	29.011	59.839	1.00	84.26
4415	CD	ARG	467	90.397	30.202	59.616	1.00	96.57
4416	NE CZ	ARG	467	90.163	30.938	60.853	1.00	103.82
4417 4418	CZ NH1	ARG ARG	467 467	89.074 88.109	31.660 31.749	61.099 60.192	$\frac{1.00}{1.00}$	107.61 108.04
4419	NH2	ARG	467	88.946	32.284	62.261	1.00	109.38
4420	H	ARG	467	93.769	27.556	58.623	1.00	25.00
4421	HE	ARG	467	90.848	30.890	61.554	1.00	25.00
4422	1HH1	ARG	467	88.194	31.273	59.318	1.00	25.00
4423	2HH1	ARG	467	87.293	32.293	60.390	1.00	25.00
4424	1HH2	ARG	467	89.669	32.209	62.950	1.00	25.00
4425	2HH2	ARG	467	88.130	32.827	62.457	1.00	25.00
4426	N	ASP	468	91.789	25.726	56.704	1.00	52.60
4427	CA	ASP	468	91.287	24.832	55.660	1.00	51.80
4428	С	ASP	468 468	91.404	23.354	56.032	1.00	52.54 57.40
4429 4430	O CB	ASP ASP	468 468	90.488 92.026	22.574 25.111	55.767 54.346	1.00 $1.00$	57.49 49.09
4430	СБ	ASP	468	92.020	24.513	53.133	1.00	52.94
4432	OD1	ASP	468	90.127	24.170	53.218	1.00	55.18
4433	OD2	ASP	468	91.983	24.402	52.076	1.00	54.77
4434	H	ASP	468	92.480	26.362	56.496	1.00	25.00
4435	N	TYR	469	92.524	22.973	56.646	1.00	52.44
4436	CA	TYR	469	92.755	21.581	57.040	1.00	49.37
4437	С	TYR	469	92.458	21.283	58.511	1.00	48.25
4438	О	TYR	469	92.316	20.121	58.894	1.00	46.98

TABLE 11-continued

	Struc	In the	e Absence of			ene Synth	ase	
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
4439	CB	TYR	469	94.200	21.165	56.730	1.00	46.99
4440 4441	CG CD1	TYR	469	94.546 94.994	21.129 22.273	55.260	1.00	47.76
4441	CD1 CD2	TYR TYR	469 469	94.431	19.947	54.602 54.524	$\frac{1.00}{1.00}$	51.80 50.13
4443	CE1	TYR	469	95.317	22.246	53.248	1.00	56.40
4444	CE2	TYR	469	94.753	19.908	53.167	1.00	54.09
4445	CZ	TYR	469	95.195	21.063	52.537	1.00	57.04
4446	OH	TYR	469	95.514	21.044	51.198	1.00	62.08
4447	H	TYR	469	93.216	23.642	56.837	1.00	25.00
4448 4449	HH N	TYR GLY	469 470	95.802 92.371	21.916 22.328	50.922 59.328	1.00 $1.00$	25.00 49.37
4450	CA	GLY	470	92.371	22.147	60.746	1.00	51.47
4451	C	GLY	470	93.332	21.551	61.425	1.00	52.90
4452	O	GLY	470	93.247	20.499	62.064	1.00	54.39
4453	Н	GLY	470	92.471	23.228	58.968	1.00	25.00
4454	N	ILE	471	94.467	22.238	61.300	1.00	53.10
4455	CA	ILE	471	95.728	21.771	61.874	1.00	48.51
4456 4457	C O	ILE ILE	471 471	96.521 96.230	22.877 24.063	62.571 62.408	$\frac{1.00}{1.00}$	49.16 48.02
4458	СВ	ILE	471	96.617	21.131	60.781	1.00	44.19
4459	CG1	ILE	471	96.816	22.116	59.621	1.00	43.37
4460	CG2	ILE	471	95.991	19.829	60.288	1.00	41.20
4461	CD1	ILE	471	97.608	21.558	58.449	1.00	38.24
4462	H	ILE	471	94.452	23.094	60.822	1.00	25.00
4463	N	SER	472	97.517	22.475	63.357	1.00	53.02
4464	CA	SER	472	98.371	23.414	64.085	1.00	57.32
4465 4466	C O	SER SER	472 472	99.352 99.689	24.117	63.146 62.084	$\frac{1.00}{1.00}$	60.76 61.94
4467	СВ	SER	472	99.089	23.587 22.672	65.180	1.00	59.87
4468	OG	SER	472	99.873	21.568	64.653	1.00	59.66
4469	Н	SER	472	97.697	21.518	63.453	1.00	25.00
4470	HG	SER	472	100.314	21.103	65.376	1.00	25.00
4471	N	THR	473	99.838	25.287	63.557	1.00	61.86
4472	CA	THR	473	100.794	26.053	62.755	1.00	63.28
4473	С	THR	473	101.959	25.160	62.340	1.00	66.44
4474 4475	O CB	THR THR	473 473	102.374 101.366	25.158 27.248	61.179 63.547	1.00 1.00	66.96 62.70
4476	OG1	THR	473	101.300	28.083	63.997	1.00	63.68
4477	CG2	THR	473	102.306	28.068	62.677	1.00	63.27
4478	H	THR	473	99.529	25.654	64.404	1.00	25.00
4479	HG1	THR	473	99.682	27.608	64.552	1.00	25.00
4480	N	LYS	474	102.454	24.380	63.296	1.00	65.27
4481	CA	LYS	474	103.568	23.470	63.065	1.00	67.87
4482 4483	C O	LYS LYS	474 474	103.248 104.051	22.508 22.325	61.922 61.001	$\frac{1.00}{1.00}$	65.57 66.11
4484	СВ	LYS	474	103.863	22.686	64.349	1.00	71.33
4485	CG	LYS	474	105.150	21.875	64.320	1.00	77.07
4486	CD	LYS	474	105.422	21.234	65.673	1.00	77.42
4487	CE	LYS	474	106.776	20.544	65.698	1.00	78.55
4488	NZ	LYS	474	107.067	19.962	67.037	1.00	75.57
4489	H	LYS	474	102.058	24.422	64.186	1.00	25.00
4490 4491	1HZ 2HZ	LYS LYS	474 474	106.336 107.998	19.263 19.500	67.278 67.020	$\frac{1.00}{1.00}$	25.00 25.00
4492	3HZ	LYS	474	107.070	20.720	67.750	1.00	25.00
4493	N	GLU	475	102.047	21.944	61.960	1.00	61.44
4494	CA	GLU	475	101.612	20.998	60.945	1.00	57.77
4495	C	GLU	475	101.378	21.683	59.599	1.00	53.01
4496	O	GLU	475	101.623	21.091	58.545	1.00	55.03
4497	CB	GLU	475	100.352	20.282	61.418	1.00	59.71
4498 4499	CG	GLU	475 475	100.104	18.950	60.737	1.00	73.81
4500	CD OE1	GLU GLU	475 475	98.994 98.562	18.148 18.513	61.399 62.518	$\frac{1.00}{1.00}$	84.47 88.32
4501	OE2	GLU	475	98.555	17.144	60.797	1.00	88.90
4502	H	GLU	475	101.423	22.181	62.675	1.00	25.00
4503	N	ALA	476	100.931	22.936	59.637	1.00	46.25
4504	CA	ALA	476	100.681	23.703	58.420	1.00	43.91
4505	С	ALA	476	102.003	23.972	57.712	1.00	44.05
4506 4507	O CP	ALA	476 476	102.124	23.774	56.501	1.00	42.50
4507 4508	CB H	ALA ALA	476 4776	99.984 100.761	25.018 23.358	58.749 60.501	$\frac{1.00}{1.00}$	34.16 25.00
4508 4509	н N	MET	477	103.000	24.402	58.480	1.00	45.42
4510	CA	MET	477	104.321	24.689	57.932	1.00	46.57
4511	С	MET	477	104.954	23.414	57.395	1.00	46.73
4512	O	MET	477	105.640	23.434	56.369	1.00	48.81

TABLE 11-continued

	In the Absence of Bound Substrate							
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
4513	CB	MET	477	105.217	25.331	58.990	1.00	41.20
4514	CG	MET	477	104.699	26.674	59.459	1.00	45.53
4515	SD	MET	477	105.842	27.539	60.529	1.00	50.24
4516	CE	MET	477	105.403	29.229	60.204	1.00	44.87
4517	H	MET	477	102.838	24.531	59.436	1.00	25.00
4518	N	ALA	478	104.689	22.301	58.071	1.00	43.88
4519	CA	ALA	478	105.214	21.012	57.646	1.00	40.24
4520	C	ALA	473	104.608	20.664	56.288	1.00	39.48
4521	O	ALA	478	105.301	20.158	55.404	1.00	42.61
4522	СВ	ALA	478	104.887	19.941	58.673	1.00	38.44
4523	H	ALA	473	104.140	22.350	58.883	1.00	25.00
4524 4525	N CA	LYS	479 479	103.324	20.969	56.113	1.00	37.41
4525 4526	CA	LYS LYS	479 479	102.642 103.214	20.700 21.598	54.850 53.754	1.00 1.00	36.91 33.62
4527	C O	LYS	479 479	103.214	21.398	52.616	1.00	32.37
4528	СВ	LYS	479	101.136	20.931	54.986	1.00	39.38
4529	CG	LYS	479	100.338	20.573	53.736	1.00	46.00
4530	CD	LYS	479	98.850	20.797	53.947	1.00	51.63
4531	CE	LYS	479	98.273	19.858	55.003	1.00	53.61
4532	NZ	LYS	479	98.180	18.451	54.525	1.00	57.01
4533	H	LYS	479	102.820	21.3777	56.852	1.00	25.00
4534	1HZ	LYS	479	99.126	18.106	54.269	1.00	25.00
4535	2HZ	LYS	479	97.776	17.853	55.274	1.00	25.00
4536	3HZ	LYS	479	97.561	18.416	53.688	1.00	25.00
4537	N	PHE	480	103.502	22.845	54.107	1.00	30.92
4538	CA	PHE	480	104.067	23.790	53.157	1.00	31.88
4539	C	PHE	480	105.457	23.356	52.714	1.00	34.12
4540	O	PHE	480	105.812	23.493	51.540	1.00	37.15
4541	CB	PHE	480	104.107	25.198	53.749	1.00	29.35
4542	CG	PHE	480	102.902	26.028	53.408	1.00	37.28
4543	CD1	PHE	480	101.662	25.427	53.190	1.00	37.60
4544	CD2	PHE	480	103.008	27.411	53.283	1.00	36.73
4545	CE1	PHE	480	100.540	26.192	52.850	1.00	36.20
4546	CE22	PHE	480	101.898	28.185	52.942	1.00	38.45
4547	CZ	PHE	480	100.665	27.574	52.726	1.00	36.96
4548 4549	H N	PHE	480 481	103.315	23.138	55.024 53.641	1.00 1.00	25.00
4549 4550	CA	GLN GLN	481	106.238 107.573	22.810 22.352	53.292	1.00	33.79 35.43
4551	CA	GLN	481	107.373	21.180	52.323	1.00	35.55
4552	o	GLN	481	108.200	21.103	51.347	1.00	35.63
4553	СВ	GLN	481	108.368	21.930	54.524	1.00	46.60
4554	CG	GLN	481	109.844	21.688	54.210	1.00	70.92
4555	CD	GLN	481	110.583	20.933	55.302	1.00	83.38
4556	OE1	GLN	481	110.036	20.658	56.371	1.00	93.92
4557	NE2	GLN	481	111.836	20.588	55.032	1.00	88.62
4558	H	GLN	481	105.919	22.723	54.562	1.00	25.00
4559	1HE2	GLN	481	112.316	20.105	55.735	1.00	25.00
4560	2HE2	GLN	481	112.220	20.824	54.166	1.00	25.00
4561	N	ASN	482	106.486	20.297	52.561	1.00	33.51
4562	CA	ASN	482	106.272	19.146	51.682	1.00	36.28
4563	C	ASN	482	105.950	19.606	50.267	1.00	36.07
4564	O	ASN	482	106.380	18.989	49.288	1.00	35.83
4565	CB	ASN	482	105.140	18.252	52.200	1.00	40.65
4566	CG CD1	ASN	482	105.535	17.465	53.436	1.00	54.20
4567	CD1 ND2	ASN	482 482	106.698 104.565	17.095	53.607 54.307	$\frac{1.00}{1.00}$	57.37 59.54
4568 4569	H	ASN ASN	482	104.303	17.204 20.415	53.350	1.00	25.00
4570	n 1HD2	ASN	482	103.916	16.698	55.105	1.00	25.00
4571	2HD2	ASN	482	103.666		54.121	1.00	25.00
4572	N	MET	483	105.000	20.698	50.163	1.00	33.64
4573	CA	MET	483	104.831	21.250	48.866	1.00	29.91
4574	C	MET	483	106.080	21.757	48.152	1.00	27.30
4575	Ö	MET	433	106.240	2[].556	46.947	1.00	32.31
4576	СВ	MET	483	103.814	22.373	49.036	1.00	29.58
4577	CG	MET	483	102.488	21.916	49.626	1.00	32.65
4578	SD	MET	483	101.388	23.311	49.943	1.00	37.42
4579	CE	MET	483	100.988	23.770	48.275	1.00	33.87
4580	H	MET	483	104.879	21.134	50.982	1.00	25.00
4581	N	ALA	484	106.979	22.385	48.903	1.00	24.24
4582	CA	ALA	484	108.226	22.895	48.339	1.00	25.37
4583	С	ALA	484	109.086	21.724	47.845	1.00	27.23
4584	O	ALA	484	109.696	21.792	46.772	1.00	26.13
4585 4596	CB	ALA	484	108.979	23.703	49.385	1.00	20.73
4586	Н	ALA	484	106.799	22.516	49.859	1.00	26.00

TABLE 11-continued

		In the	e Absence o	f Bound S	ubstrate			
Atom Typ	e Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
4587	N	GLU	485	109.103	20.642	48.622	1.00	27.25
4588	CA	GLU	485	109.864	19.437	48.289	1.00	29.59
4589	С	GLU	485	109.317	18.807	47.015	1.00	26.53
4590	O	GLU	485	110.070	18.386	46.139	1.00	32.01
4591	CB	GLU	485	109.792	18.425	49.437	1.00	39.77
4592	CG	GLU	485	110.327	18.942	50.779	1.00	61.11
4593	CD	GLU	485	110.170	17.953	51.934	1.00	69.27
4594	OE1	GLU	485	109.663	16.830	51.716	1.00	73.83
4595	OE2	GLU	485	110.561	18.302	53.073	1.00	69.26
4596	H	GLU	485	108.574	20.683	49.449	1.00	25.00
4597	N	THR	486	107.997	18.755	46.917	1.00	26.56
4598 4599	CA C	THR	486 486	107.323 107.673	18.207	45.749 44.520	1.00 1.00	26.51 24.78
4600	o	THR THR	486	107.073	19.050 18.514	43.437	1.00	24.70
4601	СВ	THR	486	107.901	18.211	45.973	1.00	30.57
4602	OG1	THR	486	105.463	17.277	47.010	1.00	30.11
4603	CG2	THR	486	105.403	17.855	44.694	1.00	27.79
4604	H	THR	486	107.446	19.085	47.660	1.00	25.00
4605	HG1	THR	486	105.755	16.396	46.782	1.00	25.00
4606	N	ALA	487	107.672	20.367	44.712	1.00	20.69
4607	CA	ALA	487	107.980	21.319	43.651	1.00	19.32
4608	C	ALA	487	109.409	21.127	43.141	1.00	18.49
4609	Ō	ALA	487	109.654	21.149	41.929	1.00	15.86
4610	СВ	ALA	487	107.768	22.750	44.152	1.00	15.08
4611	H	ALA	487	107.454	20.716	45.604	1.00	25.00
4612	N	TRP	488	110.349	20.909	44.058	1.00	20.11
4613	CA	TRP	488	111.736	20.695	43.661	1.00	19.04
4614	С	TRP	488	111.856	19.461	42.781	1.00	20.42
4615	O	TRP	488	112.555	19.486	41.768	1.00	23.27
4616	CB	TRP	488	112.656	20.590	44.879	1.00	19.13
4617	CG	TRP	488	113.256	21.905	45.262	1.00	20.79
4618	CD1	TRP	488	113.017	22.619	46.402	1.00	19.35
4619	CD2	TRP	488	114.173	22.689	44.481	1.00	18.53
4620	NE1	TRP	488	113.723	23.801	46.376	1.00	21.34
4621	CE2	TRP	488	114.441	23.869	45.210	1.00	17.70
4622	CE3	TRP	488	114.793	22.507	43.237	1.00	18.03
4623	CZ2	TRP	488	115.305	24.863	44.736	1.00	16.71
4624	CZ3	TRP	488	115.654	23.499	42.765	1.00	15.24
4625	CH2	TRP	488 488	115.899	24.659 20.905	43.515	1.00	14.14
4626 4627	H HE1	TRP TRP	488	110.109 113.699	24.482	45.009 47.075	1.00 1.00	25.00 25.00
4628	nei N	LYS	489	111.136	24.462 18.399	43.138	1.00	23.00
4629	CA	LYS	489	111.150	17.175	42.345	1.00	17.89
4630	C	LYS	489	110.604	17.476	40.961	1.00	19.66
4631	Ö	LYS	489	111.091	16.947	39.960	1.00	22.89
4632	СВ	LYS	489	110.351	16.069	43.019	1.00	17.77
4633	CG	LYS	489	110.922	15.624	44.344	1.00	15.98
4634	CD	LYS	489	110.074	14.540	44.972	1.00	22.31
4635	CE	LYS	489	110.525	14.254	46.392	1.00	24.45
4636	NZ	LYS	489	109.694	13.199	47.029	1.00	26.46
4637	H	LYS	489	110.589	18.443	43.952	1.00	25.00
4638	1HZ	LYS	489	108.703	13.510	47.052	1.00	25.00
4639	2HZ	LYS	489	109.772	12.320	46.482	1.00	25.00
4640	3HZ	LYS	489	110.028	13.037	47.997	1.00	25.00
4641	N	ASP	490	109.590	18.338	40.906	1.00	20.77
4642	CA	ASP	490	108.991	18.721	39.630	1.00	21.97
4643	С	ASP	490	110.008	19.479	38.786	1.00	24.19
4644	O	ASP	490	110.098	19.264	37.575	1.00	21.17
4645	CB	ASP	490	101.739	19.585	39.837	1.00	26.18
4646	CG	ASP	490	106.561	18.799	40.395	1.00	29.44
4647	OD1	ASSP	490	106.524	17.562	40.236	1.00	35.64
4648	OD2	ASP	490	105.657	19.425	40.982	1.00	29.87
4649	H	ASP	490	109.245	18.731	41.738	1.00	25.00
4650 4651	N CA	ILE ILE	491 491	110.776	20.362	39.419 38.692	$\frac{1.00}{1.00}$	20.68 20.37
4651 4652	CA	ILE	491 491	111.789 112.810	21.120 20.148	38.092	1.00	19.25
4653	Ö	ILE	491	113.158	20.148	36.934	1.00	20.39
4654	СВ	ILE	491	112.508	22.153	39.595	1.00	22.00
4655	CG1	ILE	491	111.540	23.278	39.975	1.00	21.39
4656	CG2	ILE	491	113.737	22.716	38.877	1.00	21.00
4657	CD1	ILE	491	112.159	24.376	40.803	1.00	22.99
4658	Н	ILE	491	110.657	20.508	40.379	1.00	25.00
4659	N	ASN	492	113.239	19.198	38.943	1.00	19.07
4660	CA	ASN	492	114.216	18.196	38.529	1.00	18.67

TABLE 11-continued

		In th	e Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
4661	C	ASN	492	113.700	17.404	37.332	1.00	20.34
4662	O	ASN	492	114.446	17.133	36.393	1.00	20.40
4663	CB	ASN	492	114.567	17.271	39.699	1.00	16.62
4664	CG	ASN	492	115.269	18.007	40.839	1.00	18.56
4665	OD1	ASN	492	115.924	19.035	40.625	1.00	15.51
4666	ND2	ASN	492	115.140	17.484	42.050	1.00	15.73
4667	H	ASN	492	112.899	19.181	39.862	1.00	25.00
4668	1HD2	ASN	492	115.583	17.939	42.793	1.00	25.00
4669	2HD2	ASN	492	114.613	16.661	42.164	1.00	25.00
4670	N	GLU	493	112.412	17.073	37.341	1.00	21.12
4671	CA	GLU	493	111.816	16.341	36.225	1.00	22.19
4672 4673	C O	GLU GLU	493 493	111.736 111.958	17.225 16.755	34.985 33.869	1.00 1.00	24.12 26.10
4674	СВ	GLU	493	111.936	15.850	36.578	1.00	19.71
4675	CG	GLU	493	110.410	14.831	37.690	1.00	30.24
4676	CD	GLU	493	109.056	14.143	37.849	1.00	25.61
4677	OE1	GLU	493	109.030	14.460	37.100	1.00	36.10
4678	OE2	GLU	493	108.953	13.268	38.728	1.00	35.77
4679	H	GLU	493	111.859	17.328	38.111	1.00	25.00
4680	N	GLY	494	111.423	18.504	35.194	1.00	24.81
4681	CA	GLY	494	111.311	19.451	34.096	1.00	18.06
4682	C	GLY	494	112.614	19.686	33.352	1.00	25.75
4683	O	GLY	494	112.605	20.176	32.217	1.00	25.47
4684	H	GLY	494	111.263	18.819	36.107	1.00	25.00
4685	N	LEU	495	113.735	19.350	33.986	1.00	24.09
4686	CA	LEU	495	115.047	19.523	33.367	1.00	23.57
4687	C	LEU	495	115.465	18.331	32.503	1.00	23.66
4688	O	LEU	495	116.385	18.445	31.700	1.00	25.21
4689	CB	LEU	495	116.111	19.781	34.439	1.00	21.29
4690	CG	LEU	495	115.968	21.063	35.270	1.00	24.69
4691	CD1	LEU	495	116.913	21.024	36.459	1.00	15.49
4692	CD2	LEU	495	116.230	22.287	34.409	1.00	21.41
4693	H	LEU	495	113.681	18.985	34.893	1.00	25.00
4694	N	LEU	496	114.781	17.200	32.651	1.00	22.59
4695	CA	LEU	496	115.118	15.996	31.889	1.00	20.47
4696	C	LEU	496	114.749	16.049	30.409	1.00	24.48
4697	O	LEU	496	113.692	16.556	30.033	1.00	22.73
4698	CB	LEU	496	114.504	14.758	32.548	1.00	20.18
4699 4700	CG CD1	LEU	496 496	115.016	14.454	33.959 34.524	1.00 1.00	23.38
4700	CD1 CD2	LEU LEU	496 496	114.276 116.523	13.265 14.187	33.938	1.00	21.31 20.33
4701	H	LEU	496 496	114.020	17.172	33.267	1.00	25.00
4702	N	ARG	497	115.642	15.530	29.573	1.00	26.43
4704	CA	ARG	497	115.443	15.501	28.128	1.00	31.12
4705	C	ARG	497	114.347	14.498	27.766	1.00	32.68
4706	Ö	ARG	497	114.217	13.457	28.411	1.00	27.55
4707	CB	ARG	497	116.757	15.124	27.431	1.00	30.06
4708	CG	ARG	497	117.883	16.155	27.626	1.00	38.94
4709	CD	ARG	497	119.217	15.505	27.851	1.00	37.77
4710	NE	ARG	497	120.087	15.584	26.683	1.00	50.61
4711	CZ	ARG	497	121.282	16.173	26.676	1.00	51.72
4712	NH1	ARG	497	121.754	16.744	27.777	1.00	46.52
4713	NH2	ARG	497	122.023	16.166	25.575	1.00	51.65
4714	H	ARG	497	116.457	15.140	29.946	1.00	25.00
4715	HE	ARG	497	119.773	15.180	25.847	1.00	25.00
4716	1HH1	ARG	497	121.213	16.733	28.615	1.00	25.00
4717	2HH1	ARG	497	122.653	17.183	27.766	1.00	25.00
4718	1HH2	ARG	497	121.685	15.718	24.748	1.00	25.00
4719	2HH2	ARG	497	122.920	16.608	25.576	1.00	25.00
4720	N	PRO	498	113.542	14.798	26.731	1.00	34.48
4721	CA	PRO	498	113.595	16.005	25.897	1.00	31.05
4722	С	PRO	498	112.886	17.179	26.568	1.00	32.44
4723	O	PRO	498	111.757	17.040	27.043	1.00	32.35
4724 4725	CB CG	PRO PRO	498 498	112.831 112.768	15.587 14.079	24.635 24.707	$\frac{1.00}{1.00}$	30.63 38.95
4725 4726	CD	PRO	498 498	112.768	13.830	24.707	1.00	38.95 34.35
4727	N N	THR	498 499	112.595	18.332	26.612	1.00	33.10
4727	CA	THR	499	112.940	19.513	27.218	1.00	30.18
4729	CA	THR	499	112.940	20.223	26.170	1.00	31.00
4730	Ö	THR	499	112.369	20.172	24.974	1.00	33.56
4731	CB	THR	499	114.016	20.474	27.795	1.00	25.39
4732	OG1	THR	499	115.004	20.752	26.798	1.00	24.84
4733	CG2	THR	499	114.703	19.843	28.996	1.00	22.98
4734	H	THR	499	114.435	18.421	26.223	1.00	25.00

TABLE 11-continued

	In the Absence of Bound Substrate							
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
4735	HG1	THR	499	115.646	21.357	27.117	1.00	25.00
4736	N	PRO	500	110.963	20.844	26.600	1.00	31.06
4737 4738	CA C	PRO PRO	500 500	110.053 110.705	21.558 22.740	25.692 24.967	$\frac{1.00}{1.00}$	32.31 34.94
4738 4739	0	PRO	500	110.703	23.075	23.843	1.00	39.31
4740	СВ	PRO	500	108.916	21.994	26.620	1.00	29.71
4741	CG	PRO	500	109.576	22.086	27.968	1.00	28.06
4742	CD	PRO	500	110.460	20.876	27.984	1.00	23.70
4743	N	VAL	501	111.642	23.398	25.641	1.00	32.94
4744	CA	VAL	501	112.390	24.523	25.078	1.00	33.22
4745 4746	C O	VAL VAL	501 501	113.858 114.154	24.257 23.347	25.421 26.204	1.00 1.00	33.52 33.00
4747	СВ	VAL	501	111.959	25.887	25.686	1.00	29.14
4748	CG1	VAL	501	110.515	26.198	25.330	1.00	26.33
4749	CG2	VAL	501	112.153	25.887	27.195	1.00	24.60
4750	H	VAL	501	111.875	23.112	26.540	1.00	25.00
4751	N	SER	502	114.775	25.026	24.844	1.00	30.56
4752 4753	CA C	SER SER	502 502	116.194 116.485	24.832 25.025	25.128 26.611	$\frac{1.00}{1.00}$	33.20 32.05
4754	Ö	SER	502	115.869	25.869	27.265	1.00	34.57
4755	СВ	SER	502	117.039	25.807	24.316	1.00	35.54
4756	OG	SER	502	116.837	25.601	22.934	1.00	56.52
4757	H	SER	502	114.502	25.729	24.224	1.00	25.00
4758	HG	SER	502	117.098	24.709	22.686	1.00	25.00
4759	N	THR	503	117.443	24.260	27.126	1.00	29.15
4760 4761	CA C	THR THR	503 503	117.836 118.166	24.333 25.771	28.530 28.927	$\frac{1.00}{1.00}$	33.23 31.90
4762	ŏ	THR	503	117.977	26.177	30.078	1.00	32.13
4763	CB	THR	503	119.058	23.443	28.797	1.00	38.70
4764	OG1	THR	503	118.767	22.110	28.366	1.00	51.69
4765	CG2	THR	503	119.395	23.420	30.278	1.00	40.66
4766	H HG1	THR	503 503	117.884	23.611 22.075	26.547	1.00 $1.00$	25.00 25.00
4767 4768	N	THR GLU	504	118.560 118.637	26.542	27.436 27.956	1.00	27.88
4769	CA	GLU	504	118.982	27.935	28.184	1.00	31.30
4770	C	GLU	504	117.801	28.706	28.789	1.00	31.46
4771	O	GLU	504	117.987	29.643	29.568	1.00	29.72
4772	CB	GLU	504	119.396	28.578	26.863	1.00	32.18
4773	CG	GLU	504	119.754	30.042	26.997	1.00	44.47
4774 4775	CD OE1	GLU GLU	504 504	120.045 119.634	30.714 30.183	25.672 24.618	1.00 1.00	47.35 49.18
4776	OE2	GLU	504	120.683	31.788	25.691	1.00	48.31
4777	Н	GLU	504	118.771	26.169	27.067	1.00	25.00
4778	N	PHE	505	116.588	28.274	28.464	1.00	27.10
4779	CA	PHE	505	115.390	28.936	28.957	1.00	23.02
4780	C	PHE	505	114.809	28.314	30.218	1.00	24.14
4781 4782	O CS	PHE PHE	505 505	113.888 114.356	28.869 29.036	30.818 27.335	$\frac{1.00}{1.00}$	22.77 28.22
4783	CG	PHE	505	114.888	29.711	26.602	1.00	28.58
4784	CD1	PHE	505	115.307	31.039	26.651	1.00	28.23
4785	CD2	PHE	505	115.048	29.001	25.417	1.00	28.81
4786	CE1	PHE	505	115.884	31.646	25.539	1.00	25.11
4787	CE2	PHE	505	115.623	29.597	24.300	1.00	28.43
4788	CZ	PHE	505	118.043 116.489	30.922	24.362 27.881	1.00	29.72 25.00
4789 4790	H N	PHE LEU	505 506	115.367	27.504 27.182	30.641	$\frac{1.00}{1.00}$	25.00
4791	CA	LEU	606	114.915	26.516	31.862	1.00	20.00
4792	C	LEU	506	115.763	26.980	33.054	1.00	20.28
4793	O	LEU	506	115.270	27.120	34.176	1.00	21.38
4794	СВ	LEU	506	115.033	24.995	31.732	1.00	17.84
4795	CG CD1	LEU	506	114.265	24.277	30.621	1.00	23.70
4796 4797	CD1 CD2	LEU LEU	506 506	114.409 112.797	22.781 24.671	30.832 30.645	1.00 $1.00$	19.27 20.22
4798	H	LEU	506	116.092	26.791	30.120	1.00	25.00
4799	N	THR	507	117.040	27.237	32.796	1.00	24.00
4800	CA	THR	507	117.968	27.666	33.837	1.00	21.87
4801	С	THR	507	117.508	28.894	34.634	1.00	21.92
4802	O	THR	507	117.636	28.913	35.858	1.00	25.77
4803 4804	CB OG1	THR	507	119.382	27.870	33.260	1.00	22.57
4804 4805	OG1 CG2	THR THR	507 507	119.728 120.400	26.728 28.014	32.405 34.381	$\frac{1.00}{1.00}$	23.76 20.42
4806	H	THR	507	117.372	27.127	31.883	1.00	25.00
4807	HG1	THR	507	119.106	26.824	31.739	1.00	25.00
4808	N	PRO	508	116.960	29.928	33.963	1.00	17.26

TABLE 11-continued

		In the	Absence o	f Bound Subs	trate		
Atom Type	Atom	Residue	Residue #	X	Y Z	occ	B-factor
4809	CA	PRO	508	116.503 31	.114 34.69	98 1.00	18.03
4810	C	PRO	508	115.423 30			19.69
4811	O	PRO	508	1155.417 31			22.29
4812	CB	PRO	508	115.943 31			18.28
4813	CG	PRO	508	116.841 31			16.25
4814	CD	PRO	508	118.909 30			14.75
4815 4816	N CA	ILE ILE	509 509	114.528 29 113.435 29			21.35 22.73
4817	C	ILE	509	114.024 28			19.96
4818	Ö	ILE	509	113.627 28			22.40
4819	СВ	ILE	509	112.450 28			18.26
4820	CG1	ILE	509	111.867 29			20.40
4821	CG2	ILE	509	111.360 27	.998 36.3	51 1.00	12.67
4822	CD1	ILE	509	111.082 28			22.89
4823	H	ILE	509	114.597 29			25.00
4824	N	LEU	510	114.989 27			21.34
4825	CA	LEU	510	115.684 27			19.45
4826	С	LEU	510	116.420 27			18.03
4827 4828	O CB	LEU LEU	510 510	116.372 27 116.693 26			23.54
4829	CG	LEU	510	117.747 25			18.99 17.88
4830	CD1	LEU	510	117.092 24			13.70
4831	CD2	LEU	510	118.660 24			12.82
4832	Н	LEU	510	115.234 27			25.00
4833	N	ASN	511	117.070 28			16.84
4834	CA	ASN	511	117.816 29	.940 39.29	93 1.00	18.52
4835	C	ASN	511	116.918 30	.774 40.18	35 1.00	19.49
4836	O	ASN	511	117.299 31			21.10
4837	CB	ASN	511	118.704 30			14.99
4838	CG	ASN	511	119.926 30			19.61
4839	OD1	ASN	511	120.276 29			22.94
4840	ND2	ASN	511	120.562 30			17.54
4841 4842	H 1HD2	ASN ASN	511 511	117.040 29 121.341 30			25.00 25.00
4843	2HD2	ASN	511	120.230 31			25.00
4844	N	LEU	512	115.705 31			18.46
4845	CA	LEU	512	114.751 31			15.00
4846	C	LEU	512	114.415 31			16.15
4847	O	LEU	512	114.304 31	.561 42.8	72 1.00	24.00
4848	CB	LEU	512	113.484 32	.114 39.72	27 1.00	17.19
4849	CG	LEU	5112	113.569 33			16.79
4850	CD1	LEU	512	112.331 33			22.39
4851	CD2	LEU	512	113.702 34			14.37
4852	H	LEU	512	115.450 30			25.00
4853	N CA	ALA	513	114.279 29			19.31
4854 4855	CA C	ALA ALA	513 513	113.979 28 115.178 28			18.01 18.19
4856	Ö	ALA	513	115.176 28			16.19
4857	СВ	ALA	513	113.654 27			12.64
4858	H	ALA	513	114.362 29			25.00
4859	Ñ	ARG	514	116.381 28			19.70
4860	CA	ARG	514	117.609 28			18.80
4861	C	ARG	514	117.696 30		34 1.00	17.46
4862	O	ARG	514	118.041 29			21.48
4863	CB	ARG	514	118.832 28			13.68
4864	CG	ARG	514	118.981 27			14.32
4865	CD	ARG	514	120.084 27			17.33
4866	NE	ARG	514	120.490 25			16.97
4867 4868	CZ NH1	ARG ARG	514 514	121.107 25 121.398 26			19.03 15.08
4869	NH2	ARG	514	121.398 20			15.36
4870	H	ARG	514	116.440 28			25.00
4871	HE	ARG	514	120.282 25			25.00
4872	1HH1	ARG	514	121.159 27			25.00
4873	2HH1	ARG	514	121.862 26			25.00
4874	1HH2	ARG	514	121.245 23			25.00
4875	2HH2	ARG	514	121.913 23			25.00
4876	N	ILE	515	117.330 31			20.52
4877	CA	ILE	515	117.352 32			22.89
4878	С	ILE	515	116.489 32			25.55
4879	O CP	ILE	515 515	116.851 32			28.26
4880 4881	CB CG1	ILE ILE	515 515	116.863 33			18.86
4881 4882	CG2	ILE	515 515	117.857 33 116.695 34			18.35 23.30
1002	002	·	515	110.000 07	17.0.	1.00	25.50

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
4883	CD1	ILE	515	117.408	34.863	41.885	1.00	19.17
4884	H	ILE	515	117.045	31.141	43.257	1.00	25.00
4885	N	VAL	516	115.372	31.637	46.082	1.00	27.24
4886	CA	VAL	516	114.467	31.463	47.220	1.00	26.85
4887	С	VAL	516	115.229	30.838	48.378	1.00	30.63
4888	O	VAL	516	115.219	31.354	49.496	1.00	28.87
4889	CB	VAL	516	113.280	30.512	46.881	1.00	28.84
4890 4891	CG1 CG2	VAL VAL	516 516	112.433 112.423	30.248 31.096	48.122 45.776	$\frac{1.00}{1.00}$	23.42 22.08
4892	H	VAL	516	115.144	31.219	45.225	1.00	25.00
4893	N	GLU	517	115.910	29.736	48.085	1.00	35.61
4894	CA	GLU	517	116.680	28.997	49.081	1.00	41.24
4895	C	GLU	517	117.696	29.890	49.796	1.00	42.08
4896	O	GLU	517	117.872	29.789	51.009	1.00	46.37
4897	CB	GLU	517	117.385	27.802	48.424	1.00	41.58
4898	CG	GLU	517	116.496	26.950	47.503	1.00	52.96
4899	CD	GLU	517	115.344	26.242	48.223	1.00	59.58
4900	OE1	GLU	517	115.593	25.557	49.236	1.00	60.41
4901	OE2	GLU	517	114.187	26.352	47.762	1.00	63.04
4902	H	GLU	517	115.899	29.407	47.161	1.00	25.00
4903 4904	N CA	VAL	518 518	118.314	30.799 31.714	49.050 49.600	1.00	40.18 42.32
4904	CA	VAL VAL	518	119.310 118.704	32.885	50.386	$\frac{1.00}{1.00}$	42.32 47.45
4906	ŏ	VAL	518	119.269	33.326	51.389	1.00	48.49
4907	СВ	VAL	518	120.219	32.251	48.474	1.00	41.14
4908	CG1	VAL	518	121.133	33.350	48.986	1.00	39.83
4909	CG2	VAL	518	121.034	31.108	47.896	1.00	45.30
4910	H	VAL	518	118.097	30.844	48.095	1.00	25.00
4911	N	THR	519	117.563	33.387	49.923	1.00	45.49
4912	CA	THR	519	116.899	34.505	50.577	1.00	44.49
4913	C	THR	519	116.183	34.117	51.884	1.00	44.03
4914	O	THR	519	115.983	34.964	52.754	1.00	41.48
4915	CB	THR	519	115.868	35.165	49.622	1.00	45.40
4916	OG1	THR	519	116.518	35.516	48.394	1.00	47.45
4917 4918	CG2 H	THR THR	519 519	115.283 117.161	36.424 32.996	50.240 49.123	$\frac{1.00}{1.00}$	50.81 25.00
4919	HG1	THR	519	116.872	34.719	47.980	1.00	25.00
4920	N	TYR	520	115.827	32.843	52.034	1.00	49.74
4921	CA	TYR	520	115.130	32.385	53.240	1.00	54.71
4922	C	TYR	520	115.783	31.213	53.984	1.00	60.58
4923	O	TYR	520	115.129	30.191	54.209	1.00	65.16
4924	CB	TYR	520	113.686	31.988	52.905	1.00	52.58
4925	CG	TYR	520	112.886	33.018	52.142	1.00	53.27
4926	CD1	TYR	520	112.885	33.027	50.748	1.00	54.77
4927	CD2	TYR	520	112.105	33.962	52.809	1.00	50.10
4928	CE1	TYR	520	112.127	33.948	50.032	1.00	57.50
4929	CE2	TYR	520 520	111.340	34.889	52.102	1.00	54.19
4930	CZ	TYR	520 520	111.357	34.873 35.777	50.713 49.999	1.00	56.86 58.70
4931 4932	OH H	TYR TYR	520	110.604 116.029	32.189	51.330	$\frac{1.00}{1.00}$	58.70 25.00
4933	HH	TYR	520	110.726	35.626	49.057	1.00	25.00
4934	N	ILE	521	117.056	31.340	54.350	1.00	67.35
4935	CA	ILE	521	117.729	30.269	55.091	1.00	74.89
4936	С	ILE	521	117.425	30.428	56.583	1.00	75.44
4937	O	ILE	521	117.194	29.397	57.255	1.00	76.20
4938	CB	ILE	521	119.276	30.258	54.856	1.00	75.70
4939	CG1	ILE	521	119.586	29.919	53.394	1.00	76.18
4940	CG2	ILE	521	119.953	29.222	55.766	1.00	77.50
4941	CD1	ILE	521	121.064	29.755	53.080	1.00	71.61
4942	H	ILE	521	117.546	32.156	54.145	1.00	25.00
4943	N	VAL	533	120.428	39.967	55.248	1.00	55.02
4944 4945	CA C	VAL VAL	533 533	120.478 121.277	38.584 38.505	54.683 53.373	$\frac{1.00}{1.00}$	57.02 55.80
4946	o	VAL	533	122.075	37.588	53.181	1.00	56.73
4947	СВ	VAL	533	119.048	37.995	54.485	1.00	56.30
4948	CG1	VAL	533	118.225	38.868	53.539	1.00	56.90
4949	CG2	VAL	533	119.125	36.552	53.986	1.00	50.74
4950	1H	VAL	533	119.970	40.608	54.578	1.00	25.00
4951	2H	VAL	533	119.880	39.938	56.132	1.00	25.00
4952	3H	VAL	533	121.396	40.274	55.462	1.00	25.00
4953	N	LEU	534	121.095	39.483	52.491	1.00	49.61
4954	CA	LEU	534	121.812	39.490	51.218	1.00	48.50
4955	С	LEU	534	123.194	40.138	51.300	1.00	47.52
4956	O	LEU	534	124.075	39.830	50.496	1.00	44.98

TABLE 11-continued

		In th	e Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	z	OCC	B-factor
4957	CB	LEU	534	120.983	40.190	50.137	1.00	48.86
4958	CG	LEU	534	119.659	39.533	49.744	1.00	50.00
4959	CD1	LEU	534	119.054	40.290	48.567	1.00	46.00
4960	CD2	LEU	534	119.886	38.066	49.384	1.00	42.36
4961	H	LEU	534	120.456	40.200	52.652	1.00	25.00
4962	N	LYS	535	123.382	40.993	52.303	1.00	45.93
4963	CA	LYS	535	124.633	41.722	52.510	1.00	45.11
4964	С	LYS	535	125.921	40.923	52.284	1.00	43.08
4965 4966	O CB	LYS LYS	535 535	126.729	41.288	51.428 53.895	1.00 1.00	42.36
4967	CG CG	LYS	535 535	124.651 125.855	42.385 43.288	54.130	1.00	46.19 54.44
4968	CD	LYS	535	125.868	43.868	55.536	1.00	57.76
4969	CE	LYS	535	127.075	44.774	55.747	1.00	61.61
4970	NZ	LYS	535	127.099	45.378	57.111	1.00	62.66
4971	H	LYS	535	122.651	41.156	52.915	1.00	25.00
4972	1HZ	LYS	535	127.134	44.621	57.824	1.00	25.00
4973	2HZ	LYS	535	127.936	45.986	57.207	1.00	25.00
4974	3HZ	LYS	535	126.239	45.946	57.252	1.00	25.00
4975	N	PRO	536	126.115	39.809	53.019	1.00	39.15
4976	CA	PRO	536	127.337	39.020	52.829	1.00	37.51
4977	С	PRO	536	127.564	38.579	51.386	1.00	33.92
4978	O	PRO	536	128.684	38.644	50.877	1.00	33.28
4979	CB CG	PRO	536 536	127.128	37.827 37.724	53.770	1.00 $1.00$	38.16 42.17
4980 4981	CD	PRO PRO	536	125.638 125.233	39.164	53.893 54.008	1.00	38.53
4982	N	HIS	537	126.488	38.181	50.714	1.00	33.98
4983	CA	HIS	537	126.575	37.730	49.327	1.00	34.56
4984	C	HIS	537	126.929	38.877	48.390	1.00	34.75
4985	O	HIS	537	127.742	38.714	47.479	1.00	29.05
4986	CB	HIS	537	125.264	37.071	48.900	1.00	34.41
4987	CG	HIS	537	124.917	35.855	49.703	1.30	41.27
4988	ND1	HIS	537	123.749	35.746	50.426	1.00	43.98
4989	CD2	HIS	537	125.601	34.705	49.917	1.00	37.03
4990	CE1	HIS	537	123.726	34.584	51.053	1.00	40.28
4991 4992	NE2 H	HIS HIS	537 537	124.838 125.618	33.933 38.225	50.760 51.154	1.00 1.00	39.05 25.00
4993	HD1	HIS	537	123.030	36.418	50.483	1.00	25.00
4994	HE2	HIS	537	125.072	33.038	51.088	1.00	25.00
4995	N	ILE	538	126.333	40.040	48.634	1.00	34.95
4996	CA	ILE	538	126.596	41.225	47.829	1.00	35.08
4997	C	ILE	538	128.063	41.612	47.969	1.00	36.46
4998	O	ILE	538	128.703	41.999	46.990	1.00	38.58
4999	CB	ILE	538	125.701	42.406	48.263	1.00	35.99
5000	CG1	ILE	538	124.230	42.072	47.983	1.00	37.36
5001	CG2	ILE	538	126.124	43.681	47.542	1.00	34.76
5002	CD1	ILE	538	123.248	43.112 40.113	48.460 49.385	1.00	36.39
5003 5004	H N	ILE ILE	538 539	125.708 128.588	41.491	49.385	1.00 $1.00$	25.00 34.70
5005	CA	ILE	539	129.979	41.807	49.473	1.00	33.68
5006	C	ILE	539	130.912	40.828	48.769	1.00	34.74
5007	Ö	ILE	539	131.868	41.239	48.093	1.00	30.49
5008	CB	ILE	539	130.253	41.761	51.004	1.00	35.37
5009	CG1	ILE	539	129.559	42.939	51.686	1.00	33.35
5010	CG2	ILE	539	131.749	41.790	51.285	1.00	32.80
5011	CD1	ILE	539	129.684	42.933	53.189	1.00	34.58
5012	H	ILE	539	127.999	41.207	49.913	1.00	25.00
5013	N	ASN	540	130.603	39.538	48.864	1.00	33.09
5014	CA	ASN	540	131.440	38.505	48.263	1.00	33.80
5015 5016	C O	ASN ASN	540 540	131.355 132.298	38.498 38.166	46.749 46.065	1.00 $1.00$	34.25 33.48
5017	CB	ASN	540	131.047	37.127	48.775	1.00	33.03
5018	CG	ASN	540	131.463	36.902	50.198	1.00	39.77
5019	OD1	ASN	540	130.776	36.219	50.965	1.00	45.10
5020	ND2	ASN	540	132.581	37.502	50.579	1.00	36.91
5021	H	ASN	540	129.781	39.280	49.311	1.00	25.00
5022	1HD2	ASN	540	132.850	37.386	51.503	1.00	25.00
5023	2HD2	ASN	540	133.079	38.025	49.919	1.00	25.00
5024	N	LEU	541	130.185	38.821	46.253	1.00	30.66
5025	CA	LEU	541	129.997	38.848	44.821	1.00	31.93
5026 5027	C	LEU	541 541	130.262	40.166 40.129	44.110 42.977	1.00	33.86
5027 5028	O CB	LEU LEU	541 541	130.805 128.600	38.308	44.486	1.00 $1.00$	30.07 34.62
5028	CG	LEU	541	128.194	36.907	44.990	1.00	35.64
5030	CD1	LEU	541	126.882	36.542	44.345	1.00	31.26

TABLE 11-continued

	In the Absence of Bound Substrate							
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
5031	CD2	LEU	541	129.256	35.866	44.669	1.00	29.53
5032	H	LEU	541	129.518	38.998	46.982	1.00	25.00
5033 5034	N CA	LEU LEU	542 542	129.910 130.075	41.305 42.581	44.706 44.033	$\frac{1.00}{1.00}$	35.33 39.16
5035	CA	LEU	542 542	131.084	43.566	44.635	1.00	42.69
5036	o	LEU	542	131.361	44.614	44.055	1.00	45.28
5037	CB	LEU	542	128.721	43.258	43.921	1.00	37.88
5038	CG	LEU	542	127.685	42.494	43.105	1.00	37.82
5039	CD1	LEU	542	126.275	42.902	43.505	1.00	37.78
5040	CD2	LEU	542	127.947	42.728	41.619	1.00	33.54
5041 5042	H N	LEU VAL	542 543	129.525 131.590	41.317 43.264	45.569 45.822	$\frac{1.00}{1.00}$	25.00 40.06
5043	CA	VAL	543	132.536	44.167	46.483	1.00	39.62
5044	C	VAL	543	133.900	43.601	46.457	1.00	40.35
5045	O	VAL	543	134.834	44.117	45.766	1.00	36.40
5046	CB	VAL	543	132.112	44.458	47.951	1.00	38.67
5047	CG1	VAL	543	133.154	45.323	48.643	1.00	41.60
5048 5049	CG2 H	VAL VAL	543 543	130.762 131.348	45.137 42.421	47.966 46.245	$\frac{1.00}{1.00}$	33.55 25.00
5050	N	ASP	544	134.175	42.421	47.191	1.00	39.19
5051	CA	ASP	544	135.485	41.887	47.274	1.00	37.12
5052	С	ASP	544	135.802	40.970	46.112	1.00	38.65
5053	O	ASP	544	134.991	40.124	45.739	1.00	42.40
5054	CB	ASP	544	135.609	41.070	48.566	1.00	37.00
5055	CG OD1	ASP	544	135.384	41.894	49.812 49.803	1.00	42.35
5056 5057	OD1 OD2	ASP ASP	544 544	135.659 134.933	43.114 41.304	50.813	$\frac{1.00}{1.00}$	49.35 50.35
5058	H	ASP	544	133.427	42.119	47.665	1.00	25.00
5059	N	SER	545	136.984	41.153	45.543	1.00	36.71
5060	CA	SER	545	137.444	40.303	44.464	1.00	39.73
5061	C	SER	545	138.200	39.158	45.142	1.00	38.96
5062 5063	O	SER	545 545	138.585	39.269	46.310 43.540	1.00	40.93 43.38
5063 5064	CB OG	SER SER	545 545	138.379 139.362	41.084 41.790	44.280	$\frac{1.00}{1.00}$	51.44
5065	Н	SER	545	137.544	41.896	45.832	1.00	25.00
5066	HG	SER	545	139.870	41.166	44.808	1.00	25.00
5067	N	ILE	546	138.377	38.048	44.442	1.00	36.92
5068	CA	ILE	548	139.109	36.920	45.011	1.00	40.59
5069	С	ILE	546	140.602	37.261	44.954	1.00	45.93
5070 5071	O CB	ILE ILE	546 548	141.117 138.839	37.620 35.612	43.889 44.226	1.00 1.00	46.41 36.34
5072	CG1	ILE	548	137.346	35.288	44.264	1.00	34.10
5073	CG2	ILE	546	139.629	34.456	44.829	1.00	31.04
5074	CD1	ILE	546	136.979	34.011	43.525	1.00	36.46
5075	H	ILE	546	138.009	37.990	43.533	1.00	25.00
5076	N	LYS	547	141.282	37.201	46.009	1.00	52.49
5077 5078	CA C	LYS LYS	547 547	142.706 143.483	37.502 36.450	46.134 45.353	$\frac{1.00}{1.00}$	58.52 60.95
5079	Ö	LYS	547	143.488	35.273	45.713	1.00	60.42
5080	СВ	LYS	547	143.217	37.599	47.572	1.00	59.36
5081	CG	LYS	547	144.684	38.023	47.659	1.00	68.28
5082	CD	LYS	547	145.065	38.553	49.037	1.00	72.08
5083	CE	LYS	547	146.486	39.105	49.029	1.00	74.86
5084	NZ	LYS	547 547	146.796	39.880	50.265	1.00	78.47
5085 5086	H 1HZ	LYS LYS	547 547	140.806 146.680	36.948 39.273	46.910 51.099	$\frac{1.00}{1.00}$	25.00 25.00
5087	2HZ	LYS	547	147.770	40.240	50.222	1.00	25.00
5088	3HZ	LYS	547	146.139	40.685	50.334	1.00	25.00
5089	N	ILE	548	144.086	36.890	44.254	1.00	67.72
5090	CA	ILE	548	144.868	36.018	43.381	1.00	76.79
5091	С	ILE	548	146.198	35.622	44.025	1.00	83.68
5092 5093	O CB	ILE ILE	548 548	146.583 145.120	34.440 36.678	43.897 41.986	1.00 $1.00$	86.24 75.68
5093	CG1	ILE	548	145.604	38.125	42.152	1.00	73.68 78.47
5095	CG2	ILE	543	143.855	36.623	41.137	1.00	68.49
5096	CD1	ILE	548	145.930	38.827	40.831	1.00	78.93
5097	OXT	ILE	548	146.823	36.492	44.672	1.00	92.78
5098	H	ILE	548	144.032	37.836	44.045	1.00	25.00
5099 5100	H	ILE	548	104.105	26.025	£2.020	1.00	64.63
5100 5101	MG MG	MG MG	851 852	104.185 102.138	36.235 43.657	53.030 49.009	$\frac{1.00}{1.00}$	61.83 62.23
5101	O	HOH	601	107.742	22.057	32.406	1.00	15.11
5102	Ö	НОН	602	122.540	22.695	37.531	1.00	32.44
5104	Ö	НОН	603	127.188	14.109	43.835	1.00	23.85

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
5105	O	НОН	604	123.257	32.177	37.651	1.00	25.21
5106	O	HOH	605	131.975	36.814	38.945	1.00	20.08
5107	O	HOH	606	130.320	38.579	40.729	1.00	28.69
5108	O	HOH	607	124.735	33.181	39.810	1.00	19.46
5109	O	НОН	608	119.958	22.714	50.725	1.00	24.82
5110	0	HOH	609	125.172	22.654	40.253	1.00	21.47
5111	0	HOH	610	106.047	21.994	29.826	1.00	26.03
5112 5113	0 0	НОН НОН	611 612	123.659 129.924	29.782 22.165	47.444 49.955	1.00 $1.00$	22.10 20.33
5114	Ö	HOH	613	117.254	16.672	36.732	1.00	18.88
5115	ŏ	НОН	614	131.911	22.935	48.204	1.00	23.59
5116	Ö	НОН	615	123.421	30.030	35.911	1.00	23.89
5117	O	НОН	616	128.952	30.316	38.829	1.00	22.41
5118	O	НОН	617	98.347	33.326	40.948	1.00	28.07
5119	O	HOH	618	126.062	19.250	36.922	1.00	29.11
5120	O	HOH	619	133.788	33.099	36.415	1.00	20.10
5121	O	HOH	620	127.252	22.013	48.848	1.00	24.10
5122	O	HOH	621	123.122	19.043	45.472	1.00	19.68
5123	0	НОН	622	124.636	25.767	41.845	1.00	42.37
5124	0	HOH	623	138.021	26.937	54.497	1.00	33.32
5125 5126	O O	НОН НОН	624 625	130.604 119.735	16.213	44.273 55.175	1.00	25.46 23.51
5127	Ö	HOH	626	109.560	17.425 43.332	32.386	$\frac{1.00}{1.00}$	27.79
5128	ŏ	НОН	627	104.016	36.817	39.018	1.00	24.34
5129	ŏ	НОН	628	134.051	35.256	29.604	1.00	37.22
5130	O	НОН	629	107.947	18.792	36.023	1.00	35.84
5131	O	HOH	630	129.821	19.576	48.096	1.00	29.63
5132	O	HOH	631	104.550	21.758	41.675	1.00	38.10
5133	O	HOH	632	111.970	10.709	47.161	1.00	23.86
5134	O	HOH	633	125.976	29.448	50.341	1.00	26.42
5135	O	НОН	634	97.143	36.787	48.102	1.00	35.12
5136	0	НОН	635	121.582	36.805	25.111	1.00	35.51
5137	0	HOH	636	113.756	26.801	22.571	1.00	30.58
5138	0	HOH	637	124.698	19.485	28.803	1.00	29.60
5139 5140	O O	НОН НОН	638 639	130.563 121.706	25.567 39.646	43.476 27.124	$\frac{1.00}{1.00}$	29.93 32.61
5141	Ö	HOH	640	104.749	34.099	30.683	1.00	28.14
5142	ŏ	НОН	641	111.751	8.174	35.080	1.00	34.23
5143	Ö	НОН	642	120.339	31.400	41.487	1.00	52.69
5144	O	HOH	643	95.163	26.623	43.384	1.00	36.83
5145	O	НОН	644	137.113	41.980	40.124	1.00	30.35
5146	O	HOH	645	116.126	11.318	49.986	1.00	25.34
5147	O	HOH	646	110.165	35.328	17.495	1.00	37.81
5148	O	HOH	647	118.054	20.287	30.749	1.00	33.12
5149	O	НОН	648	115.899	40.354	30.351	1.00	29.82
5150	0	НОН	649	113.524	54.000	32.295	1.00	30.14
5151	O O	НОН НОН	650 651	127.950	27.982	37.184 30.127	1.00	28.39
5152 5153	0	HOH	652	108.770 112.843	18.109 23.036	50.127	$\frac{1.00}{1.00}$	36.94 41.87
5154	Ö	HOH	653	132.804	32.747	50.167	1.00	34.56
5155	ŏ	НОН	654	99.278	32.670	36.214	1.00	31.88
5156	ŏ	HOH	655	93.100	36.093	41.777	1.00	39.13
5157	O	НОН	656	114.575	17.087	50.058	1.00	29.96
5158	O	НОН	657	134.890	18.651	45.599	1.00	29.79
5159	O	HOH	658	134.764	16.354	47.235	1.00	41.87
5160	O	HOH	659	138.146	19.452	46.210	1.00	40.62
5161	O	НОН	660	113.498	7.243	37.601	1.00	44.14
5162	O	НОН	661	118.735	25.324	49.539	1.00	32.46
5163	0	НОН	662	121.072	19.323	57.037	1.00	28.13
5164	0	HOH	663	120.647	52.139	31.728	1.00	31.21
5165 5166	0 0	НОН НОН	664 665	125.201 103.040	27.805 17.910	35.886 41.249	$\frac{1.00}{1.00}$	35.41 34.74
5167	Ö	HOH	666	92.281	23.719	49.317	1.00	36.36
5168	Ö	HOH	667	120.731	30.312	30.736	1.00	40.91
5169	ŏ	HOH	668	111.010	16.805	31.260	1.00	37.18
5170	Ö	НОН	669	98.374	30.892	39.496	1.00	39.09
5171	Ö	НОН	670	142.913	20.086	59.043	1.00	40.89
5172	O	НОН	671	120.070	4.238	32.203	1.00	32.10
5173	O	НОН	672	116.885	14.360	38.230	1.00	19.20
5174	O	HOH	673	135.198	31.364	38.159	1.00	21.99
5175	O	HOH	674	130.652	23.815	45.653	1.00	22.37
5176	O	НОН	675	116.184	18.170	25.042	1.00	33.65
5177	0	НОН	676	102.763	37.505	36.535	1.00	29.50
5178	О	НОН	677	113.482	17.709	47.318	1.00	24.10

TABLE 11-continued

		In the	Absence of	Bound S	ubstrate			
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
5179	O	HOH	678	128.292	24.082	47.295	1.00	27.62
5180	O	HOH	679	128.934	20.011	39.747	1.00	26.34
5181	O	HOH	680	129.840	32.556	48.799	1.00	34.07
5182	O	HOH	681	115.123	17.894	45.342	1.00	23.02
5183	O	HOH	682	134.875	11.928	61.810	1.00	24.68
5184	O	HOH	683	140.837	17.873	38.782	1.00	33.65
5185	O	HOH	684	135.724	8.315	55.152	1.00	37.93
5186	O	НОН	685	131.660	25.7655	6.520	1.00	36.71
5187	O	НОН	686	148.447	27.966	42.675	1.00	38.11
5188	O	НОН	687	110.190	10.176	45.195	1.00	35.74
5189	0	HOH	688	109.091	17.883	25.410	1.00	38.94
5190	0	HOH	6889	104.860	34.526	28.030	1.00	38.81
5191	0	НОН	690	102.070	36.177	27.889	1.00	35.60
5192	0	НОН	691	118.113	11.174	28.782	1.00	38.94
5193 5104	0	HOH	692 693	131.635	20.640	62.725	1.00	33.60
5194 5195	0	НОН НОН	694	136.344	35.530 31.406	31.124 33.335	$\frac{1.00}{1.00}$	36.08 31.14
5196	o	НОН	695	120.257 102.005	32.616	56.124	1.00	33.30
5197	Ö	НОН	696	124.575	21.994	35.468	1.00	36.59
5198	Ö	НОН	697	101.923	20.169	46.398	1.00	40.37
5199	Ö	НОН	698	129.243	49.171	40.765	1.00	49.17
5200	ŏ	НОН	699	139.196	35.578	48.616	1.00	31.26
5201	Ö	НОН	700	134.064	15.022	43.146	1.00	40.48
5202	ŏ	НОН	701	128.514	31.051	51.675	1.00	39.32
5203	ŏ	НОН	702	112.958	10.222	36.694	1.00	47.07
5204	ŏ	НОН	703	109.649	15.841	28.459	1.00	35.43
5205	Ō	НОН	704	140.094	42.685	39.958	1.00	41.31
5206	Ō	НОН	705	86.608	31.749	55.350	1.00	36.82
5207	O	НОН	706	128.605	34.147	28.351	1.00	35.79
5208	O	НОН	707	87.075	34.369	56.433	1.00	42.04
5209	O	HOH	708	89.030	34.345	44.620	1.00	40.07
5210	O	HOH	709	104.535	51.407	27.998	1.00	39.44
5211	O	HOH	710	120.125	34.187	24.397	1.00	63.74
5212	O	HOH	711	100.184	37.778	52.580	1.00	43.18
5213	O	HOH	712	109.218	37.444	46.111	1.00	37.68
5214	O	HOH	713	139.550	20.401	60.539	1.00	40.82
5215	O	HOH	714	140.612	17.7933	52.684	1.00	42.33
5216	O	HOH	715	120.330	21.170	32.392	1.00	31.20
5217	O	HOH	716	100.372	35.917	30.033	1.00	43.22
5218	O	HOH	717	120.163	23.899	33.930	1.00	33.67
5219	O	HOH	718	146.383	28.556	40.921	1.00	38.01
5220	O	НОН	719	109.966	20.788	31.041	1.00	38.62
5221	O	НОН	720	105.493	40.925	45.887	1.00	35.53
5222	0	НОН	721	119.171	27.937	23.152	1.00	55.39
5223	0	HOH	722	124.424	41.390	25.938	1.00	43.52
5224	0	НОН	723	102.779	17.993	48.134	1.00	38.38
5225	0	НОН	724 725	112.387	5.685 25.140	33.453	1.00	48.35
5226 5227	0	НОН НОН	725 726	151.082 127.089	21.203	44.349 29.049	$\frac{1.00}{1.00}$	35.50 45.21
5228	Ö	НОН	727	133.178	5.551	47.734	1.00	39.38
5229	Ö	НОН	728	151.127	34.628	33.927	1.00	42.02
5230	ŏ	HOH	729	150.405	22.240	44.559	1.00	38.43
5231	Ö	НОН	730	131.660	2.1074	7.933	1.00	37.78
5232	ŏ	НОН	731	135.465	8.584	52.047	1.00	40.15
5233	Ō	НОН	732	147.814	29.664	45.229	1.00	44.50
5234	O	HOH	733	140.989	33.094	47.707	1.00	43.19
5235	O	HOH	734	103.951	49.441	25.596	1.00	38.72
5236	O	HOH	735	86.471	53.747	29.731	1.00	43.56
5237	O	HOH	736	134.470	31.1682	5.54.6	1.00	52.39
5238	O	HOH	737	122.918	25.464	36.469	1.00	42.39
5239	O	HOH	738	99.309	33.456	31.178	1.00	48.32
5240	O	HOH	739	91.548	47.290	47.278	1.00	45.43
5241	O	HOH	740	92.024	43.380	40.690	1.00	42.02
5242	O	HOH	741	149.190	38.195	52.530	1.00	47.74
5243	0	HOH	742	153.088	41.575	36.804	1.00	46.51
5244	O	НОН	743	138.714	31.651	53.657	1.00	43.64
5245	O	НОН	744	143.900	19.054	51.722	1.00	40.32
5246	O	НОН	745	138.795	15.536	49.608	1.00	43.79
5247	0	HOH	746	124.711	3.430	56.077	1.00	44.40
5248	0	НОН	747	145.969	30.921	42.825	1.00	39.08
5249	0	НОН	748	134.979	10.249	59.470	1.00	35.78
5250 5251	0	НОН	749	133.932	40.151	29.911	1.00	41.40
5251 5252	0	НОН НОН	750 751	114.521 129.614	21.309 38.180	22.697 25.426	1.00 $1.00$	38.72 39.89
3434	0	11011	731	127.014	20.100	20.720	1.00	57.07

TABLE 11-continued

In the Absence of Bound Substrate										
Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor		
5253	O	НОН	752	111.6443	13.087	29.735	1.00	45.90		
5254	0	HOH	753 754	104.216	21.388	44.848	1.00	33.35		
5255 5256	0	HOH	754 755	110.986	12.520	49.459	1.00	49.32		
5256 5257	0	HOH	755 756	139.600 113.295	40.725	48.728	1.00	46.07		
5257 5258	O O	НОН НОН	756 757	127.101	9.448 23.382	29.832 34.156	$\frac{1.00}{1.00}$	35.78 48.02		
5259	o	НОН	758	127.101	18.490	63.251	1.00	46.33		
5260	Ö	нон	759	130.420	26.867	25.702	1.00	40.40		
5261	Ö	НОН	760	122.231	3.237	35.918	1.00	44.61		
5262	Ö	НОН	761	128.310	26.484	40.968	1.00	32.14		
5263	ŏ	НОН	762	88.443	24.530	48.586	1.00	57.07		
5264	ŏ	НОН	763	103.542	23.739	25.080	1.00	45.05		
5265	0	НОН	764	116.278	57.331	34.559	1.00	42.40		
5266	O	HOH	765	120.787	5.886	61.156	1.00	43.73		
5267	O	HOH	766	142.631	40.352	42.775	1.00	65.94		
5268	O	HOH	767	124.244	13.057	63.666	1.00	43.68		
5269	O	HOH	768	101.830	22.900	29.735	1.00	36.47		
5270	O	HOH	769	137.190	5.022	37.071	1.00	50.65		
5271	O	HOH	770	135.078	34.403	50.639	1.00	51.53		
5272	O	HOH	771	103.266	58.719	26.225	1.00	46.58		
5273	O	HOH	772	144.319	16.861	24.565	1.00	53.32		
5274	O	НОН	773	127.856	47.718	31.019	1.00	45.45		
5275	0	HOH	774	95.530	18.110	49.546	1.00	52.47		
5276	0	НОН	775	148.435	20.165	43.831	1.00	49.25		
5277	0	НОН	776 777	118.026	13.535	59.021	1.00	48.41		
5278 5279	Ö	НОН НОН	778	110.119 110.457	43.903 61.356	16.201 39.879	$\frac{1.00}{1.00}$	37.10 44.66		
5280	Ö	НОН	779	105.313	56.879	27.692	1.00	51.08		
5281	Ö	НОН	780	106.267	19.656	28.049	1.00	45.55		
5282	Ö	НОН	781	122.226	20.789	29.638	1.00	45.73		
5283	ŏ	НОН	782	107.680	19.165	33.248	1.00	35.37		
5284	Ō	НОН	783	141.434	30.527	58.190	1.00	56.49		
5285	O	HOH	784	121.953	27.180	30.544	1.00	43.22		
5286	O	HOH	785	116.050	27.492	52.913	1.00	59.86		
5287	O	HOH	786	115.271	11.494	53.629	1.00	47.46		
5288	O	HOH	787	136.166	43.700	43.430	1.00	44.89		
5289	O	HOH	788	123.135	5.923	32.296	1.00	61.24		
5290	O	HOH	789	148.342	38.089	38.232	1.00	41.22		
5291	O	НОН	790	112.195	39.980	44.065	1.00	44.26		
5292	O	НОН	791	108.340	50.773	20.100	1.00	62.55		
5293	O	НОН	792	126.140	29.670	29.775	1.00	38.87		
5294 5295	0	НОН	793	122.347	26.176	27.904	1.00	47.43		
5295 5296	O O	HOH	794 795	105.375 146.608	13.283 19.061	37.860 33.529	1.00 $1.00$	40.63 50.53		
5290 5297	0	НОН НОН	793 796	112.240	28.192	56.028	1.00	54.08		
5298	Ö	НОН	797	106.519	16.717	37.160	1.00	39.17		
5299	ŏ	НОН	798	122.257	-2.147	57.632	1.00	59.87		
5300	ŏ	НОН	799	105.969	47.469	20.174	1.00	42.44		
5301	Ō	НОН	800	124.201	23.387	29.951	1.00	51.85		
5302	O	HOH	801	104.010	26.139	23.199	1.00	57.02		
5303	O	HOH	802	106.547	37.540	47.839	1.00	46.00		
5304	O	HOH	803	126.083	27.795	33.246	1.00	45.66		
5305	O	HOH	804	93.229	25.530	63.301	1.00	50.45		
5306	O	НОН	805	126.637	14.627	66.291	1.00	54.63		
5307	O	НОН	806	117.649	48.031	30.248	1.00	44.41		
5308	0	HOH	807	112.889	34.483	45.820	1.00	41.77		
5309	0	НОН	808	143.749	8.474	39.051	1.00	58.35		
5310	0	HOH	809	117.223	16.467	56.527	1.00	54.55		
5311 5312	0 0	НОН НОН	810 811	136.640 130.573	48.794 47.831	42.640 52.219	1.00 $1.00$	59.70 43.65		
5313	Ö	НОН	812	119.790	22.620	53.732	1.00	49.88		
5314	Ö	НОН	813	105.220	9.911	43.334	1.00	53.82		
5315	ŏ	НОН	814	94.459	22.230	65.891	1.00	53.43		
5316	Ö	НОН	815	145.893	33.119	447.904	1.00	50.15		
5317	ŏ	НОН	816	137.540	19.003	49.581	1.00	32.04		
5318	O	НОН	817	127.395	18.676	22.177	1.00	58.02		
5319	Ō	НОН	818	135.930	19.361	20.695	1.00	61.65		
5320	O	HOH	819	122.368	-4.865	43.028	1.00	43.72		
5321	O	HOH	820	117.352	52.131	24.538	1.00	49.67		
5322	O	HOH	821	129.874	51.577	33.814	1.00	58.12		
5323	O	HOH	822	129.360	28.179	34.594	1.00	43.67		
5324	O	HOH	823	97.243	40.051	31.308	1.00	40.94		
5325	O	НОН	824	119.361	23.189	24.691	1.00	55.59		
5326	О	HOH	825	105.947	8.433	39.961	1.00	47.78		

TABLE 11-continued

Structural (	Coordinates	of Tobacco	5-Epi-Aristolo	chene Synthase
	In the A	bsence of B	Sound Substrate	

Atom Type	Atom	Residue	Residue #	X	Y	Z	OCC	B-factor
5327	O	НОН	826	124.177	-6.929	48.285	1.00	50.47
5328	Ō	НОН	827	143.743	41.219	49.977	1.00	54.42
5329	O	HOH	828	117.815	15.765	23.926	1.00	47.10
5330	0	HOH	829	106.852	11.509	45.366	1.00	59.91
5331	O	HOH	830	114.340	49.442	45.031	1.00	54.21
5332	O	НОН	831	107.212	10.319	38.018	1.00	47.91
5333	O	HOH	832	89.843	54.539	37.711	1.00	55.79
5334	O	HOH	833	115.120	21.415	49.941	1.00	40.64
5335	O	HOH	834	119.324	14.942	62.472	1.00	63.27
5336	O	HOH	835	149.479	14.241	50.723	1.00	65.18
5337	O	HOH	836	99.208	46.311	26.331	1.00	59.48
5338	O	HOH	837	146.479	34.108	25.046	1.00	49.79
5339	O	HOH	838	117.731	49.616	19.065	1.00	60.85
5340	O	HOH	839	115.539	6.301	34.276	1.00	51.97
5341	O	HOH	840	97.213	27.831	34.233	1.00	45.30
5342	O	HOH	841	89.788	22.728	43.919	1.00	61.79
5343	O	HOH	842	147.830	32.323	40.885	1.00	46.95
5344	O	HOH	843	132.462	17.381	68.762	1.00	50.53
5345	O	HOH	844	140.816	13.261	39.613	1.00	50.48
5346	O	HOH	845	131.788	48.689	43.107	1.00	55.44
5347	O	HOH	846	106.451	38.430	52.704	1.00	44.59
5348	O	HOH	847	112.522	3.225	51.067	1.00	62.24
5349	O	HOH	848	116.588	33.059	17.286	1.00	51.54
5350	O	HOH	849	121.984	13.530	21.831	1.00	59.69
5351	O	HOH	850	121.351	34.646	19.580	1.00	63.69
5352	O	HOH	853	119.444	26.300	52.657	1.00	48.12
5353	O	HOH	854	119.223	18.972	28.280	1.00	43.53
5354	O	HOH	855	109.476	29.077	61.498	1.00	46.95
5355	O	HOH	856	96.378	36.846	50.773	1.00	37.88
5356	O	HOH	857	96.918	46.467	51.605	1.00	69.73
5357	O	HOH	858	97.861	35.983	32.096	1.00	48.71
5358	O	HOH	859	105.582	44.217	22.626	1.00	52.96
5359	O	HOH	860	111.207	54.577	33.852	1.00	44.86
5360	O	HOH	861	106.475	45.773	50.620	1.00	52.70
5361	O	HOH	862	136.750	45.222	40.123	1.00	53.92
5362	O	HOH	863	134.438	43.600	31.414	1.00	51.51
5363	O	HOH	864	147.130	24.676	49.884	1.00	42.49
5364	O	HOH	865	126.425	22.757	59.405	1.00	54.25
5365	O	HOH	866	135.514	7.098	48.245	1.00	59.13
5366	O	HOH	867	114.942	1.622	48.125	1.00	56.08
5367	O	HOH	868	119.740	-4.108	46.312	1.00	51.35
5368	O	HOH	869	134.478	8.308	29.219	1.00	53.23
5369	O	HOH	870	127.297	14.232	21.009	1.00	54.19
5370	O	НОН	871	134.315	17.294	22.547	1.00	59.58
5371	O	HOH	872	130.159	26.543	36.441	1.00	34.46
5372	Ö	НОН	873	136.207	18.694	43.344	1.00	35.20
5373	Ö	НОН	874	134.779	10.368	41.428	1.00	45.81
5374	Ö	НОН	875	137.054	3.899	33.453	1.00	51.47
5375	0	НОН	876	145.762	17.318	28.638	1.00	52.42
5376	0		877		20.944	29.342	1.00	
33/0	U	НОН	0//	146.344	∠0.944	29.342	1.00	47.62

## TABLE 12

	bits (419), Expect = 5e-41 = 88/270 (32%), Positives = 152/270 (55%), Gaps = 5/270 (1%)	
Query: 1	DRVVECYFWALGVYFEPQYSQARVMLVKTISMISIVDDTFDAYGTVKELEAYTDAIQRWD DR+VECYFW G+ O++ AR+M+ K ++I+++DD +D YGT++BLE +TD I+RWD	60
Sbjct: 316	DRLVECYFWNTGIIEPRQHASARIMMGKVNALITVIDDIYDVYGTLEELEQFTDLIRRWD	375
Query: 61	INEIDRLPDYMKISYKAILDLYKDYEKELSSAGRSHIVCHAIERMKEVVRNYNVESTWFI IN ID+LPDYM++ + A+ + D ++ +++ + + + Y VE+ WF	120
Sbjct: 376	INSIDQLPDYMQLCFLALNNFVDDTSYDVMKEKGVNVIPYLRQSWVDLADKYMVEARWFY	435
Query: 121	EGYMPPVSEYLSNALATTTYYYLATTSYLGM-KSATEQDFEWLSKNPKILEASVIICRVI G+ P + EYL N+ + + + T + + S T++ + L K ++ S + R+	179
Sbjct: 436	GGHKPSLEEYLENSWQSISGPCMLTHIFFRVTDSFTKETVDSLYKYHDLVRWSSFVLRLA	495
Query: 180	DDTATYEVEKSRGQIATGIECCMRDYGISTKEAMAKFQNMAETAWKDIN-EGLLRPTPVS DD T E SRG + ++C M DY S EA + + WK +N E + + +P	238
Sbjct: 496		555

## TABLE 12-continued

Query:	239	TEFLTPILNLARIVEVTYIHNLDGYTHP	
		+F+ ++L R+ ++ Y HN DG+ HP	
Sbjct:	556	KDFIGCAVDLGRMAQLMY-HNGDGHGTQHP	

### TABLE 13

		bits (289), Expect = 1e-25 = 77/270 (28%), Positives = 126/270 (46%), Gaps = 6/270 (2%)	
Query:	3	VAEVYFSSATFEP-EYSATRIAFTKIGCLQVLFDDMADIFATLDELKSFTEGVKRWDTSL V +++ FEP ++ R I L + DD+ D++ TLDEL+ FT+ KRWDT	61
Sbjct:	318	VESFFWAVGMFEPHQHGYQRKMAATIIVLATVIDDIYDVYGTLDELELFTDTFKRWDTES	377
Query:	62	$ \begin{array}{llllllllllllllllllllllllllllllllllll$	121
Sbjct:	378	TRLPYYMOLCYWGVHNYISDAAYDILKEHGFFCLQYLRKSVVDLVEAYFHEAKWYHSGY	437
Query:	122	IPTFEEYLKTYAISVGLGPCTLQPILLMGELVKDDVVEKVHYPSNMFELVSLSWRLTN P+ +EYL	179
Sbjct:	438	TPSLDEYLNIAKISVA-SPAIISPTYFTFANASHDTAVIDSLYQYHDILCLAGIILRLPD	496
Query:	180	DTKTYQAEKARGQQASGIACYMKDNPGATEEDAIKHICRVVDRALKEASFEYFKPSNDIP D T E ARG I CYMK+ A+EE+A++H+ ++ A K+ + P	239
Sbjct:	497	DLGTSYFELARGDVPKTIQCYMKET-NASEEEAVEHVKFLIREAWKDMN-TAIAAGYPFP	554
Query:	240	MGCKSFIFNLRLCVQIFYKFIDGYGIANEE	269
Sbjct:	555	G + N+ Q Y DG+G+ + + DGMVAGAANIGRVAQFIYLHGDGFGVQHSK	584

### TABLE 14

	0 bits (299), Expect + 6e-27 s = 70/272 (25%), Positives = 137/272 (49%), Gaps = 3/272 (1%)	
Quer <b>y:</b> 2	RVVECYFWALGVYFEPQYSQARVMLVKTISMISIVDDTFDAYUTVKELEAYTDAIQRWDI R VE Y W + FEP++S++R+ KT + +++DD +D + T+ E++ T+ ++RWD+	6
Sbjct: 296	RHVEYYSWVVMCIFEPEFSESRIAFAKTAILCTVLDDLYDTHATLHELKIMTEGVRRWDL	355
Query: 62	NEIDRLPDYMKISYKAILDLYKDYEKELSSAGRSHIVCHAIERMKEVVRNYNVESTWFIE + D LPDY+KI+++ + + E+ + K + +Y E+ W	121
Sbjct: 356	S SLTDDLPDYIKIAFQFFFNTVNELIVEIVKRQGRDMTTIVKDCWKRYIESYLQEAEWIAT	415
Query: 122	P. GYMPPVSEYLSNALATTTYYYLATTSYLGM-KSATEQDFEWLSKNPKILEASVIICRVID G++P +EY+ N +A++ L L + K + E + KIL+ + R+ D	180
Sbjct: 416	G GHIPTFNEYIKNGMASSGMCILNLNPLLLLDKLLPDNILEQIHSPSKILDLLELTGRIAD	475
Quer <b>y:</b> 181	DTATYEVEKSRGQIATGIECCMRDYGISTKE-AMAKFQNMAETAWKDINEGLLRPTPVST D +E EK RG++A+ ++C M++ ST E A+ + + + + + N ++ V	239
Sbjct: 476	DLKDFEDEKERGEMASSLQCYMKENPESTVENALNHIKGILNRSLEEFNWEFMKQDSVPM	535
Query: 240	) EFLTPILNLARIVEVTYIHNLDGYTHPEKVLK N+ R ++ Y + DG +K +K	271
Sbjct: 536	CCKKFTFNIGRGLQFIYKYR-DGLYISDKEVK	566

### TABLE 15

Score = 221 bits (557), Expect = 4e-57 Identities = 20/263 (42%), Positive = 178/283 (62%, Gaps = 6/293 (2%)										
Query:	5	EFYFWMAAAISEPEFSGSRVAFTKIAILMTMLDDLYDTHGTLDQLKIFTEGVRRWDVSLV E YF A+ I EPEFS R +TK + +LDDLYD HG+LD LK+FTE V+RWD+SLV	64							
Sbjct:	589		648							
Query:	65	${\tt EGLPDEMKIAFEFWLKTSNELIAEAVKAQGQDMAAYIRKNAWERYLEAYLQ0AEWIATGH}$	124							
Sbjct:	649	+ +P MKI F + T N++ E + QG+D+ YI +N W+ LEAY ++AEW + DQMPQQMKICFVGFYNTFNDIAKEGRERQGRDVLGYI-QNVWKVQLEAYTKEAEWSEAKY	707							
Query:	125	~	184							
Sbjct:	708	$ \begin{array}{llllllllllllllllllllllllllllllllllll$	767							

### TABLE 15-continued

Query: 185	RDFQAEKDHGDL-SCIECYLKDHPESTVEDALNHVNGLLGNCLLEMNWKFLKKQDSVPLS + +OAE+ G++ S I+CY+KDHP+ + E+AL HV ++ N L E+N +F+ + +P	243
Sbjct: 768	KTYQAERGQGEVASAIQCYMKDHPKISEEEALQHVYSVMENALEELNREFVNNKIPDI	825
Query: 244	CKKYSFHVLARSIQFMYNQGDGFSISNKV-IKDQVQKVLIVPV K+ F AR +O Y OGDG ++S+ + IK+ V+ L PV	285
Sbjct: 826	YKRLVFET-ARIMQLFYMQGDGLTLSHDMEIKEHVKNCLFQPV	867

#### SEQUENCE LISTING

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	gat Asp														147	
	gct Ala														195	
	gca Ala														243	
	gaa Glu 75														291	
	ttg Leu														339	
	tct Ser														387	
	cct Pro														435	
	tct Ser														483	
	cat His 155														531	
	tcc Ser														579	
	agg Arg														627	

						cga Arg										675
						gtg Val										723
						cac His 240										771
						ttt Phe										819
_	-	_	_	_		ttt Phe		-			_					867
				_	-	gtc Val	_		-	_				_		915
_		-	-	-		ttt Phe	-	-				-		-		963
	-			-	-	ata Ile 320		-		-			-		-	1011
						aaa Lys										1059
	_	_		_	_	gaa Glu	_		_	_		-				1107
						aga Arg										1155
						att Ile										1203
		_		_		gca Ala 400										1251
						aag Lys										1299
						att Ile										1347
						acg Thr										1395
						tgc C <b>y</b> s										1443
						ttt Phe 480										1491
						ctt Leu										1539
						ctt Leu										1587

#### -continued

515 cac aat cta gat gga tac act cat ccg gag aaa gtc tta aaa cct cac 1635 His Asn Leu Asp Gly Tyr Thr His Pro Glu Lys Val Leu Lys Pro His 530 att att aac cta ctt gtg gac tcc atc aaa att tga 1671 Ile Ile Asn Leu Leu Val Asp Ser Ile Lys Ile <210> SEQ ID NO 2 <211> LENGTH: 548 <212> TYPE: PRT <213> ORGANISM: Nicotiana tabacum <400> SEQUENCE: 2 Met Ala Ser Ala Ala Val Ala Asn Tyr Glu Glu Glu Ile Val Arg Pro 10 Val Ala Asp Phe Ser Pro Ser Leu Trp Gly Asp Gln Phe Leu Ser Phe Ser Ile Lys Asn Gln Val Ala Glu Lys Tyr Ala Gln Glu Ile Glu Ala 35  $\phantom{\bigg|}40\phantom{\bigg|}40\phantom{\bigg|}45\phantom{\bigg|}$ Leu Lys Glu Gln Thr Arg Asn Met Leu Leu Ala Thr Gly Met Lys Leu 50 60Ala Asp Thr Leu Asn Leu Ile Asp Thr Ile Glu Arg Leu Gly Ile Ser 65 70 75 80 Tyr His Phe Glu Lys Glu Ile Asp Asp Ile Leu Asp Gln Ile Tyr Asn Gln Asn Ser Asn Cys Asn Asp Leu Cys Thr Ser Ala Leu Gln Phe Arg \$100\$Leu Leu Arg Gln His Gly Phe Asn Ile Ser Pro Glu Ile Phe Ser Lys Phe Gln Asp Glu Asn Gly Lys Phe Lys Glu Ser Leu Ala Ser Asp Val 130  $$135\$ Leu Gly Leu Leu Asn Leu Tyr Glu Ala Ser His Val Arg Thr His Ala Asp Asp Ile Leu Glu Asp Ala Leu Ala Phe Ser Thr Ile His Leu Glu Ser Ala Ala Pro His Leu Lys Ser Pro Leu Arg Glu Gln Val Thr His Ala Leu Glu Gln Cys Leu His Lys Gly Val Pro Arg Val Glu Thr Arg 195  $\phantom{\bigg|}200\phantom{\bigg|}205\phantom{\bigg|}$ Leu Leu Arg Phe Ala Lys Leu Asp Phe Asn Leu Leu Gln Met Leu His Lys Gln Glu Leu Ala Gln Val Ser Arg Trp Trp Lys Asp Leu Asp Phe Val Thr Thr Leu Pro Tyr Ala Arg Asp Arg Val Val Glu Cys Tyr Phe  $260 \hspace{1cm} 265 \hspace{1cm} 270 \hspace{1cm}$ Trp Ala Leu Gly Val Tyr Phe Glu Pro Gln Tyr Ser Gln Ala Arg Val Met Leu Val Lys Thr Ile Ser Met Ile Ser Ile Val Asp Asp Thr Phe Asp Ala Tyr Gly Thr Val Lys Glu Leu Glu Ala Tyr Thr Asp Ala Ile 315 Gln Arg Trp Asp Ile Asn Glu Ile Asp Arg Leu Pro Asp Tyr Met Lys

												COII	CTII	ueu			
				325					330					335			
Ile	Ser	Tyr	Lys 340	Ala	Ile	Leu	Asp	Leu 345	Tyr	Lys	Asp	Tyr	Glu 350	Lys	Glu		
Leu	Ser	Ser 355	Ala	Gly	Arg	Ser	His 360	Ile	Val	Cys	His	Ala 365	Ile	Glu	Arg		
Met	<b>Lys</b> 370	Glu	Val	Val	Arg	Asn 375	Tyr	Asn	Val	Glu	Ser 380	Thr	Trp	Phe	Ile		
Glu 385	Gly	Tyr	Thr	Pro	Pro 390	Val	Ser	Glu	Tyr	Leu 395	Ser	Asn	Ala	Leu	Ala 400		
Thr	Thr	Thr	Tyr	Tyr 405	Tyr	Leu	Ala	Thr	Thr 410	Ser	Tyr	Leu	Gly	Met 415	Lys		
Ser	Ala	Thr	Glu 420	Gln	Asp	Phe	Glu	Trp 425	Leu	Ser	Lys	Asn	Pro 430	Lys	Ile		
Leu	Glu	Ala 435	Ser	Val	Ile	Ile	Cys 440	Arg	Val	Ile	Asp	Asp 445	Thr	Ala	Thr		
Tyr	Glu 450	Val	Glu	Lys	Ser	Arg 455	Gly	Gln	Ile	Ala	Thr	Gly	Ile	Glu	Cys		
Cys 465		Arg	Asp	Tyr	Gly 470		Ser	Thr	Lys	Glu 475		Met	Ala	Lys	Phe 480		
	Asn	Met	Ala	Glu 485		Ala	Trp	Lys	Asp		Asn	Glu	Gly	Leu 495			
Arg	Pro	Thr	Pro		Ser	Thr	Glu	Phe		Thr	Pro	Ile	Leu 510		Leu		
Ala	Arg	Ile 515		Glu	Val	Thr	<b>Tyr</b> 520		His	Asn	Leu	Asp 525		Tyr	Thr		
His		Glu	Lys	Val	Leu			His	Ile	Ile			Leu	Val	Asp		
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		CATI			(1	1644	)										
		t ca			a++	ac-	227	+=+	us -	a	ne~	a++	a++	cac	ccc	ло	
		tca Ser														48	
		gac Asp	Phe					Trp					Leu			96	
tac	a++	gat	20	cac	a++	aca	ass	25	+a+	ac+	Cas	aea	30 a++	asa	aca	144	
		Asp 35														144	
		gaa Glu														192	
		aca Thr														240	
65	1				70		r			75	,		1		80		
		ttt Phe														288	

cam acc tom acc togs act get titg togs acc tot gos out come tits cog of clin Abn Ser Abn Cys Abn Abp Leu Cys Thr Ser Ala Leu Gin The Arg 100 100 100 100 100 100 100 100 100 10	_																	
Leu Leu Arg oln His Gly Phe Asn Ile Ser Pro Glu Ile Phe Ser Lys 115  tto cas gat gas aast ggc aas tto aag gag tot ott got agt gat gtc Phe Gln Asp Glu Asn Gly Lys Phe Lys Glu Ser Leu Ala Ser Asp Val 130  tta gga tta tta aac ttg tat gas gct tca cat gta agg act cat gct Leu Glu Leu Leu Asn Leu Tyr Glu Ala Ser His Val Arg Thr His Ala 160  gac gat atc tta gas gcc gca ctt gct tto tcc act atc cat ctt gas Asp Asp Ile Leu Glu Asp Ala Leu Ala Phe Ser Thr Ile His Leu Glu Asp Ala Leu Ala Phe Ser Thr Ile His Leu Glu Fr His 160  gac gat atc tta gas gcc gca ctt gct tto tcc act atc cat ctt gas Asp Asp Ile Leu Glu Asp Ala Leu Ala Phe Ser Thr Ile His Leu Glu Fr His 160  gcc ctt gag cas atg ttg cac aag ggt gtt cct aga gtc gag acc cgs Ala Leu Glu Gln Cys Leu His Lys Gly Val Pro Arg Val Glu Gln Thr Arg 205  tto tto atc tca tca atc tat gac aag gga caa atc gas aat aat gtg Phe Phe Ile Ser Ser Ile Tyr Asp Lys Glu Gln Ser Lys Asn Asn Val 2215  tta ctt cat tca tca tat gat gat ttc aac ttg ctc cag atg ttg cac Leu Leu Arg Phe Ala Lys Leu Asp Phe Asn Leu Gln Met Leu His Lys Leu Asp Phe Asn Leu Gln Met Leu His Lys Gly Val Pro Arg Val Glu Chn Met Leu His Lys Glo Gln Ser Lys Asn Asn Val 2215  gaa aca aca ctt gct cas gta tca agg tgg tgg asa gat ttg gat ttt Cay Gln Glu Leu Ala Gln Val Ser Arg Trp Trp Lys Asp Leu Asp Phe 230  aaa caa gas ctt gct cas gta tca agg tgg tgg asa gat ttg gat ttt Lys Gln Glu Leu Ala Gln Val Ser Arg Trp Trp Lys Asp Leu Asp Phe 245  gag gas tta gga gtt tat ttt gag cct caa tac tct caa gct cgc gc Gl agg gas ct agg gat ta gat gat gat gat gat gat gat					Asn					Cys					Gln			336
Phe Gln Asp Glu Asn Gly Lys Phe Lys Glu Ser Leu Ala Ser Asp Val 135 135 140 135 140 135 140 135 140 135 140 135 140 135 140 135 140 155 160 160 151 145 160 155 160 155 160 165 165 165 165 165 165 165 165 165 165				Arg					Asn					Ile				384
Leu Gily Leu Leu Asn Leu Tyr Glu Ala Ser His Val Arg Thr His Ala 150 150 160 155 160 155 160 155 160 155 160 155 160 155 160 155 160 155 160 155 160 155 165 165 165 170 170 170 170 170 170 170 170 170 170		ne	Gln					Lys					Leu					432
Asp Asp Ile Leu Glu Asp Ala Leu Ala Phe Ser Thr Ile His Leu Glu 175  tot goa got coa cat ttg aaa tot coa ott agg gag caa gtg aca cat Ser Ala Ala Pro His Leu Lys Ser Pro Leu Arg Glu Gln Val Thr His 180  goc ott gag caa tgt ttg cac aag ggt gtt cot aga gtc gag acc cga Ala Leu Glu Gln Cys Leu His Lys Gly Val Pro Arg Val Glu Thr Arg 195  tot tot ato toa toa ato tat gac aag gat cat cga aga aca at gtg Phe Phe Phe Ile Ser Ser Ile Tyr Asp Lys Glu Gln Ser Lys Asn Asn Val 210  tta ott cga ttt goc aaa ttg gat ttc aac ttg ctc cag atg ttg cac 225  tta ott cga ttt goc aaa ttg gat ttc aac ttg ctc cag atg ttg cac 240  aaa caa gaa ctt got caa gta tca agg tgg tgg aaa gat ttg gat ttc Lys Gln Glu Leu Ala Gln Val Ser Arg Trp Trp Lys Asp Leu Asp Phe 245  gta aca aca ctt cca tat gct aga gat cga gta gtt gaa tgg tac ttt Val Thr Thr Leu Pro Tyr Ala Arg Asp Arg Val Val Glu Cys Try Phe 260  gag gca tta gga gtt tat ttt gag cot caa tac tot caa gct cgc gtc Glu Ala Leu Gly Val Tyr Phe Glu Pro Gln Tyr Ser Gln Ala Arg Val 275  atg ct tac ggt aca gt aca gt aca atg att tog att gat gat gac acc tt Met Leu Val Lys Thr 11e Ser Met Ile Ser Ile Val Asp Asp App Thr Phe 290  gat got tac ggt aca gt aca gat aca gac ctt gag gac aca ctt 280  gat got tac ggt aca gt aca gat aca gac ctt gag gat aca aca gat gcc aca Asp Ala Tyr Gly Thr Val Lys Glu Leu Glu Ala Tyr Thr Asp Ala Ile 305  aca aga tgg gat atc aca gca att cta gat ctc aca gg tta aca gaa gat 229  gat gct tac ggt aca gat aca gat ct aca gat tta gac aca gaa 230  caa aga tgg gat atc aca caa aca act gac ct cac aca gat gcc ata 310  aca aga tgg gat atc aca gt aca gt aca gac ctt gca aca gat aca 330  act agt tat aca gct att cta gat ctc tac aca gg tta aca aca 330  act agt tat aca gct att cta gat ctc tac aca ggat tat gaa aca 330  act aga tat aca gct att cta gat ctc tac aca ggat tat gaa 330  act aca gaa gat gta aga act tat aca gat ctc gca aca aca 330  act aca aca aca cta tac ctc gca aca act gca aca tag aca 330  act aca cac act att cac ctc gca aca act gca cta gca 330  act acc aca tat tac ctc gca cac aca acc cta ttg gc	Le	eu					Leu					His					Ala	480
Ser Åla Åla Pro His Leù Lys Ser Pro Leu Arg Glú Gln Val Thr His 180 cc ctt gag caa tgt ttg cac aag ggt gtt cct aag gtc gag acc cga Ala Eue Glu Gln Cys Leu His Lys Gly Val Pro Arg Val Glu Thr Arg 200 ctt ctc atc atc tca tca atc tat gac aag gaa caa tcg aag aat aat gtg phe phe 11e Ser Ser Ile Tyr Asp Lys Glu Gln Ser Lys Asn Asn Val 210 c200 ctt ctc dat cta ga aat tag gat ttc aac ttg ctc cag atg ttg cac 221 cac Leu Leu Arg Phe Ala Lys Leu Asp Phe Asn Leu Leu Gln Met Leu His 2230 caa aca aga act gad ctt gct caa ggt ttg ga aag gat ttg gat ttt 2235 cac 235 cac 244 cac 235 cac 244 cac 235 cac 244 cac 245 cac 245 cac 245 cac 245 cac 255 ca						Glu					Phe					Leu		528
Ala Leu Glu Gln Cys Leu His Lys Gly Val Pro Arg Val Glu Thr Arg 200  ttc ttc atc tca atc tat gac aag gaa caa tcg aag aat aat gtg Phe Phe Ile Ser Ser Ile Tyr Asp Lys Glu Gln Ser Lys Asn Asn Val 210  tta ctt cga ttt gcc aaa ttg gat ttc aac ttg ctc cag atg ttg cac Leu Leu Arg Phe Ala Lys Leu Asp Phe Asn Leu Leu Gln Met Leu His 220  aaa caa gaa ctt gct caa gta tca agg tgg tgg aaa gat ttg gat ttt Lys Gln Glu Leu Ala Gln Val Ser Arg Trp Trp Lys Asp Leu Asp Phe 250  gta aca aca ctt cca tat gct agg gat cga gta gtt gaa tgc tac ttt Val Thr Thr Leu Pro Tyr Ala Arg Asp Arg Val Val Glu Cys Tyr Phe 260  gag gca tta gga gtt tat ttt gag cct caa atc tc caa gct cgc gtc Glu Ala Leu Gly Val Tyr Phe Glu Pro Gln Tyr Ser Gln Ala Arg Val 275  atg ctc gtt aag acc ata caa atg att tcg att gt gat acc acc ttt Met Leu Val Lys Thr Ile Ser Met Ile Ser Ile Val Asp Asp Thr Phe 290  gat gct tac ggt aca gtt aaa gaa ctt gag gca tac aca gat gcc ata Asp Asp Ala Tyr Gly Thr Val Lys Glu Leu Glu Ala Tyr Thr Asp Ala Ile 305  atc agg gat atc aca gat att caa gat ctg gc ct cat aca gat gcc ata Asp Ala Tyr Gly Thr Val Lys Glu Leu Glu Ala Tyr Thr Asp Ala Ile 305  atc agt tat aaa gct att cta gat ctc tac aag gat tac aaa gaa caa gaa aga aga aga leu Pro Asp Tyr Met Lys 325  atc agt tat aaa gct att cta gat ctc tac aag gat tac gaa aga aga l104  Leu Ser Tyr Lys Ala Ile Leu Asp Leu Tyr Lys Asp Tyr Glu Lys Glu 355  atc agt tat aaa gct att cta gat ctc tac aag gat tac aaa aga aga l104  Leu Ser Ser Ala Gly Arg Ser His Ile Val Cys His Ala Ile Glu Arg 355  atg aaa gaa gta gta aga att tat aat gtc gg tca aca aca gat gaa aga l104  Leu Ser Ser Ala Gly Arg Ser His Ile Val Cys His Ala Ile Glu Arg 355  atg aaa gaa ta atg cca cct gtt tct gaa tac cta agc aat gca cta gca l104  Met Lys Glu Val Val Arg Asp Tyr Asn Val Glu Lys Glu 395  atg aaa gaa ta atg cca cct gtt tct gaa tac cta agc aat gca cta gca l206  act acc aca tat tac tac ctc gcg aca acc tcg tat ttg gcg atg aac acc tag acc acc acc acc acc acc acc gcd gat acc acc acc acc acc acc acc acc acc gcd gat acc acc acc acc acc acc acc a			-	_	Pro		_			Pro					Val			576
Phe Phe Tile Ser Ser Ile Tyr Asp Lys Glu Gln Ser Lys Asn Asn Val  210  211  The act cga the goe as a try gat the aso they asn Leu Leu Gln Met Leu His 225  230  230  230  230  230  230  230				Glu					Lys					Val				624
Leu Leu Arg Phe Ala Lys Leu Asp Phe Asn Leu Leu Gln Met Leu His 240  aaa caa gaa ctt gct caa gta tca agg tgg gg aaa gat ttg gat ttt Lys Gln Glu Leu Ala Gln Val Ser Arg Trp Trp Lys Asp Leu Asp Phe 255  gta aca aca ctt cca tat gct aga gat cga gta gtt gaaa tgc tac ttt Val Thr Thr Leu Pro Tyr Ala Arg Asp Arg Val Val Glu Cys Tyr Phe 270  gag gca tta gga gtt tat ttt gga cct caa tac tct caa gct cgc gtc Glu Ala Arg Val 275  atg ctc gtt aag acc ata tca atg att tcg att gga gac acc ttt Met Leu Val Lys Thr Ile Ser Met Ile Ser Ile Val Asp Asp Asp Thr Phe 290  gat gct tac ggt aca gtt aaa gac ctt gag gct acc ttt 300  gat gct tac ggt aca gtt aaa gaa ctt gag gct acc ttt 300  gat gct tac ggt aca gtt aca ag acc ata tca atg att tcg att gc gat gac acc ttt 300  gat gct tac ggt aca gtt aaa gaa ctt gag gca tac aca gat gcc ata Asp Ala Tyr Gly Thr Val Lys Glu Leu Glu Ala Tyr Thr Asp Ala Ile 305  atc agg tat acc acc gat acc gat ccc ccc gat tac acc gat gclu Ala Tyr Thr Asp Ala Ile 325  atc agt tat aaa gct atc caa gat ccc acc and glu Ile Asp Arg Leu Pro Asp Tyr Met Lys 325  atc agt tat aaa gct atc caa gat ccc acc and gat ccc acc and acc gat tac acc gat acc acc acc acc acc acc acc acc acc a		ne i	Phe					Tyr					Ser					672
Lys Gln Glu Leu Ala Gln Val Ser Arg Trp Trp Lys Asp Leu Asp Phe 255 Per 245 Per 245 Per 245 Per 246 Per 246 Per Tyr Ala Arg Asp Arg Val Val Glu Cys Tyr Phe 261 Per Tyr Ala Arg Asp Arg Val Val Glu Cys Tyr Phe 270 Per 270 Per 280 Per 270 Pe	Le	eu ∶					Lys					Leu					His	720
Val Thr Thr Leu Pro Tyr Ala Arg Asp 265 Arg Val Val Glu Cys Tyr Phe 260 Regard gas atta gga gtt tat ttt gag cct caa tac tct caa gct cgc gtc Glu Ala Leu Gly Val Tyr Phe Glu Pro Gln Tyr Ser Gln Ala Arg Val 275 Ref Cln Ala Leu Gly Val Tyr Phe Glu Pro Gln Tyr Ser Gln Ala Arg Val 285 Ref Cln Ala Arg Val 275 Ref Cln Ala Arg Val 285 Ref Cln Ala Arg Asp Thr Phe 295 Ref Cln Ala Tyr Gly Thr Val Lys Glu Leu Glu Ala Tyr Thr Asp Ala Tle 305 Ref Cln Ala Tyr Thr Asp Ala Tle 315 Ref Cln Ala Tyr Thr Asp Ala Tle 320 Ref Cln Arg Trp Asp Ile Asn Glu Ile Asp Arg Leu Pro Asp Tyr Met Lys 325 Ref Cln Arg Tyr Lys Ala Ile Leu Asp Leu Tyr Lys Asp Tyr Glu Lys Glu 340 Ref Cln Arg 345 Ref Cys Ref His Ala Ile Glu Arg 355 Ref Cln Arg Arg Leu Pro Asp Tyr Met Lys Glu 360 Ref Ref Cys Ref His Ala Ile Glu Arg 355 Ref				_		Āla		-			Trp			-	_	Asp		768
Glu Ala Leu Gly Val Tyr Phe Glu Pro Gln Tyr Ser Gln Ala Arg Val 285  atg ctc gtt aag acc ata tca atg att tcg att gtc gat gac acc ttt 912  Met Leu Val Lys Thr Ile Ser Met Ile Ser Ile Val Asp Asp Thr Phe 295  gat gct tac ggt aca gtt aaa gaa ctt gag gca tac aca gat gcc ata 320  Gaa aga tgg gat atc aca gtt aaa gaa att gat cgg ctt cct gat tac atg aaa 11008  Gln Arg Trp Asp Ile Asn Glu Ile Asp Asp Asp Tyr Met Lys 335  atc agt tat aaa gct att cta gat ctt tac aag gat tat gaa aag gaa att gat Sile Val Lys Ala Ile Leu Asp Leu Tyr Lys Asp Tyr Glu Lys Glu Lys Glu Ser Tyr Lys Ala Ile Leu Asp Leu Tyr Lys Asp Tyr Glu Lys Glu Arg 355  atg tct agt gcc gga aga tct cat att gtc tgc cat gca ata gaa aga 1104  Leu Ser Ser Ala Gly Arg Ser His 360  atg aaa gaa gta gta aga att tat aat gtc gag tca aca tag gat att att gaa aga aga 1104  Met Lys Glu Val Val Arg Asn Tyr Asn Val Glu Ser Thr Trp Phe Ile 370  gaa gga tat atg cca cct gtt tct gaa tac cta agc aat gca cta gca 1200  Glu Gly Tyr Met Pro Pro Val Ser Glu Tyr Leu Ser Asn Ala Leu Ala 385  act acc aca tat tac tac ctc gcg aca aca tcg tat ttg gc atg aag 1248  Thr Thr Thr Tyr Tyr Tyr Leu Ala Thr Thr Ser Tyr Leu Gly Met Lys					Leu					Asp					Cys			816
Met Leu Val Lys Thr Ile Ser Met Ile Ser Ile Val Asp Asp Thr Phe 290    gat gct tac ggt aca gtt aaa gaa ctt gag gca tac aca gat gcc ata 960   Asp Ala Tyr Gly Thr Val Lys Glu Leu Glu Ala Tyr Thr Asp Ala Ile 320    caa aga tgg gat atc aac gaa att gat cgg ctt cct gat tac atg aaa 1008   Gln Arg Trp Asp Ile Asn Glu Ile Asp Arg Leu Pro Asp Tyr Met Lys 335    atc agt tat aaa gct att cta gat ctc tac aag gat tat gaa aag gaa   Ile Ser Tyr Lys Ala Ile Leu Asp Leu Tyr Lys Asp Tyr Glu Lys Glu Lys Glu Lys Glu Ser Ala Gly Arg Ser His Ile Val Cys His Ala Ile Glu Arg 355    atg aaa gaa gta gta gta aga att att aat gtc gag tca aca tgg ttt att   Met Lys Glu Val Val Arg Asn Tyr Asn Val Glu Ser Thr Trp Phe Ile 370    gaa gga tat atg cca cct gtt tct gaa tac cta agc aat gca cta gca   Glu Gly Tyr Met Pro Pro Val Ser Glu Tyr Leu Ser Asn Ala Leu Ala 385   The Thr Thr Tyr Tyr Tyr Leu Ala Thr Thr Ser Tyr Leu Gly Met Lys    1248	_	_	-	Leu		_			Glu					Gln	-	_	-	864
Asp Ala Tyr Gly Thr Val Lys Glu Leu Glu Ala Tyr Thr Asp Ala Ile 320  caa aga tgg gat atc aac gaa att gat cgg ctt cct gat tac atg aaa 1008 Gln Arg Trp Asp Ile Asn Glu Ile Asp Arg Leu Pro Asp Tyr Met Lys 325  atc agt tat aaa gct att cta gat ctc tac aag gat tat gaa aag gaa 1056 Ile Ser Tyr Lys Ala Ile Leu Asp Leu Tyr Lys Asp Tyr Glu Lys Glu 340  ttg tct agt gcc gga aga tct cat att gtc tgc cat gca ata gaa aga 1104  Leu Ser Ser Ala Gly Arg Ser His Ile Val Cys His Ala Ile Glu Arg 355  atg aaa gaa gta gta aga aat tat aat gtc gag tca aca tgg ttt att Met Lys Glu Val Val Arg Asn Tyr Asn Val Glu Ser Thr Trp Phe Ile 370  gaa gga tat atg cca cct gtt tct gaa tac cta agc aat gca cta gca Glu Gly Tyr Met Pro Pro Val Ser Glu Tyr Leu Ser Asn Ala Leu Ala 385  act acc aca tat tac tac ctc gcg aca aca tcg tat ttg ggc atg aag 1248 Thr Thr Thr Tyr Tyr Tyr Leu Ala Thr Thr Ser Tyr Leu Gly Met Lys		et :	Leu	_	_			Ser	_		_		Val	-	-			912
Gln Arg Trp Asp Ile Asn Glu Ile Asp Arg Leu Pro Asp Tyr Met Lys 335 atc agt tat aaa gct att cta gat ctc tac aag gat tat gaa aag gaa 1056 Ile Ser Tyr Lys Ala Ile Leu Asp Leu Tyr Lys Asp Tyr Glu Lys Glu 340 at 1 le Leu Asp Leu Tyr Lys Asp Tyr Glu Lys Glu 350 atg gcc gga aga tct cat att gtc tgc cat gca ata gaa aga Lou Ser Ser Ala Gly Arg Ser His Ile Val Cys His Ala Ile Glu Arg 355 atg gca aga att tat aat gtc gag tca aca tgg ttt att Lys Glu Val Val Arg Asn Tyr Asn Val Glu Ser Thr Trp Phe Ile 370 and gaa gga tat atg cca cct gtt tct gaa tac cta agc aat gca cta gca aca gga gga tat atg cca cct gtt tct gaa tac cta agc aat gca cta gca l200 Glu Gly Tyr Met Pro Pro Val Ser Glu Tyr Leu Ser Asn Ala Leu Ala 385 acc aca tat tac tac ctc gcg aca aca tcg tat ttg ggc atg aag l248 Thr Thr Thr Tyr Tyr Tyr Leu Ala Thr Thr Ser Tyr Leu Gly Met Lys	As	sp.					Val					Ala					Ile	960
The Ser Tyr Lys Ala Ile Leu Asp Leu Tyr Lys Asp Tyr Glu Lys Glu 340  ttg tct agt gcc gga aga tct cat att gtc tgc cat gca ata gaa aga Ll04  Leu Ser Ser Ala Gly Arg Ser His Ile Val Cys His Ala Ile Glu Arg 355  atg aaa gaa gta gta aga aat tat aat gtc gag tca aca tgg ttt att Lys Glu Val Val Arg Asn Tyr Asn Val Glu Ser Thr Trp Phe Ile 370  gaa gga tat atg cca cct gtt tct gaa tac cta agc aat gca cta gca 1200  Glu Gly Tyr Met Pro Pro Val Ser Glu Tyr Leu Ser Asn Ala Leu Ala 385  act acc aca tat tac tac ctc gcg aca aca tcg tat ttg ggc atg aag 1248  Thr Thr Thr Tyr Tyr Tyr Leu Ala Thr Thr Ser Tyr Leu Gly Met Lys						Ile					Arg					Met		1008
Leu Ser Ser Ala Gly Arg Ser His Ile Val Cys His Ala Ile Glu Arg 355 and gas gas at tat aat gtc gag tca aca tgg ttt att 2370 and Val Val Arg Asn Tyr Asn Val Glu Ser Thr Trp Phe Ile 370 and Glu Gly Tyr Met Pro Pro Val Ser Glu Tyr Leu Ser Asn Ala Leu Ala 385 and acc aca tat tac tac ctc gcg aca aca tcg tat ttg ggc atg aag Trans Ala Cys His Ala Ile Glu Arg 365 and 395 and acc aca tat tac tac ctc gcg aca aca tcg tat ttg ggc atg aag 1248 Thr Thr Thr Tyr Tyr Tyr Leu Ala Thr Thr Ser Tyr Leu Gly Met Lys					Lys					Leu					Glu			1056
Met Lys Glu Val Val Arg Asn Tyr Asn Val Glu Ser Thr Trp Phe Ile 370  gaa gga tat atg cca cct gtt tct gaa tac cta agc aat gca cta gca Glu Gly Tyr Met Pro Pro Val Ser Glu Tyr Leu Ser Asn Ala Leu Ala 385  act acc aca tat tac tac ctc gcg aca aca tcg tat ttg ggc atg aag Thr Thr Tyr Tyr Tyr Leu Ala Thr Thr Ser Tyr Leu Gly Met Lys		-		Ser	-		_		His		_	_		Ala		-	_	1104
Glu Gly Tyr Met Pro Pro Val Ser Glu Tyr Leu Ser Asn Ala Leu Ala 385 390 395 400  act acc aca tat tac tac ctc gcg aca aca tcg tat ttg ggc atg aag Thr Thr Tyr Tyr Tyr Leu Ala Thr Thr Ser Tyr Leu Gly Met Lys		et :	Lys					Asn					Ser					1152
Thr Thr Thr Tyr Tyr Tyr Leu Ala Thr Thr Ser Tyr Leu Gly Met Lys	Ğ]	lu					Pro					Leu					Ala	1200
						${\tt Tyr}$					Thr	_		_		Met	_	1248

	gcc Ala															1296
	gaa Glu															1344
	gag Glu 450															1392
_	atg Met	_	-								_	_	-			1440
	aat Asn	_	_			_			-			_				1488
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Tyr	His	Phe	Glu	Lys 85	Glu	Ile	Asp	Glu	Ile 90	Leu	Asp	Gln	Ile	Tyr 95	Asn	
Gln	Asn	Ser	Asn 100	Cys	Asn	Asp	Leu	C <b>y</b> s 105	Thr	Ser	Ala	Leu	Gln 110	Phe	Arg	
Leu	Leu	Arg 115	Gln	His	Gly	Phe	Asn 120	Ile	Ser	Pro	Glu	Ile 125	Phe	Ser	Lys	
Phe	Gln	Asp	Glu	Asn	Gly		Phe	Lys	Glu	Ser	Leu 140	Ala	Ser	Asp	Val	
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Leu 145	130 Gly	Leu	Leu	Asn	Leu 150		Glu	Ala	Ser	His 155		Arg	Thr	His	Ala 160	
145					150	Tyr				155	Val				160	

												con	tın	uea	
			180					185					190		
Ala	Leu	Glu 195	Gln	Cys	Leu	His	L <b>y</b> s 200	Gly	Val	Pro	Arg	Val 205	Glu	Thr	Arg
Phe	Phe 210	Ile	Ser	Ser	Ile	Tyr 215	Asp	Lys	Glu	Gln	Ser 220	Lys	Asn	Asn	Val
Leu 225	Leu	Arg	Phe	Ala	L <b>y</b> s 230	Leu	Asp	Phe	Asn	Leu 235	Leu	Gln	Met	Leu	His 240
Lys	Gln	Glu	Leu	Ala 245	Gln	Val	Ser	Arg	Trp 250	Trp	Lys	Asp	Leu	Asp 255	Phe
Val	Thr	Thr	Leu 260	Pro	Tyr	Ala	Arg	Asp 265	Arg	Val	Val	Glu	C <b>y</b> s 270	Tyr	Phe
Glu	Ala	Leu 275	Gly	Val	Tyr	Phe	Glu 280	Pro	Gln	Tyr	Ser	Gln 285	Ala	Arg	Val
Met	Leu 290	Val	Lys	Thr	Ile	Ser 295	Met	Ile	Ser	Ile	Val 300	Asp	Asp	Thr	Phe
Asp 305	Ala	Tyr	Gly	Thr	Val 310	Lys	Glu	Leu	Glu	Ala 315	Tyr	Thr	Asp	Ala	Ile 320
Gln	Arg	Trp	Asp	Ile 325	Asn	Glu	Ile	Asp	Arg 330	Leu	Pro	Asp	Tyr	Met 335	Lys
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Leu	Ser	Ser 355	Ala	Gly	Arg	Ser	His 360	Ile	Val	Сув	His	Ala 365	Ile	Glu	Arg
Met	L <b>y</b> s 370	Glu	Val	Val	Arg	Asn 375	Tyr	Asn	Val	Glu	Ser 380	Thr	Trp	Phe	Ile
Glu 385	Gly	Tyr	Met	Pro	Pro 390	Val	Ser	Glu	Tyr	Leu 395	Ser	Asn	Ala	Leu	Ala 400
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Cys 465	Met	Arg			Gly 470				Lys			Met		_	Phe 480
Gln	Asn	Met	Ala	Glu 485	Thr	Ala	Trp	Lys	Asp 490	Ile	Asn	Glu	Gly	Leu 495	Leu
Arg	Pro	Thr	Pro 500	Val	Ser	Thr	Glu	Phe 505	Leu	Thr	Pro	Ile	Leu 510	Asn	Leu
Ala	Arg	Ile 515	Val	Glu	Val	Thr	<b>Ty</b> r 520	Ile	His	Asn	Leu	Asp 525	Gly	Tyr	Thr
His	Pro 530	Glu	Lys	Val	Leu	L <b>y</b> s 535	Pro	His	Ile	Ile	Asn 540	Leu	Leu	Val	Asp
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Ile	Ser	Tyr	L <b>y</b> s 340	Ala	Ile	Leu	Asp	Leu 345	Tyr	Lys	Asp	Tyr	Glu 350	Lys	Glu
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Tyr	Glu 450	Val	Glu	Lys	Ser	Arg 455	Gly	Gln	Ile	Ala	Thr 460	Gly	Ile	Glu	Cys

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ttc caa gat gaa Phe Gln Asp Glu 130		s Phe Lys Gl			32
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gac gat atc tta Asp Asp Ile Leu			e Ser Thr Ile		8 !
tct gca gct cca Ser Ala Ala Pro 180	His Leu Ly				76
gcc ctt gag caa	tgt ttg ca	c aag ggt gt	t cct aga gtc	gag acc cga 62	24

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Ala	Leu	Glu 195	Gln	Cys	Leu	His	L <b>y</b> s 200	Gly	Val	Pro	Arg	Val 205	Glu	Thr	Arg	
	ttc Phe 210															672
	ctt					ttg					ctc					720
Leu 225	Leu	Arg	Phe	Ala	Lys 230	Leu	Asp	Phe	Asn	Leu 235	Leu	GIn	Met	Leu	H1s 240	
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	aca Thr		Leu					Asp					Cys			816
tgg	gca	tta	260 gga	gtt	tat	ttt	gag	265 cct	caa	tac	tct	caa	270 gct	cgc	gtc	864
Trp	Ala	Leu 275	Gly	Val	Tyr	Phe	Glu 280	Pro	Gln	Tyr	Ser	Gln 285	Ala	Arg	Val	
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-	gct Ala				-		-			_			-	-		960
	aga Arg															1008
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Ser	Ile	Asp 35	Asn	Gln	Val	Ala	Glu 40	Lys	Tyr	Ala	Gln	Glu 45	Ile	Glu	Ala	
	L <b>y</b> s 50					55					60			_		
A1a 65	Asp	Thr	Leu	Asn	Leu 70	Ile	Asp	Ile	Ile	GIu 75	Arg	Leu	Gly	Ile	Ser 80	
Tyr	His	Phe	Glu	Lys 85	Glu	Ile	Asp	Glu	Ile 90	Leu	Asp	Gln	Ile	Tyr 95	Asn	
Gln	Asn	Ser	Asn 100	Cys	Asn	Asp	Leu	C <b>y</b> s 105	Thr	Ser	Ala	Leu	Gln 110	Phe	Arg	
Leu	Leu	Arg 115	Gln	His	Gly	Phe	Asn 120	Ile	Ser	Pro	Glu	Ile 125	Phe	Ser	Lys	
Phe	Gln 130	Asp	Glu	Asn	Gly	Lys 135	Phe	Lys	Glu	Ser	Leu 140	Ala	Ser	Asp	Val	
Leu 145	Gly	Leu	Leu	Asn	Leu 150	Tyr	Glu	Ala	Ser	His 155	Val	Arg	Thr	His	Ala 160	
Asp	Asp	Ile	Leu	Glu 165	Asp	Ala	Leu	Ala	Phe 170	Ser	Thr	Ile	His	Leu 175	Glu	
Ser	Ala	Ala	Pro 180	His	Leu	Lys	Ser	Pro 185	Leu	Arg	Glu	Gln	Val 190	Thr	His	
Ala	Leu	Glu 195	Gln	Cys	Leu	His	<b>Lys</b> 200	Gly	Val	Pro	Arg	Val 205	Glu	Thr	Arg	
Phe	Phe 210	Ile	Ser	Ser	Ile	Tyr 215	Asp	Lys	Glu	Gln	Ser 220	Lys	Asn	Asn	Val	
Leu 225	Leu	Arg	Phe	Ala	L <b>y</b> s 230	Leu	Asp	Phe	Asn	Leu 235	Leu	Gln	Met	Leu	His 240	
Lys	Gln	Glu	Leu	Ala 245	Gln	Val	Ser	Arg	Trp 250	Trp	Lys	Asp	Leu	Asp 255	Phe	
Val	Thr	Thr	Leu 260	Pro	Tyr	Ala	Arg	Asp 265	Arg	Val	Val	Glu	C <b>y</b> s 270	Tyr	Phe	
Trp	Ala	Leu 275	Gly	Val	Tyr	Phe	Glu 280	Pro	Gln	Tyr	Ser	Gln 285	Ala	Arg	Val	
Met	Leu 290	Val	Lys	Thr	Ile	Ser 295	Met	Ile	Ser	Ile	Val 300	Asp	Asp	Thr	Phe	
Asp 305	Ala	Tyr	Gly	Thr	Val 310	Lys	Glu	Leu	Glu	Ala 315	Tyr	Thr	Asp	Ala	Ile 320	

Gln	Arg	Trp	Asp	Ile 325	Asn	Glu	Ile	Asp	Arg 330	Leu	Pro	Asp	Tyr	Met 335	Lys	
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Leu	Ser	Ser 355	Ala	Gly	Arg	Ser	His 360	Ile	Val	Cys	His	Ala 365	Ile	Glu	Arg	
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Ser	Ala	Thr	Glu 420	Gln	Asp	Phe	Glu	Trp 425	Leu	Ser	Lys	Asn	Pro 430	Lys	Ile	
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Tyr	Glu 450	Val	Glu	Lys	Ser	Arg 455	Gly	Gln	Ile	Ala	Thr 460	Gly	Ile	Glu	Cys	
C <b>y</b> s 465	Met	Arg	Ąsp	Tyr	Gly 470	Ile	Ser	Thr	Lys	Glu 475	Ala	Met	Ala	Lys	Phe 480	
Gln	Asn	Met	Ala	Glu 485	Thr	Ala	Trp	Lys	Asp 490	Ile	Asn	Glu	Gly	Leu 495	Leu	
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Ala	Arg	Ile 515	Val	Glu	Val	Thr	<b>Tyr</b> 520	Ile	His	Asn	Leu	Asp 525	Gly	Phe	Thr	
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tac																288

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	ctc Leu															384
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	gga Gly															480
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	gca Ala	_			_											576
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	ctt Leu															720
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	gct Ala															960
	aga Arg															1008
	agt Ser															1056
	tct Ser															1104
	aaa Lys 370															1152
	gga Gly															1200
act	acc	aca	tat	tac	tac	ctc	gcg	aca	aca	tcg	tat	ttg	ggc	atg	aag	1248

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Ser	Ile	Asp 35	Asn	Gln	Val	Ala	Glu 40	Lys	Tyr	Ala	Gln	Glu 45	Ile	Glu	Ala		
	<b>Lys</b> 50					55					60						
65	Asp				70		-			75	•		-		80		
	His			85					90		_			95			
	Asn		100	_				105					110		-		
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Leu 145		Asp Leu	Leu	Asn	Leu 150	135 Tyr	Glu	Ala	Ser	His 155	140 Val	Arg	Thr	His	Ala 160		

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Ala	Leu	Glu 195	Gln	Cys	Leu	His	L <b>y</b> s 200	Gly	Val	Pro	Arg	Val 205	Glu	Thr	Arg
Phe	Phe 210	Ile	Ser	Ser	Ile	Tyr 215	Asp	Lys	Glu	Gln	Ser 220	Lys	Asn	Asn	Val
Leu 225	Leu	Arg	Phe	Ala	Lys 230	Leu	Asp	Phe	Asn	Leu 235	Leu	Gln	Met	Leu	His 240
Lys	Gln	Glu	Leu	Ala 245	Gln	Val	Ser	Arg	Trp 250	Trp	Lys	Asp	Leu	Asp 255	Phe
Val	Thr	Thr	Leu 260	Pro	Tyr	Ala	Arg	Asp 265	Arg	Val	Val	Glu	C <b>y</b> s 270	Tyr	Phe
Ser	Ala	Leu 275	Gly	Val	Tyr	Phe	Glu 280	Pro	Gln	Tyr	Ser	Gln 285	Ala	Arg	Val
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Gln	Arg	Trp	Asp	Ile 325	Asn	Glu	Ile	Asp	Arg 330	Leu	Pro	Asp	Tyr	Met 335	Lys
Ile	Ser	Tyr	L <b>y</b> s 340	Ala	Ile	Leu	Asp	Leu 345	Tyr	Lys	Asp	Tyr	Glu 350	Lys	Glu
Leu	Ser	Ser 355	Ala	Gly	Arg	Ser	His 360	Ile	Val	Сув	His	Ala 365	Ile	Glu	Arg
Met	Lys 370	Glu	Val	Val	Arg	Asn 375	Tyr	Asn	Val	Glu	Ser 380	Thr	Trp	Phe	Ile
Glu 385	Gly	Tyr	Met	Pro	Pro 390	Val	Ser	Glu	Tyr	Leu 395	Ser	Asn	Ala	Leu	Ala 400
Thr	Thr	Thr	Tyr	Tyr 405	Tyr	Leu	Ala	Thr	Thr 410	Ser	Tyr	Leu	Gly	Met 415	Lys
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				485	Thr			_	490					495	
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		515			Val		520					525			
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														tac Tyr 95		288
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-	-			-	-	_		-						ctt Leu 175	-	528
	_	-			_						-			aca Thr		576
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		_		_		_	-			_		_	_	ttg Leu		720
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												gat Asp			912	
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Ser	Ile	Asp 35	Asn	Gln	Val	Ala	Glu 40	Lys	Tyr	Ala	Gln	Glu 45	Ile	Glu	Ala
Leu	<b>Ly</b> s 50	Glu	Gln	Thr	Arg	Ser 55	Met	Leu	Leu	Ala	Thr 60	Gly	Arg	Lys	Leu
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Tyr	His	Phe	Glu	Lys 85	Glu	Ile	Asp	Glu	Ile 90	Leu	Asp	Gln	Ile	<b>Ty</b> r 95	Asn
Gln	Asn	Ser	Asn 100	Сув	Asn	Asp	Leu	C <b>y</b> s 105	Thr	Ser	Ala	Leu	Gln 110	Phe	Arg
Leu	Leu	Arg 115	Gln	His	Gly	Phe	Asn 120	Ile	Ser	Pro	Glu	Ile 125	Phe	Ser	Lys
Phe	Gln 130	Asp	Glu	Asn	Gly	Lys 135	Phe	Lys	Glu	Ser	Leu 140	Ala	Ser	Asp	Val
Leu 145	Gly	Leu	Leu	Asn	Leu 150	Tyr	Glu	Ala	Ser	His 155	Val	Arg	Thr	His	Ala 160
Asp	Asp	Ile	Leu	Glu 165	Asp	Ala	Leu	Ala	Phe 170	Ser	Thr	Ile	His	Leu 175	Glu
Ser	Ala	Ala	Pro 180	His	Leu	Lys	Ser	Pro 185	Leu	Arg	Glu	Gln	Val 190	Thr	His
Ala	Leu	Glu 195	Gln	Сув	Leu	His	L <b>y</b> s 200	Gly	Val	Pro	Arg	Val 205	Glu	Thr	Arg
Phe	Phe 210	Ile	Ser	Ser	Ile	Tyr 215	Asp	Lys	Glu	Gln	Ser 220	Lys	Asn	Asn	Val
Leu 225	Leu	Arg	Phe	Ala	L <b>y</b> s 230	Leu	Asp	Phe	Asn	Leu 235	Leu	Gln	Met	Leu	His 240
Lys	Gln	Glu	Leu	Ala 245	Gln	Val	Ser	Arg	Trp 250	Trp	Lys	Asp	Leu	<b>Asp</b> 255	Phe
Val	Thr	Thr	Leu 260	Pro	Tyr	Ala	Arg	Asp 265	Arg	Val	Val	Glu	C <b>y</b> s 270	Tyr	Phe
Trp	Ala	Leu 275	Gly	Val	Tyr	Phe	Glu 280	Pro	Gln	Tyr	Ser	Gln 285	Ala	Arg	Val
Met	Leu 290	Val	Lys	Thr	Ile	Ser 295	Met	Ile	Ser	Ile	Val 300	Asp	Asp	Thr	Phe
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Gln	Arg	Trp	Asp	Ile 325	Asn	Glu	Ile	Asp	Arg 330	Leu	Pro	Asp	Tyr	Met 335	Lys
Ile	Ser	Tyr	Lys 340	Ala	Ile	Leu	Asp	Leu 345	Tyr	Lys	Asp	Tyr	Glu 350	Lys	Glu
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Met	L <b>y</b> s 370	Glu	Val	Val	Arg	Asn 375	Tyr	Asn	Val	Glu	Ser 380	Thr	Trp	Phe	Ile
Glu 385	Gly	Tyr	Met	Pro	Pro 390	Val	Ser	Glu	Tyr	Leu 395	Ser	Asn	Ala	Leu	Ala 400
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	gga atg agt agg cga ggg aaa tct atc act cct Gly Met Ser Arg Arg Gly Lys Ser Ile Thr Pro 40 45	146
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	ıe					att Ile										722
_	la	-				acc Thr 245				_	-	_		_		770
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						ttt Phe										1442

Cys GJy His Arg Ile Ser Ala Leu Gln Pro Ile Leu Thr Met Asp Ile 495  ccc ttt cct gat cat atc ctc aag gaa gtt gac ttc cca tca aag ctt Pro Phe Pro Asp His Ile Leu Lys Glu Val Asp Phe Pro Ser Lys Leu 500  aac gac ttg gca tgt gcc atc ctt cga tta cga ggt gat acg cgg tgc Asn Asp Leu Ala Cys Ala Ile Leu Arg Leu Arg Gly Asp Thr Arg Cys 515  tac aag gcg gac agg gct cgt gga gaa gaa gct tcc tct ata tca tgt Tyr Lys Ala Asp Arg Ala Arg Gly Glu Glu Ala Ser Ser Ile Ser Cys 530  tat atg aaa gac aat cct gga gta tca gag gaa gat gct ctc gat cat Tyr Met Lys Asp Asn Pro Gly Val Ser Glu Glu Asp Ala Leu Asp His 555  atc aac gcc atg atc agt gac gta atc aac gga gta tca aat tgg gaa ctt Ile Asn Ala Met Ile Ser Asp Val Ile Lys Gly Leu Asn Trp Glu Leu 560  ctc aaa cca gac atc aat gtt ccc atc tcg gcg aag aaa cat gct ttt Leu Lys Pro Asp Ile Asn Val Pro Ile Ser Ala Lys Lys His Ala Phe 590	1490 1538 1586 1634 1682
Pro Phe Pro Asp His Ile Leu Lys Glu Val Asp Phe Pro Ser Lys Leu 510  aac gac ttg gca tgt gcc atc ctt cga tta cga ggt gat acg cgg tgc Asn Asp Leu Ala Cys Ala Ile Leu Arg Leu Arg Gly Asp Thr Arg Cys 525  tac aag gcg gac agg gct cgt gga gaa gaa gct tcc tct ata tca tgt Tyr Lys Ala Asp Arg Ala Arg Gly 535  tat atg aaa gac aat cct gga gta tca gag gaa gat gct ctc gat cat Tyr Met Lys Asp Asn Pro Gly Val Ser Glu Glu Ala Ser Ser Ile Ser Cys 540  atc aac gcc atg atc agt gac gta atc aag gga gaa gat gct ctc gat cat Tyr Met Lys Asp Asn Pro Gly Val Ser Glu Glu Asp Ala Leu Asp His 545  atc aac gcc atg atc agt gac gta atc aag gga gta tta aat tgg gaa ctt Ile Asn Ala Met Ile Ser Asp Val Ile Lys Gly Leu Asn Trp Glu Leu 560  ctc aaa cca gac atc aat gtt ccc atc tcg gcg aag aaa cat gct ttt Leu Lys Pro Asp Ile Asn Val Pro Ile Ser Ala Lys Lys His Ala Phe 580  gac atc gcc aga gct ttc cat tac ggc tac aaa tac cga gac ggc tac Asp Ile Ala Arg Ala Phe His Tyr Gly Tyr Lys Tyr Arg Asp Gly Tyr	1586 1634 1682
Asn Asp Leu Ala Cys Ala Ile Leu Arg Leu Arg Gly Asp Thr Arg Cys 515  tac aag gcg gac agg gct cgt gga gaa gaa gct tcc tct ata tca tgt Tyr Lys Ala Asp Arg Ala Arg Gly Glu Glu Ala Ser Ser Ile Ser Cys 530  tat atg aaa gac aat cct gga gta tca gag gaa gat gct ctc gat cat Tyr Met Lys Asp Asn Pro Gly Val Ser Glu Glu Asp Ala Leu Asp His 545  atc aac gcc atg atc agt gac gta atc aaa gga tta aat tgg gaa ctt Ile Asn Ala Met Ile Ser Asp Val Ile Lys Gly Leu Asn Trp Glu Leu 560  ctc aaa cca gac atc aat gtt ccc atc tcg gcg aag aaa cat gct ttt Leu Lys Pro Asp Ile Asn Val Pro Ile Ser Ala Lys Lys His Ala Phe 580  gac atc gcc aga gct ttc cat tac ggc tac aaa tac cga gac ggc tac Asp Ile Ala Arg Ala Phe His Tyr Gly Tyr Lys Tyr Arg Asp Gly Tyr	1634 1682
Tyr Lys Ala Asp Arg Ala Arg Gly Glu Glu Ala Ser Ser Ile Ser Cys 530    tat atg aaa gac aat cct gga gta tca gag gaa gat gct ctc gat cat Tyr Met Lys Asp Asn Pro Gly Val Ser Glu Glu Asp Ala Leu Asp His 545    atc aac gcc atg atc agt gac gta atc aaa gga tta aat tgg gaa ctt Ile Asn Ala Met Ile Ser Asp Val Ile Lys Gly Leu Asn Trp Glu Leu 575    ctc aaa cca gac atc aat gtt ccc atc tcg gcg aag aaa cat gct ttt Leu Lys Pro Asp Ile Asn Val Pro Ile Ser Ala Lys Lys His Ala Phe 580    gac atc gcc aga gct ttc cat tac ggc tac aaa tac cga gac ggc tac Asp Ile Ala Arg Ala Phe His Tyr Gly Tyr Lys Tyr Arg Asp Gly Tyr	1682
Tyr Met Lys Asp Asn Pro Gly Val Ser Glu Glu Asp Ala Leu Asp His 545  atc aac gcc atg atc agt gac gta atc aaa gga tta aat tgg gaa ctt Ile Asn Ala Met Ile Ser Asp Val Ile Lys Gly Leu Asn Trp Glu Leu 560  ctc aaa cca gac atc aat gtt ccc atc tcg gcg aag aaa cat gct ttt Leu Lys Pro Asp Ile Asn Val Pro Ile Ser Ala Lys Lys His Ala Phe 580  gac atc gcc aga gct ttc cat tac ggc tac aaa tac cga gac ggc tac Asp Ile Ala Arg Ala Phe His Tyr Gly Tyr Lys Tyr Arg Asp Gly Tyr	
Ile Asn Ala Met Ile Ser Asp Val Ile Lys Gly Leu Asn Trp Glu Leu 575  ctc aaa cca gac atc aat gtt ccc atc tcg gcg aag aaa cat gct ttt Leu Lys Pro Asp Ile Asn Val Pro Ile Ser Ala Lys Lys His Ala Phe 580  gac atc gcc aga gct ttc cat tac ggc tac aaa tac cga gac ggc tac Asp Ile Ala Arg Ala Phe His Tyr Gly Tyr Lys Tyr Arg Asp Gly Tyr	1730
Leu Lys Pro Asp Ile Asn Val Pro Ile Ser Ala Lys Lys His Ala Phe 580 580 580 590  gac atc gcc aga gct ttc cat tac ggc tac aaa tac cga gac ggc tac Asp Ile Ala Arg Ala Phe His Tyr Gly Tyr Lys Tyr Arg Asp Gly Tyr	
Asp Ile Ala Arg Ala Phe His Tyr Gly Tyr Lys Tyr Arg Asp Gly Tyr	1778
	1826
agc gtt gcc aac gtt gaa acg aag agt ttg gtc acg aga acc ctc ctt Ser Val Ala Asn Val Glu Thr Lys Ser Leu Val Thr Arg Thr Leu Leu 610 615 620	1874
gaa tot gtg oot ttg tag caacagotca aatotatgoo ctatgotatg Glu Ser Val Pro Leu 625	1922
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taataaagtt gtaatttaaa aaaaaaaaa aaaaaa 20	2018
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Glu 145	Ile	Lys	Ser	Ala	Leu 150	Asp	Tyr	Val	Tyr	Ser 155	Tyr	Trp	Gly	Glu	Asn 160
Gly	Ile	Gly	Суѕ	Gl <b>y</b> 165	Arg	Glu	Ser	Val	Val 170	Thr	Asp	Leu	Asn	Ser 175	Thr
Ala	Leu	Gly	Leu 180	Arg	Thr	Leu	Arg	Leu 185	His	Gly	Tyr	Pro	Val 190	Ser	Ser
Asp	Val	Phe 195	Lys	Ala	Phe	Lys	Gly 200	Gln	Asn	Gly	Gln	Phe 205	Ser	Суѕ	Ser
Glu	Asn 210	Ile	Gln	Thr	Asp	Glu 215	Glu	Ile	Arg	Gly	Val 220	Leu	Asn	Leu	Phe
Arg 225	Ala	Ser	Leu	Ile	Ala 230	Phe	Pro	Gly	Glu	L <b>y</b> s 235	Ile	Met	Asp	Glu	Ala 240
Glu	Ile	Phe	Ser	Thr 245	Lys	Tyr	Leu	Lys	Glu 250	Ala	Leu	Gln	Lys	Ile 255	Pro
Val	Ser	Ser	Leu 260	Ser	Arg	Glu	Ile	Gly 265	Asp	Val	Leu	Glu	Tyr 270	Gly	Trp
His	Thr	<b>Ty</b> r 275	Leu	Pro	Arg	Leu	Glu 280	Ala	Arg	Asn	Tyr	Ile 285	Gln	Val	Phe
Gly	Gln 290	Asp	Thr	Glu	Asn	Thr 295	Lys	Ser	Tyr	Val	L <b>y</b> s 300	Ser	Lys	Lys	Leu
Leu 305	Glu	Leu	Ala	Lys	Leu 310	Glu	Phe	Asn	Ile	Phe 315	Gln	Ser	Leu	Gln	<b>Lys</b> 320
Arg	Glu	Leu	Glu	Ser 325	Leu	Val	Arg	Trp	Trp 330	Lys	Glu	Ser	Gly	Phe 335	Pro
Glu	Met	Thr	Phe 340	Суѕ	Arg	His	Arg	His 345	Val	Glu	Tyr	Tyr	Thr 350	Leu	Ala
Ser	Сув	Ile 355	Ala	Phe	Glu	Pro	Gln 360	His	Ser	Gly	Phe	Arg 365	Leu	Gly	Phe
Ala	L <b>y</b> s 370	Thr	Суѕ	His	Leu	Ile 375	Thr	Val	Leu	Asp	Asp 380	Met	Tyr	Asp	Thr
Phe 385	Gly	Thr	Val	Asp	Glu 390	Leu	Glu	Leu	Phe	Thr 395	Ala	Thr	Met	Lys	Arg 400
Trp	Asp	Pro	Ser	Ser 405	Ile	Asp	Суѕ	Leu	Pro 410	Glu	Tyr	Met	Lys	Gly 415	Val
Tyr	Ile	Ala	Val 420	Tyr	Asp	Thr	Val	Asn 425	Glu	Met	Ala	Arg	Glu 430	Ala	Glu
Glu	Ala		Gly			Thr							Ala	Trp	Glu
Ala	<b>Ty</b> r 450	Ile	Asp	Ser	Tyr	Met 455	Gln	Glu	Ala	Arg	Trp 460	Ile	Ala	Thr	Gly
<b>Ty</b> r 465	Leu	Pro	Ser	Phe	Asp 470	Glu	Tyr	Tyr	Glu	Asn 475	Gly	Lys	Val	Ser	C <b>y</b> s 480
Gly	His	Arg	Ile	Ser 485	Ala	Leu	Gln	Pro	Ile 490	Leu	Thr	Met	Asp	Ile 495	Pro
Phe	Pro	Asp	His 500	Ile	Leu	Lys	Glu	Val 505	Asp	Phe	Pro	Ser	L <b>y</b> s 510	Leu	Asn
Asp	Leu	Ala 515	Суѕ	Ala	Ile	Leu	Arg 520	Leu	Arg	Gly	Asp	Thr 525	Arg	Суѕ	Tyr
Lys	Ala 530	Asp	Arg	Ala	Arg	Gl <b>y</b> 535	Glu	Glu	Ala	Ser	Ser 540	Ile	Ser	Суѕ	Tyr
Met 545	Lys	Asp	Asn	Pro	Gly 550	Val	Ser	Glu	Glu	<b>A</b> sp 555	Ala	Leu	Asp	His	Ile 560

Asn	Ala	Met	Ile	Ser 565	Asp	Val	Ile	Lys	Gl <b>y</b> 570	Leu	Asn	Trp	Glu	Leu 575	Leu	
Lys	Pro	Asp	Ile 580	Asn	Val	Pro	Ile	Ser 585		Lys	Lys	His	Ala 590	Phe	Asp	
Ile	Ala	Arg 595	Ala	Phe	His	Tyr	Gly 600	Tyr	Lys	Tyr	Arg	Asp 605	Gly	Tyr	Ser	
Val	Ala 610	Asn	Val	Glu	Thr	L <b>y</b> s 615	Ser	Leu	Val	Thr	Arg 620	Thr	Leu	Leu	Glu	
Ser 625	Val	Pro	Leu													
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			ICE:						-							
agaq	gagag	gag a	aggaa	agga	aa g	attaa								agt o Ser V		52
-			_				-			_		-		caa Gln		100
														agt Ser		148
														act Thr 55		196
-	_	_								_		_	-	aac Asn		244
														att Ile		292
-			_	_		_	Val	_	_	_	-			gaa Glu	_	340
														atg Met		388
														tct Ser 135		436
														gaa Glu		484
-							-					_	-	cat His		532
														gag Glu		580
gag			_	-		_	-	-	acc Thr	_		_	_	caa	_	628

_													con	tin	ued			
						ctg Leu											676	
						acc Thr											724	
						ctt Leu											772	
	ro					att Ile											820	
Tı						ccc Pro 270											868	
			_			att Ile	-		_				-				916	
						tgg Trp											964	
						ctg Leu											1012	
				-	-	cat His		-				-	-				1060	
As						gtg Val 350											1108	
		-	_		-	caa Gln			_			-	_		-		1156	
				_		ctt Leu		-		_		_	_			-	1204	
						gat Asp											1252	
						ccc Pro					Ser		Val				1300	
As						gag Glu 430											1348	
						ttg Leu											1396	
						ata Ile											1444	
						ttg Leu											1492	
						ctt Leu											1540	
Vá						gtg Val 510											1588	

tac aat qca tcq qaq qcq qaq qcq cqq aaq cac qtq aaa tqq ctq ata	
Tyr Asn Ala Ser Glu Ala Glu Ala Arg Lys His Val Lys Trp Leu Ile 525 530 535	1636
gcg gag gtg tgg aag aag atg aat gcg gag agg gtg tcg aag gat tct Ala Glu Val Trp Lys Lys Met Asn Ala Glu Arg Val Ser Lys Asp Ser 540 545 550	1684
cca ttc ggc aaa gat ttt ata gga tgt gca gtt gat tta gga agg atg Pro Phe Gly Lys Asp Phe Ile Gly Cys Ala Val Asp Leu Gly Arg Met 555 560 565	1732
gcg cag ttg atg tac cat aat gga gat ggg cac ggc aca caa cac cct Ala Gln Leu Met Tyr His Asn Gly Asp Gly His Gly Thr Gln His Pro 570 575 580	1780
att ata cat caa caa atg acc aga acc tta ttc gag ccc ttt gca tga Ile Ile His Gln Gln Met Thr Arg Thr Leu Phe Glu Pro Phe Ala 585 590 595	1828
gagatgatga cgagccatcg tttacttact taaattctac caaagttttt cgaaggcata	1888
gttcgtaatt tttcaagcac caataaataa ggagaatcgg ctcaaacaaa cgtggcattt	1948
gccaccacgt gagcacaagg gagagtctgt cgtcgtttat ggatgaacta ttcaattttt	2008
atgcatgtaa taattaagtt caagttcaag agccttctgc atatttaact atgtatttga	2068
atttatcgag tgtgattttc tgtctttggc aacatatatt tttgtcatat gtggcatctt	2128
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1 5 10 15  Asn Leu Thr Thr Cys Leu Gln Pro Ser His Phe Lys Ser Ser Pro Lys 25  Leu Leu Ser Ser Thr Asn Ser Ser Ser Arg Ser Arg Leu Arg Val Tyr	
1 5 10 15  Asn Leu Thr Thr Cys Leu Gln Pro Ser His Phe Lys Ser Ser Pro Lys 30  Leu Leu Ser Ser Thr Asn Ser Ser Ser Ser Arg Ser Arg Leu Arg Val Tyr 45  Cys Ser Ser Ser Gln Leu Thr Thr Glu Arg Arg Ser Gly Asn Tyr Asn	
Asn Leu Thr Thr Cys Leu Gln Pro Ser His Phe Lys Ser Ser Pro Lys 30    Leu Leu Ser Ser Thr Asn Ser Ser Ser Arg Ser Arg Leu Arg Val Tyr 45    Cys Ser Ser Ser Gln Leu Thr Thr Glu Arg Arg Ser Gly Asn Tyr Asn Fro Ser Arg Trp Asp Val Asn Phe Ile Gln Ser Leu Leu Ser Asp Tyr	
Asn Leu Thr Thr Cys Leu Gln Pro Ser His Phe Lys Ser Ser Pro Lys 30 Leu Leu Leu Ser Ser Thr Asn Ser Ser Ser Arg Ser Gly Asn Tyr Asn Ser Ser Arg Trp Asp Val Asn Phe Ile Gln Ser Leu Leu Ser Asp Tyr Robert Ser Glu Asp Lys His Val Ile Arg Ala Ser Glu Leu Val Thr Leu Val	
1	
1	
10 15  Asn Leu Thr Thr Cys Leu Gln Pro 25 His Phe Lys Ser Ser Pro Lys 30  Leu Leu Ser Ser Thr Asn Ser Ser Ser Arg Ser Arg Leu Arg Val Tyr Asn 55 Ser Ser Ser Gln Leu Thr Thr Glu Arg Arg Ser Gly Asn Tyr Asn 55 Ser Arg Trp Asp Val Asn Phe Ile Gln Ser Leu Leu Ser Asp Tyr 80  Lys Glu Asp Lys His Val Ile Arg Ala Ser Glu Leu Val Thr Leu Val Ser Ser Ser Ser Ser Ser Ser Ser Thr Asp 105 Thr Ser Leu Leu Ser Asp Tyr 80  Lys Met Glu Leu Glu Lys Glu Thr Asp Gln Ile Arg Gln Leu Val Thr Leu Val 110  Leu Leu Ser Ser Ser Ser Ser Ile Tyr Leu Asp His His Tyr Tyr Lys	
1	
1	

Glu	Gly 210	Glu	Thr	Thr	Leu	Glu 215	Ser	Ala	Arg	Glu	Phe 220	Ala	Thr	Lys	Phe
Leu 225	Glu	Glu	Lys	Val	Asn 230	Glu	Gly	Gly	Val	Asp 235	Gly	Asp	Leu	Leu	Thr 240
Arg	Ile	Ala	Tyr	Ser 245	Leu	Asp	Ile	Pro	Leu 250	His	Trp	Arg	Ile	<b>Lys</b> 255	Arg
Pro	Asn	Ala	Pro 260	Val	Trp	Ile	Glu	Trp 265	Tyr	Arg	Lys	Arg	Pro 270	Asp	Met
Asn	Pro	Val 275	Val	Leu	Glu	Leu	Ala 280	Ile	Leu	Asp	Leu	Asn 285	Ile	Val	Gln
Ala	Gln 290	Phe	Gln	Glu	Glu	Leu 295	Lys	Glu	Ser	Phe	Arg 300	Trp	Trp	Arg	Asn
Thr 305	Gly	Phe	Val	Glu	Lys 310	Leu	Pro	Phe	Ala	Arg 315	Asp	Arg	Leu	Val	Glu 320
Суѕ	Tyr	Phe	Trp	Asn 325	Thr	Gly	Ile	Ile	Glu 330	Pro	Arg	Gln	His	Ala 335	Ser
Ala	Arg	Ile	Met 340	Met	Gly	Lys	Val	Asn 345	Ala	Leu	Ile	Thr	Val 350	Ile	Asp
Asp	Ile	Tyr 355	Asp	Val	Tyr	Gly	Thr 360	Leu	Glu	Glu	Leu	Glu 365	Gln	Phe	Thr
Asp	Leu 370	Ile	Arg	Arg	Trp	Asp 375	Ile	Asn	Ser	Ile	Asp 380	Gln	Leu	Pro	Asp
Tyr 385	Met	Gln	Leu	Cys	Phe 390	Leu	Ala	Leu	Asn	Asn 395	Phe	Val	Asp	Asp	Thr 400
Ser	Tyr	Asp	Val	Met 405	Lys	Glu	Lys	Gly	Val 410	Asn	Val	Ile	Pro	<b>Ty</b> r 415	Leu
Arg	Gln	Ser	Trp 420	Val	Asp	Leu	Ala	Asp 425	Lys	Tyr	Met	Val	Glu 430	Ala	Arg
Trp	Phe	Tyr 435	Gly	Gly	His	Lys	Pro 440	Ser	Leu	Glu	Glu	Tyr 445	Leu	Glu	Asn
Ser	Trp 450	Gln	Ser	Ile	Ser	Gly 455	Pro	Сув	Met	Leu	Thr 460	His	Ile	Phe	Phe
Arg 465	Val	Thr	Asp	Ser	Phe 470	Thr	Lys	Glu	Thr	Val 475	Asp	Ser	Leu	Tyr	L <b>y</b> s 480
-		_		485	Arg	_			490			-		495	Ī
Asp	Leu	Gly	Thr 500	Ser	Val	Glu	Glu	Val 505	Ser	Arg	Gly	Asp	Val 510	Pro	Lys
Ser	Leu	Gln 515	Cys	Tyr	Met	Ser	Asp 520	Tyr	Asn	Ala	Ser	Glu 525	Ala	Glu	Ala
Arg	<b>Ly</b> s 530	His	Val	Lys	Trp	Leu 535	Ile	Ala	Glu	Val	Trp 540	Lys	Lys	Met	Asn
Ala 545	Glu	Arg	Val	Ser	L <b>y</b> s 550	Asp	Ser	Pro	Phe	Gly 555	Lys	Asp	Phe	Ile	Gl <b>y</b> 560
Cys	Ala	Val	Asp	Leu 565	Gly	Arg	Met	Ala	Gln 570	Leu	Met	Tyr	His	Asn 575	Gly
Asp	Gly	His	Gly 580	Thr	Gln	His	Pro	Ile 585	Ile	His	Gln	Gln	Met 590	Thr	Arg
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<210> SEQ ID NO 23 <211> LENGTH: 1967

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		agt Ser														147
		caa Gln	_								-	-				195
		cct Pro														243
		aag L <b>y</b> s 80														291
		aat Asn														339
		att Ile														387
		atc Ile														435
		gaa Glu														483
		tac Tyr 160														531
		aag Lys														579
		tta Leu														627
-		cta Leu			-	-	-		-				-			675
		cta Leu														723
		gag Glu 240														771
		att Ile														819
	-	cta Leu	-		_	-			_	_		_				867

270 275 280 285	
caa gag ctc aaa gag gcc tct agg tgg tgg aat agt acg ggt ctt gtc Gln Glu Leu Lys Glu Ala Ser Arg Trp Trp Asn Ser Thr Gly Leu Val 290 295 300	915
cac gag ctt ccc ttt gtg aga gat agg att gtg gaa tgc tac tac tgg His Glu Leu Pro Phe Val Arg Asp Arg Ile Val Glu Cys Tyr Tyr Trp 305 310 315	963
acg aca gga gtg gtt gag cgt cgt gaa cat gga tac gag agg ata atg Thr Thr Gly Val Val Glu Arg Arg Glu His Gly Tyr Glu Arg Ile Met 320 325 330	1011
ctc acc aaa ata aat gct ctt gtt aca aca ata gac gat gtc ttt gat Leu Thr Lys Ile Asn Ala Leu Val Thr Thr Ile Asp Asp Val Phe Asp 335 340 345	1059
att tat ggt acg ctt gaa gag cta caa cta ttc aca act gct att caa Ile Tyr Gly Thr Leu Glu Glu Leu Gln Leu Phe Thr Thr Ala Ile Gln 350 360 365	1107
aga tgg gat att gaa tca atg aag caa ctc cct cct tac atg caa ata Arg Trp Asp Ile Glu Ser Met Lys Gln Leu Pro Pro Tyr Met Gln Ile 370 375 380	1155
tgt tat ctt gct ctc ttc aac ttt gtg aat gag atg gct tat gat act Cys Tyr Leu Ala Leu Phe Asn Phe Val Asn Glu Met Ala Tyr Asp Thr 385 390 395	1203
ctt agg gat aaa ggt ttc aac tcc acc cca tat cta cga aaa gcg tgg Leu Arg Asp Lys Gly Phe Asn Ser Thr Pro Tyr Leu Arg Lys Ala Trp 400 405 410	1251
gtt gat ttg gtt gag tca tat cta ata gag gca aag tgg tac tac atg Val Asp Leu Val Glu Ser Tyr Leu Ile Glu Ala Lys Trp Tyr Tyr Met 415 420 425	1299
gga cat aaa cct agt ttg gaa gaa tat atg aag aat agt tgg ata tca Gly His Lys Pro Ser Leu Glu Glu Tyr Met Lys Asn Ser Trp Ile Ser 430 435 440 445	1347
atc gga ggc atc ccc att cta tct cat cta ttt ttc cgg cta aca gat Ile Gly Gly Ile Pro Ile Leu Ser His Leu Phe Phe Arg Leu Thr Asp 450 455 460	1395
tcg att gag gaa gag gat gct gag agt atg cat aaa tac cat gat att Ser Ile Glu Glu Glu Asp Ala Glu Ser Met His Lys Tyr His Asp Ile 465 470 475	1443
gtt cgt gca tcg tgt act att cta agg ctt gct gat gat atg gga aca Val Arg Ala Ser Cys Thr Ile Leu Arg Leu Ala Asp Asp Met Gly Thr 480 485 490	1491
tcg ctg gat gag gtg gag aga ggc gac gtg ccc aaa tca gtt cag tgc Ser Leu Asp Glu Val Glu Arg Gly Asp Val Pro Lys Ser Val Gln Cys 495 500 505	1539
tac atg aat gag aag aat gct tcg gaa gaa gaa gcg cga gag cat gtg Tyr Met Asn Glu Lys Asn Ala Ser Glu Glu Glu Ala Arg Glu His Val 510 515 520 525	1587
cga tca ctc ata gac caa aca tgg aag atg atg aac aag gaa atg atg Arg Ser Leu Ile Asp Gln Thr Trp Lys Met Met Asn Lys Glu Met Met 530 535 540	1635
acg tca tca ttt tcc aaa tat ttt gta caa gtt tct gct aat ctt gca Thr Ser Ser Phe Ser Lys Tyr Phe Val Gln Val Ser Ala Asn Leu Ala 545 550 555	1683
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tat gag taa ctaatcttcg cccgggttcc aaatgaatca atctgttgtg	1828

T <b>y</b> r 590	Glu														
ttg	etgtt	ccc a	acct	gatat	c aa	ataat	aatt	aga	caaa	atgt	ttct	tgtad	gg q	gtggd	ccaac
cgto	caggo	ccc a	attt	egete	ca to	gttca	ataat	. aaa	taat	aaa	acto	gttaa	atc a	ataa	caaaa
aaaa	aaaa	aaa a	aaaa	aaaa	a										
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Thr	Thr	Ile 35	Thr	Thr	Arg	Gly	Gly 40	Arg	Trp	Ala	His	C <b>y</b> s 45	Ser	Leu	Gln
Met	Gly 50	Asn	Glu	Ile	Gln	Thr 55	Gly	Arg	Arg	Thr	Gly 60	Gly	Tyr	Gln	Pro
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Val	Glu 130	Ile	Leu	Asn	Ser	L <b>y</b> s 135	Tyr	Tyr	Thr	Asn	Asn 140	Glu	Ile	Asp	Glu
Ser 145	Asp	Leu	Tyr	Ser	Thr 150	Ala	Leu	Arg	Phe	L <b>y</b> s 155	Leu	Leu	Arg	Gln	Tyr 160
Asp	Phe	Ser	Val	Ser 165	Gln	Glu	Val	Phe	Asp 170	Суѕ	Phe	Lys	Asn	Asp 175	Lys
Gly	Thr	Asp	Phe 180	Lys	Pro	Ser	Leu	Val 185	Asp	Asp	Thr	Arg	Gly 190	Leu	Leu
Gln	Leu	<b>Ty</b> r 195	Glu	Ala	Ser	Phe	Leu 200	Ser	Ala	Gln	Gly	Glu 205	Glu	Thr	Leu
His	Leu 210	Ala	Arg	Asp	Phe	Ala 215	Thr	Lys	Phe	Leu	His 220	Lys	Arg	Val	Leu
Val 225	Asp	Lys	Asp	Ile	Asn 230	Leu	Leu	Ser	Ser	Ile 235	Glu	Arg	Ala	Leu	Glu 240
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Lys	Gly	Phe	Asn	Ser 405	Thr	Pro	Tyr	Leu	Arg 410	Lys	Ala	Trp	Val	Asp 415	Leu	
Val	Glu	Ser	Tyr 420	Leu	Ile	Glu	Ala	L <b>y</b> s 425	Trp	Tyr	Tyr	Met	Gly 430	His	Lys	
Pro	Ser	Leu 435	Glu	Glu	Tyr	Met	Lys 440	Asn	Ser	Trp	Ile	Ser 445	Ile	Gly	Gly	
Ile	Pro 450	Ile	Leu	Ser	His	Leu 455	Phe	Phe	Arg	Leu	Thr 460	Asp	Ser	Ile	Glu	
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Phe 545	Ser	Lys	Tyr	Phe	Val 550	Gln	Val	Ser	Ala	Asn 555	Leu	Ala	Arg	Met	Ala 560	
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ctt Leu 30																145
tca Ser																193
cct Pro	-			_					_							241

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	ata Ile 95															337
	cat His	_	_			_			_	_				-		385
	aag Lys															433
	aat Asn															481
	ctc Leu															529
	ttc Phe 175															577
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	gat Asp	-			-						-	-		-		1105
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Arg	Gln	His	Gly	Phe 165	Asn	Ile	Ser	Gln	Asp 170	Val	Phe	Asn	Cys	Phe 175	Lys
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L <b>y</b> s 225	Leu	Asp	Glu	Gly	Gly 230	Asn	Glu	Ile	Asp	Glu 235	Asn	Leu	Leu	Leu	Trp 240
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Glu	Ala	Arg	Trp 260	Phe	Ile	Asp	Ala	<b>Ty</b> r 265	Ala	Arg	Arg	Pro	Asp 270	Met	Asn
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C <b>y</b> s 305	Phe	Pro	Glu	Lys	Leu	Dro									_
Phe	Dh o				310	FIO	Phe	Val	Arg	Asp 315	Arg	Leu	Val	Glu	320
	PHE	Trp	Ala	Val 325	310					315				Glu Tyr 335	320
Arg		-		325	310 Gly	Met	Phe	Glu	Pro 330	315 His	Gln	His	Gly	Tyr	320 Gln
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Ile Thr Met	Lys Tyr Phe 370 Gln	Met Asp 355 Lys Leu	Ala 340 Val Arg	325 Ala Tyr Trp	310 Gly Thr Gly Asp Trp 390	Met Ile Thr Thr 375 Gly	Phe Ile Leu 360 Glu Val	Glu Val 345 Asp Ser	Pro 330 Leu Glu Ile Asn	315 His Ala Leu Thr Tyr 395	Gln Thr Glu Arg 380 Ile	His Val Leu 365 Leu Ser	Gly Ile 350 Phe Pro	Tyr 335 Asp Thr	320 Gln Asp Tyr Ala 400
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Tyr His Asp Ile Leu Cys Leu Ala Gly Ile Ile Leu Arg Leu Pro Asp 485 490 495	
Asp Leu Gly Thr Ser Tyr Phe Glu Leu Ala Arg Gly Asp Val Pro Lys 500 505 510	
Thr Ile Gln Cys Tyr Met Lys Glu Thr Asn Ala Ser Glu Glu Glu Ala 515 520 525	
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Phe Ser Leu Asp Asp Lys Glu Gln Gln Lys Cys Ser Glu Thr Ile Glu 35 40 45	
Ala Leu Lys Gln Glu Ala Arg Gly Met Leu Met Ala Ala Thr Thr Pro 50 60	
Leu Gln Gln Met Thr Leu Ile Asp Thr Leu Glu Arg Leu Gly Leu Ser 70 75 80	
Phe His Phe Glu Thr Glu Ile Glu Tyr Lys Ile Glu Leu Ile Asn Ala 85 90 95	
Ala Glu Asp Asp Gly Phe Asp Leu Phe Ala Thr Ala Leu Arg Phe Arg 100 $0000000000000000000000000000000000$	
Leu Leu Arg Gln His Gln Arg His Val Ser Cys Asp Val Phe Asp Lys 115 120 125	
Phe Ile Asp Lys Asp Gly Lys Phe Glu Glu Ser Leu Ser Asn Asn Val	
Glu Gly Leu Leu Ser Leu Tyr Glu Ala Ala His Val Gly Phe Arg Glu 145 150 155 160	
Glu Arg Ile Leu Gln Glu Ala Val Asn Phe Thr Arg His His Leu Glu 165 170 175	
Gly Ala Glu Leu Asp Gln Ser Pro Leu Leu Ile Arg Glu Lys Val Lys 180 185 190	
Arg Ala Leu Glu His Pro Leu His Arg Asp Phe Pro Ile Val Tyr Ala 195 200 205	
Arg Leu Phe Ile Ser Ile Tyr Glu Lys Asp Asp Ser Arg Asp Glu Leu	

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	210					215					220				
Leu 225	Leu	Lys	Leu	Ser	L <b>y</b> s 230	Val	Asn	Phe	Lys	Phe 235	Met	Gln	Asn	Leu	Tyr 240
Lys	Glu	Glu	Leu	Ser 245	Gln	Leu	Ser	Arg	Trp 250	Trp	Asn	Thr	Trp	Asn 255	Leu
Lys	Ser	Lys	Leu 260	Pro	Tyr	Ala	Arg	Asp 265	Arg	Val	Val	Glu	Ala 270	Tyr	Val
Trp	Gly	Val 275	Gly	Tyr	His	Tyr	Glu 280	Pro	Gln	Tyr	Ser	<b>Ty</b> r 285	Val	Arg	Met
Gly	Leu 290	Ala	Lys	Gly	Val	Leu 295	Ile	Cys	Gly	Ile	Met 300	Asp	Asp	Thr	Tyr
Asp 305	Asn	Tyr	Ala	Thr	Leu 310	Asn	Glu	Ala	Gln	Leu 315	Phe	Thr	Gln	Val	Leu 320
Asp	Lys	Trp	Asp	Arg 325	Asp	Glu	Ala	Glu	Arg 330	Leu	Pro	Glu	Tyr	Met 335	Lys
Ile	Val	Tyr	Arg 340	Phe	Ile	Leu	Ser	Ile 345	Tyr	Glu	Asn	Tyr	Glu 350	Arg	Asp
Ala	Ala	L <b>y</b> s 355	Leu	Gly	Lys	Ser	Phe 360	Ala	Ala	Pro	Tyr	Phe 365	Lys	Glu	Thr
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Glu 385	Arg	Gln	Leu	Pro	Ser 390	Phe	Gln	Asp	Tyr	Val 395	Lys	Asn	Ser	Glu	L <b>y</b> s 400
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Ala	Thr	Ser 435	Thr	Ala	Met	Ile	Gly 440	Arg	Tyr	Trp	Asn	Asp 445	Thr	Ser	Ser
Gln	Leu 450	Arg	Glu	Ser	Lys	Gly 455	Gly	Glu	Met	Leu	Thr 460	Ala	Leu	Asp	Phe
His 465	Met	Lys	Glu	Tyr	Gly 470	Leu	Thr	Lys	Glu	Glu 475	Ala	Ala	Ser	Lys	Phe 480
Glu	Gly	Leu	Val	Glu 485	Glu	Thr	Trp	Lys	Asp 490	Ile	Asn	Lys	Glu	Phe 495	Ile
Ala	Thr	Thr	Asn 500	Tyr	Asn	Val	Gly	<b>A</b> rg 505	Glu	Ile	Ala	Ile	Thr 510	Phe	Leu
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Ala	<b>Ty</b> r 530	Ser	Asp	Pro	Asn	Val 535	Ala	Lys	Ala	Asn	Val 540	Val	Ala	Leu	Phe
Val 545	Asp	Ala	Ile	Val	Phe 550										
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tgc	egge	acg a	aggti	tatc	tt ga	agcti	tcct	c cat	tata	ggcc	aac	acata	atc a	atato	caaagg

 ${\tt tgccggcacg} \ {\tt aggttatctt} \ {\tt gagcttcctc} \ {\tt catataggcc} \ {\tt aacacatatc} \ {\tt atatcaaagg}$ 

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ctg cgc aag tcg ttg atc agt tca att cat gaa cat aag cct ccc tat Leu Arg Lys Ser Leu Ile Ser Ser Ile His Glu His Lys Pro Pro Tyr 15 20 25 30	158
aga aca atc cca aat ctt gga atg cgt agg cga ggg aaa tct gtc acg Arg Thr Ile Pro Asn Leu Gly Met Arg Arg Arg Gly Lys Ser Val Thr $35$ $40$ $45$	206
cct tcc atg agc atc agt ttg gcc acc gct gca cct gat gat ggt gta Pro Ser Met Ser Ile Ser Leu Ala Thr Ala Ala Pro Asp Asp Gly Val 50 55 60	254
caa aga cgc ata ggt gac tac cat tcc aat atc tgg gac gat gat ttc Gln Arg Arg Ile Gly Asp Tyr His Ser Asn Ile Trp Asp Asp Asp Phe 65 70 75	302
ata cag tct cta tca acg cct tat ggg gaa ccc tct tac cag gaa cgt Ile Gln Ser Leu Ser Thr Pro Tyr Gly Glu Pro Ser Tyr Gln Glu Arg 80 85 90	350
gct gag aga tta att gtg gag gta aag aag ata ttc aat tca atg tac Ala Glu Arg Leu Ile Val Glu Val Lys Lys Ile Phe Asn Ser Met Tyr 95 100 105 110	398
ctg gat gat gga aga tta atg agt tcc ttt aat gat ctc atg caa cgc Leu Asp Asp Gly Arg Leu Met Ser Ser Phe Asn Asp Leu Met Gln Arg 115 120 125	446
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aag aac gag ata aca tca gct ctg gat tat gtt ttc cgt tac tgg gag Lys Asn Glu Ile Thr Ser Ala Leu Asp Tyr Val Phe Arg Tyr Trp Glu 145 150 155	542
gaa aac ggc att gga tgt ggg aga gac agt att gtt act gat ctc aac Glu Asn Gly Ile Gly Cys Gly Arg Asp Ser Ile Val Thr Asp Leu Asn 160 165 170	590
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tct cca gag gtt tta aaa gct ttt caa gat caa aat gga cag ttt gta Ser Pro Glu Val Leu Lys Ala Phe Gln Asp Gln Asn Gly Gln Phe Val 195 200 205	686
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tat cgg gct tcc ctc att gcc ttc cct ggt gag aaa gtt atg gaa gaa Tyr Arg Ala Ser Leu Ile Ala Phe Pro Gly Glu Lys Val Met Glu Glu 225 230 235	782
gct gaa atc ttc tcc aca aga tat ttg aaa gaa gct cta caa aag att Ala Glu Ile Phe Ser Thr Arg Tyr Leu Lys Glu Ala Leu Gln Lys Ile 240 245 250	830
cca gtc tcc gct ctt tca caa gag ata aag ttt gtt atg gaa tat ggc Pro Val Ser Ala Leu Ser Gln Glu Ile Lys Phe Val Met Glu Tyr Gly 255 260 265 270	878
tgg cac aca aat ttg cca aga ttg gaa gca aga aat tac ata gac aca Trp His Thr Asn Leu Pro Arg Leu Glu Ala Arg Asn Tyr Ile Asp Thr 275 280 285	926
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		-			gct Ala 340			-			-				_	1118
					att Ile											1166
					cat His											1214
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					gag Glu											1310
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	_				aga Arg						_	_	_	-		1406
	-			_	tca Ser		_	-	_	_						1454
		_		_	ttt Phe	-									-	1502
	-				gca Ala								-		-	1550
					atc Ile 500											1598
					tcc Ser											1646
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	_		_		cct Pro				_	_	_	_				1742
					aat Asn											1790
					aat Asn 580											1838
					ctc Leu											1886
					gaa Glu											1934
-	tct Ser	_			taa	ctat	caaco	cat a	atcca	ataa	ta a	taag	ctcat	t		1982

339 340

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Met Ser Ile Ser Leu Ala Thr Ala Ala Pro Asp Asp Gly Val Gln Arg 50 55 60
Arg Ile Gly Asp Tyr His Ser Asn Ile Trp Asp Asp Asp Phe Ile Gln 65 70 75 80
Ser Leu Ser Thr Pro Tyr Gly Glu Pro Ser Tyr Gln Glu Arg Ala Glu 85 90 95
Arg Leu Ile Val Glu Val Lys Lys Ile Phe Asn Ser Met Tyr Leu Asp 100 105 110
Asp Gly Arg Leu Met Ser Ser Phe Asn Asp Leu Met Gln Arg Leu Trp 115 120 125
Ile Val Asp Ser Val Glu Arg Leu Gly Ile Ala Arg His Phe Lys Asn 130 135 140
Glu Ile Thr Ser Ala Leu Asp Tyr Val Phe Arg Tyr Trp Glu Glu Asn 145 150 150
Gly Ile Gly Cys Gly Arg Asp Ser Ile Val Thr Asp Leu Asn Ser Thr $165 \\ 170 \\ 175$
Ala Leu Gly Phe Arg Thr Leu Arg Leu His Gly Tyr Thr Val Ser Pro 180 185 190
Glu Val Leu Lys Ala Phe Gln Asp Gln Asn Gly Gln Phe Val Cys Ser 195 200 205
Pro Gly Gln Thr Glu Gly Glu Ile Arg Ser Val Leu Asn Leu Tyr Arg 210 215 220
Ala Ser Leu Ile Ala Phe Pro Gly Glu Lys Val Met Glu Glu Ala Glu 225 230 235 240
Ile Phe Ser Thr Arg Tyr Leu Lys Glu Ala Leu Gln Lys Ile Pro Val 245 250 255
Ser Ala Leu Ser Gln Glu Ile Lys Phe Val Met Glu Tyr Gly Trp His 260 265 270
Thr Asn Leu Pro Arg Leu Glu Ala Arg Asn Tyr Ile Asp Thr Leu Glu 275 280 285
Lys Asp Thr Ser Ala Trp Leu Asn Lys Asn Ala Gly Lys Lys Leu Leu 290 295 300
Glu Leu Ala Lys Leu Glu Phe Asn Ile Phe Asn Ser Leu Gln Gln Lys 305 310 315 320
Glu Leu Gln Tyr Leu Leu Arg Trp Trp Lys Glu Ser Asp Leu Pro Lys

												COII	CIII	ueu		
				325					330					335		
Leu	Thr	Phe	Ala 340	Arg	His	Arg	His	Val 345	Glu	Phe	Tyr	Thr	Leu 350	Ala	Ser	
Cys	Ile	Ala 355	Ile	Asp	Pro	Lys	His 360	Ser	Ala	Phe	Arg	Leu 365	Gly	Phe	Ala	
Lys	Met 370	Cys	His	Leu	Val	Thr 375	Val	Leu	Asp	Asp	Ile 380	Tyr	Asp	Thr	Phe	
Gl <b>y</b> 385	Thr	Ile	Asp	Glu	Leu 390	Glu	Leu	Phe	Thr	Ser 395	Ala	Ile	Lys	Arg	Trp 400	
Asn	Ser	Ser	Glu	Ile 405	Glu	His	Leu	Pro	Glu 410	Tyr	Met	Lys	Cys	Val 415	Tyr	
Met	Val	Val	Phe 420	Glu	Thr	Val	Asn	Glu 425	Leu	Thr	Arg	Glu	Ala 430	Glu	Lys	
Thr	Gln	Gly 435	Arg	Asn	Thr	Leu	Asn 440	Tyr	Val	Arg	Lys	Ala 445	Trp	Glu	Ala	
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Leu 465	Pro	Met	Phe	Glu	Glu 470	Tyr	His	Glu	Asn	Gl <b>y</b> 475	Lys	Val	Ser	Ser	Ala 480	
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Pro	Asp	Tyr	Ile 500	Leu	Lys	Gly	Ile	Asp 505	Phe	Pro	Ser	Arg	Phe 510	Asn	Asp	
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L <b>y</b> s 545	Asp	Asn	Pro	Gly	Ser 550	Thr	Glu	Glu	Asp	Ala 555	Leu	Asn	His	Ile	Asn 560	
Ala	Met	Val	Asn	Asp 565	Ile	Ile	Lys	Glu	Leu 570	Asn	Trp	Glu	Leu	Leu 575	Arg	
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Met 625	Leu	Phe														
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<400	)> SE	QUEN	ICE:	31												
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					gta Val										gtt	107

con can got gac the tet can agt cut tyg ggt gat ogt cut cat Arg Pro 11e Ala Asp Phe Ser Pro Ser Leu Trp Gly Asp Arg Phe His 25    to the tot toe cto gac aat ong att got gga aan tat get can agag atc																	
ser Phe Ser Leu Asp Asm Gal nie Ala Giy Lys Tyr Ala Gin Giu Tie  35 40 40 45 45 40 45 55 60 52 60 52 60 52 60 55 60 55 60 60 65 60 65 60 65 60 65 60 65 60 65 60 65 60 60 60 65 60 60 60 60 60 60 60 60 60 60 60 60 60			Ile					Pro					Asp				155
Glu Thr Leu Lya Glu Gln Ser Arg Tle Ile Leu Ser Ala Ser Ser Arag 55    aga aca ttg gct gag and ttg gat ctg ata gac att gtt gag cgc ctt   Arg Thr Leu Ala Glu Lya Leu Asp Leu Ile Asp Ile Val Glu Arg Leu   70    ggc att gct tat cat ttt gan and can ata gat gat atg ttg gat can   Gly Ile Ala Tyr His Phe Glu Lys Gln Ile Asp Asp Met Leu Asp Gln   95    ttt tac and gca gat cct and ttt gan gct cac gag tac and gat tta   Phe Tyr Lys Ala Asp Pro Asn Phe Glu Ala His Glu Tyr Asn Asp Leu   100    can act tta tcc gtt can ttt cga cta ttg aga can cat ggt tac and   Gln Thr Leu Ser Val Gln Phe Arg Leu Leu Arg Gln His Gly Tyr Asn   115    act tcc cca and ctt tt att aga ttc can agt gca and ggc and ttt   120    att tcc cca and ctt tt att aga ttc can agt gca and ggc and ttt   115    aca gan act ct ttt att aga ttc can agt gca and ggc and ttt   120    att tcc cca and ctt ttt att aga ttc can agt gca and ggc and ttt   121    122    act tcc cca and ctt ttt att aga ttc can ggt ctt ttg and tta tac gan   135    143    149    149    149    149    150    30    gcc tcg cat gta agg act cat gga gan gat att ttg gan gag gca   145    Ala Ser His Val Arg Thr His Gly Glu Ly Leu Leu Asn Leu Tyr Glu   165    gcc tcg cat gta agg act cat gga gan gat att ttg gan gag gca   215    gct ttc tct act gct cat ctt gan gan gat ctt ttg and tala Leu   175    gct ttc tct act gct cat gt gan gan gat att ttg gan gag gca   216    gct tct ct act gct cat ctt gan gan gat ctt cc   217    gct ttc tct act gct cat gt gan gan gat att ttg gan gan gca   218    cct ctg agt and gag can gtg act cat gga gan gat att ttg gan gan gca   218    grat tcc and gan gtt gan		Phe					Gln					Tyr					203
Arg Thr Leu Ala Glu Lys Leu Asp Leu Ile Asp Ile Val Glu Arg Leu Roll Glu Arg Leu Roll Glu Arg Leu Roll Glu Arg Leu Roll Glu First His Phe Glu Lys Gln Ile Asp Asp Met Leu Asp Gln Standard Stand	Glu					Gln					Leu					Arg	251
Giy Ile Ala Tyr His Phe Glu Lys Gln Ile Asp Asp Met Leu Asp Gln 85  ttt tac aaa gca gat cct aac ttt gag gct cac gag tac aat gat tta 100 100 105 Asp Pro Asn Phe Glu Ala His Glu Tyr Asn Asp Leu 100 105 110 110 110 110 110 110 110 110					Glu					Ile					Arg		299
Phe Tyr Lys Åla Åsp Pro Asn Phe Glú Åla His Glú Tyr Asn Åsp Leu 100 105 105 105 110 110 110 100 105 105				Tyr					Gln					Leu			347
Gln Thr Leu Ser Val Gln Phe Arg Leu Leu Arg Gln His Gly Tyr Asn 125  atc tcc cca aaa ctt ttt att atga ttc caa gat gca aaa ggc aaa ttt 11e Ser Pro Lys Leu Phe Ile Arg Phe Gln Asp Ala Lys Gly Lys Phe 130			Lys	_	-			Phe		_			Tyr		-		395
The Ser Pro Lys Leu Phe 118 Arg Phe Gln Asp Ala Lys Gly Lys Phe 130    aaa gaa tot ott tgt aac gac atc aag ggt cut ttg aac tat acc gaa Lys Glu Ser Leu Cys Asn Asp Ile Lys Gly Leu Leu Asn Leu Tyr Glu 150    gcc tog cat gta agg act cat gga gaa gat att ttg gaa gag gca ctt Ala Ser His Val Arg Thr His Gly Glu Asp Ile Leu Glu Glu Ala Leu 165    gct ttc tot act gct cat ctt gaa tot gaa gct cat gag gc gct cat ttg aag to ala Phe Ser Thr Ala His Leu Glu Ser Ala Ala Pro His Leu Lys Ser 180    cct ctg agt aag caa gtg aca cat gcc ctt gag caa tot ctc cat aag Pro Leu Ser Lys Gln Val Thr His Ala Leu Glu Gln Ser Leu His Lys 195    agc att cca aga gtt gag aca cgt act ttc atc tot act tat cac gaa gag gar att cca aga gtt can act gcc ctt gag caa tot ctc cat aag Pro Leu Ser Lys Gln Val Thr His Ala Leu Glu Gln Ser Leu His Lys 2005    agc att cca aga gtt gag aca cgc tac ttc atc tot atc tac gaa gag Ser Ile Pro Arg Val Glu Thr Arg Tyr Phe Ile Ser Ile Tyr Glu Glu 210    225    gag gaa cag aag aat gat gtg ttg ctt caa ttt gca aac ctg gac ttc Glu Gln Ser Leu His Lys 220    acc tta ctt cag at gat ttg cac aac aca ctt act tac tac gaa gag Ser Ile Pro Arg Val Glu Thr Arg Tyr Phe Ile Ser Ile Tyr Glu Glu Glu Glu Glu Glu Lys Asn Asp Val Leu Leu Gln Phe Ala Lys Leu Asp Phe 240    acc tta ctt cag at gtg ttg cac aca cat ttg ca act act gc gac ttc Glu		Thr			_		Phe	_			_	Gln					443
Lys Glu Ser Leu Cys Asn Asp Ile Lys Gly Leu Leu Asn Leu Tyr Glu 150    gcc tcg cat gta agg act cat gga gaa gat att ttg gaa gag gca ctt 165    gct ttc tct act gct cat ctt gaa tct gca gct cca cat ttg aag tca 170    gct ttc tct act gct cat ctt gaa tct gca gct cca cat ttg aag tca 180    gct ttc tct act gct cat ctt gaa tct gca gct cca cat ttg aag tca 180    gct ttc tct act gct cat ctt gaa tct gcg cca cat ttg aag tca 180    gct ttc tct act gct cat ctt gaa tct gca gct cca cat ttg aag tca 180    gct ttc tct act gct cat ctt gaa tct gcc ctt gag caa tct ctc cat aag 190    gct ttc ser Lys Gln Val Thr His Ala Leu Glu Gln Ser Leu His Lys 200    agc att cca aga gtt gag aca cat gcc ttc atc tct atc tac gaa gag 30    agc att cca aga gtt gag aca cac gcc tac ttc atc tct atc acc gaa gag 31    ser Ile Pro Arg Val Glu Thr Arg Tyr Phe Ile Ser Ile Tyr Glu Glu 210    215    gag gaa cag aag aat gat gtg ttg ctt caa ttt gca aaa ctg gac ttc Glu Glu Glu Gln Lys Asn Asp Val Leu Leu Gln Phe Ala Lys Leu Asp Phe 230    aac tta ctt cag atg ttg cac aaa caa gaa ctt agt gaa gta tca agg 827    Asn Leu Leu Gln Met Leu Hie Lys Gln Glu Leu Ser Glu Val Ser Arg 255    tgg tgg aaa gat ttg gat ttt gtg aca aca ctt cca tat gct agg gat    trp Trp Lys Asp Leu Asp Phe Val Thr Thr Leu Pro Tyr Ala Arg Asp 260    aga gca gtg gaa tgc tac ttt tgg aca acc ctt cca tat gct agg gat    trp Trp Lys Asp Leu Asp Phe Val Thr Thr Leu Pro Tyr Ala Arg Asp 260    aga gca gtg gaa tgc tac ttt tgg acg atg ggg gtg tat gct gaa cct Arg Ala Val Glu Cys Tyr Phe Trp Thr Met Gly Val Tyr Ala Glu Pro 280    caa tac tct cag gct cgt gtc atg ctt gct atg ctt gct aa gac atg atg att gcl    aac tct aag gat gac aca tc ggt gct atg ctt gct aaa gaa ctt 300    caa tac tct cag gct cgt aca ttc gat gct tat gcc aaa acc at acc acc acc acc acc acc ac	Ile					Phe					Asp					Phe	491
Ala Ser His Val Arg Thr His Gly Glu Asp Ile Leu Glu Glu Ala Leu 175  get tte tet act get eat cett gaa tet gea get eea cat ttg aag tea Ala Phe Ser Thr Ala His Leu Glu Ser Ala Ala Pro His Leu Lys Ser 180  cet etg agt aag caa gtg aca cat gee ett gag caa tet ett eat aag 683  Pro Leu Ser Lys Gln Val Thr His Ala Leu Glu Gln Ser Leu His Lys 200  age att eea aga gtt gag aca ege tac tte ate tet ate tac gaa gag gag eat 195  age att eea aga gtt gag aca ege tac tte ate tet ate tac gaa gag 731  Ser Ile Pro Arg Val Glu Thr Arg Tyr Phe Ile Ser Ile Tyr Glu Glu 225  gag gaa eag aag aat gat gtt gtt eett eat tte gea aaa ett gac tte 691  Glu Glu Gln Lys Ash Asp Val Leu Leu Gln Phe Ala Lys Leu Asp Phe 230  aac tta ett eag att gtt eac aaa eaa gaa ett agt gaa gta tea agp Phe 240  aac tta ett eag att gtt geat tt gtg aca aca ett eat get agg gat ste Arg 255  tgg tgg aaa gat ttg gat ttt gtg aca aca ett eat eget agg gat 875  Trp Trp Lys Asp Leu Asp Phe Val Thr Thr Leu Pro Tyr Ala Arg Asp 260  aga gag gag gtg gaa tge tac ttt tgg acg atg ggg gtt at get gaa eet 270  aga gag gag gtg gaa tge tac ttt tgg acg atg ggg gtg tat get gaa eet 270  aga gac gtg gaa tge tac ttt tgg acg atg ggg gtg tat get gaa eet 280  aga gac gtg gaa tge tac ttt tgg acg atg ggg gtg tat get gaa eet 280  aga gac gtg gaa tge tac ttt tgg acg atg ggg gtg tat get gaa eet 280  aga gac gtg gaa tge tac ttt tgg acg atg ggg gtg tat get gaa eet 280  aga gac gtg gaa tge tac ttt tgg acg atg ggg gtg tat get gaa eet 280  aga gac ata eet ea get get gte atg ett get aag act ata gea atg att 280  caa tac tet eag get gac aca tee gat get atg eet get aag act ata gea aag act ata get att 290  aac tac acc gat gac aca te each eag agg tgt gat att age aaa gaa ett 290  aac ata acc gat gac aca tee gat gat gat att age aaa gaa ett 310  aac tac acc gat gec ata eac aca acg agg tgg gat att age aaa gaa ett 280  aac ata eac acc gat gec ata eac acg agg tgg gat att age aaa gaa ett 310  aac tac acc gat gec ata eac acg agg tgg gat att age aaa tt age 61  aac tta etc acc gat gec ata eac acg agg tgg gat att age eaa					Cys					Gly					Tyr		539
Ala Phe Ser Thr Ala His Leu Glu Ser Ala Ala Pro His Leu Lys Ser 180 Cct ctg agt aag caa gtg aca cat gcc ctt gag caa tct ctc cat aag 683 Pro Leu Ser Lys Gln Val Thr His Ala Leu Glu Gln Ser Leu His Lys 205 agc att cca aga gtt gag aca cgc tac ttc atc tct atc tac gaa gag 731 Ser Ile Pro Arg Val Glu Thr Arg Tyr Phe Ile Ser Ile Tyr Glu Glu 210 225 aga gaa cag aat gat gtg ttg ctt caa ttt gca aaa ctg gat tc Glu Glu Gln Lys Asn Asp Val Leu Leu Gln Phe Ala Lys Leu Asp Phe 230 aca tta ctt cag at gat tc a gaa ctt act tac aga gad ctt act act cag at gtg ttg ctt caa ttt gca aca ctg gat tc Asn Leu Leu Gln Met Leu His Lys Gln Glu Leu Ser Glu Val Ser Arg 250 aca at act gaa gat tca agg gad gad gad gad gad tca aca cac aca aca cac ctt act act agg gad gad gad gad ser act act cac aca gad act tact cac act gad				Val					Glu					Glu			587
Pro Leu Ser Lys Gln Val Thr His Ala Leu Glu Gln Ser Leu His Lys 200 205 205 205 205 205 205 205 205 205	-		Ser		-			Glu		_	_		His	_	_		635
Ser Ile Pro Arg Val Glu Thr Arg Tyr Phe Ile Ser Ile Tyr Glu Glu 225  gag gaa cag aag aat gat gtg ttg ctt caa ttt gca aaa ctg gac ttc 779 Glu Glu Glu Lys Asn Asp Val Leu Leu Gln Phe Ala Lys Leu Asp Phe 230  aac tta ctt cag atg ttg cac aaa caa gaa ctt agt gaa gta tca agg 827 Asn Leu Leu Gln Met Leu His Lys Gln Glu Leu Ser Glu Val Ser Arg 245  tgg tgg aaa gat ttg gat ttt gtg aca aca ctt cca tat gct agg gat 875 Trp Trp Lys Asp Leu Asp Phe Val Thr Thr Leu Pro Tyr Ala Arg Asp 265  aga gca gtg gaa tgc tac ttt tgg acg atg ggg gtg tat gct gaa cct 270  aga gca gtg gaa tgc tac ttt tgg acg atg ggg gtg tat gct gaa cct 280  caa tac tct cag gct cgt gtc atg ctt gct aag act ata gca atg atg Pro 275  caa tac tct cag gct cgt gtc atg ctt gct aag act ata gca atg atg Pro 275  caa tac tct cag gct cgt gtc atg ctt gct aag act ata gca atg atg 11e Tyr Ser Gln Ala Arg Val Met Leu Ala Lys Thr Ile Ala Met Ile 290  caa tac gat gat gac aca ttc gat gct tat ggc att gtc aaa gaa ctt 1290  ser Ile Val Asp Asp Thr Phe Asp Ala Tyr Gly Ile Val Lys Glu Leu 315  gag atc tac acc gat gcc ata cag agg tgg gat att agc caa att gat 1067 Glu Ile Tyr Thr Asp Ala Ile Gln Arg Trp Asp Ile Ser Gln Ile Asp		Leu	_	_			Thr		_			Gln				_	683
Glu Glu Gln Lys Asn Asp Val Leu Leu Gln Phe Ala Lys Leu Asp Phe 230 235 240 240 235 235 240 240 235 235 240 240 240 235 250 240 240 240 240 240 240 240 240 240 24	Ser					Glu					Ile					Glu	731
Asn Leu Leu Gln Met Leu His Lys Gln Glu Leu Ser Glu Val Ser Arg 245  tgg tgg aaa gat ttg gat ttt gtg aca aca ctt cca tat gct agg gat 875  Trp Trp Lys Asp Leu Asp Phe Val Thr Thr Leu Pro Tyr Ala Arg Asp 265  aga gca gtg gaa tgc tac ttt tgg acg atg ggg gtg tat gct gaa cct 923  Arg Ala Val Glu Cys Tyr Phe Trp Thr Met Gly Val Tyr Ala Glu Pro 275  caa tac tct cag gct cgt gtc atg ctt gct aaa act ata gca atg att gln Tyr Ser Gln Ala Arg Val Met Leu Ala Lys Thr Ile Ala Met Ile 290  tct ata gta gat gac aca ttc gat gct tat ggc att gtc aaa gaa ctt 1019  Ser Ile Val Asp Asp Thr Phe Asp Ala Tyr Asp Ile Ser Gln Ile Asp  1067  Glu Ile Tyr Thr Asp Ala Ile Gln Arg Trp Asp Ile Ser Gln Ile Asp		-	_	_	Asn	_		_		Gln		-		_	Asp		779
Trp Trp Lys Asp Leu Asp Phe Val Thr Thr Leu Pro Tyr Ala Arg Asp 260 265 270  aga gca gtg gaa tgc tac ttt tgg acg atg ggg gtg tat gct gaa cct 923  Arg Ala Val Glu Cys Tyr Phe Trp Thr Met Gly Val Tyr Ala Glu Pro 275 280 285  caa tac tct cag gct cgt gtc atg ctt gct aag act ata gca atg att Gln Tyr Ser Gln Ala Arg Val Met Leu Ala Lys Thr Ile Ala Met Ile 300 305  tct ata gta gat gac aca ttc gat gct tat ggc att gtc aaa gac ctt 1019  Ser Ile Val Asp Asp Thr Phe Asp Ala Tyr Gly Ile Val Lys Glu Leu 310  gag atc tac acc gat gcc ata cag agg tgg gat att agc caa att gat 1067  Glu Ile Tyr Thr Asp Ala Ile Gln Arg Trp Asp Ile Ser Gln Ile Asp				Gln					Gln					Val			827
Arg Ala Val Glu Cys Tyr Phe Trp Thr Met Gly Val Tyr Ala Glu Pro 275  caa tac tct cag gct cgt gtc atg ctt gct aag act ata gca atg att 971 Gln Tyr Ser Gln Ala Arg Val Met Leu Ala Lys Thr Ile Ala Met Ile 290  ctct ata gta gat gac aca ttc gat gct tat ggc att gtc aaa gaa ctt 1019 Ser Ile Val Asp Asp Thr Phe Asp Ala Tyr Gly Ile Val Lys Glu Leu 310  gag atc tac acc gat gcc ata cag agg tgg gat att agc caa att gat 1067 Glu Ile Tyr Thr Asp Ala Ile Gln Arg Trp Asp Ile Ser Gln Ile Asp			Lys					Val					${\tt Tyr}$				875
Gln Tyr Ser Gln Ala Arg Val Met Leu Ala Lys Thr Ile Ala Met Ile 290 295 300 305  tct ata gta gat gac aca ttc gat gct tat ggc att gtc aaa gaa ctt Ser Ile Val Asp Asp Thr Phe Asp Ala Tyr Gly Ile Val Lys Glu Leu 310 315 320  gag atc tac acc gat gcc ata cag agg tgg gat att agc caa att gat Glu Ile Tyr Thr Asp Ala Ile Gln Arg Trp Asp Ile Ser Gln Ile Asp		Ala					Phe					Val					923
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#### -continued

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His	Ser	Phe 35	Ser	Leu	Asp	Asn	Gln 40	Ile	Ala	Gly	Lys	Tyr 45	Ala	Gln	Glu
Ile	Glu 50	Thr	Leu	Lys	Glu	Gln 55	Ser	Arg	Ile	Ile	Leu 60	Ser	Ala	Ser	Ser
Arg 65	Arg	Thr	Leu	Ala	Glu 70	Lys	Leu	Asp	Leu	Ile 75	Asp	Ile	Val	Glu	Arg 80
Leu	Gly	Ile	Ala	<b>Ty</b> r 85	His	Phe	Glu	Lys	Gln 90	Ile	Asp	Asp	Met	Leu 95	Asp
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Leu	Gln	Thr 115	Leu	Ser	Val	Gln	Phe 120	Arg	Leu	Leu	Arg	Gln 125	His	Gly	Tyr
Asn	Ile 130	Ser	Pro	Lys	Leu	Phe 135	Ile	Arg	Phe	Gln	Asp 140	Ala	Lys	Gly	Lys
Phe 145	Lys	Glu	Ser	Leu	C <b>y</b> s 150	Asn	Asp	Ile	Lys	Gl <b>y</b> 155	Leu	Leu	Asn	Leu	<b>Ty</b> r 160
Glu	Ala	Ser	His	Val 165	Arg	Thr	His	Gly	Glu 170	Asp	Ile	Leu	Glu	Glu 175	Ala
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Phe	Asn	Leu	Leu	Gln 245	Met	Leu	His	Lys	Gln 250	Glu	Leu	Ser	Glu	Val 255	Ser
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Arg	Val	Ile	Asp	Asp	Ile	Ala	Thr	Tyr	Glu	Val	Glu	Lys	Gly	Arg	Gly

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aag cag tct atc cga agg ggc ttg cca agg gtt gag gcc cgg aat ttc Lys Gln Ser Ile Arg Arg Gly Leu Pro Arg Val Glu Ala Arg Asn Phe 205 210 215	2352

Let tog dat the can get the gas too cut ase can tog the cut of the Series (19 cut) can be presented as the series of the Series (19 cut) series in annuy Series (19 cut) can be presented as the case they the case the type of the series (19 cut) can be presented as the case they are carefully care		
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geotaatact attroaattg atoacacqac tgttgctgac attratagat gettrittta 2556  ggt gg tgg aas gat tta gac ttt ace ags aas cta cca ttt gca ags 2603  Trp Trp lys Asp Lou Asp Phe Thr Asg Lys Lou Pro Phe Ala Asg 260  at aga gtg gtt gaa ggot tat ttt trg ata stg ggs gtt tro tit gaa 2651  Asp Asg Val Tel Gill Gily Try Phe Tap gle Net Gily Val Typ Phe Gil 2651  acc cca tac tct ctt ggt ags aag atg tg aca aas gcc ata gca atg 2699  Pro Gil Tyr Ser Lou Gily Asg Lys Met Lou Thr Lys Val I lie Ala Net 2795  ccc cca tac tct ggt ags aag atg tg aca aas gcc ata gca atg 2699  ggt tor att gft gat gat act tat gat toa tat gos acc tat gca atg gas 310 310  ggt toc att ggt ags aga atg ag agggagatit ttttcottt 2794  als Ser Ile Val Asp Asp Thr Tyr Asp Ser Tyr Ala Thr Tyr Asp Gilu 315  ccc cca aaa ac aasaasaagt tttggaatc ccccaagaat aggggagaat attggttt 2914  cccccaaaaa aasaasaagt tttggagatc ccccaagaat tgggtttt agg tag gat Arg Tep Asp Asg Tyr Ala Thr Tyr Asp Asg Tyr Ala Thr Tyr Asp Asg Tyr Ala Thr Tyr Asp Asg Tyr	Phe Ala Lys Ile Asp Phe Asn Leu Leu Gln Leu Leu His Arg Lys Glu	2448
gg tgg tgg aaa gat tta gac ttt aca aga aaa cta cca ttt gca aga Trp Trp Tup Aap Leu Aap Phe Tr Acg Lye Leu Pro Phe Ala Arg 260  gat aga gtg gtt gaa ggo tat ttt tgg ata atg gga gtt tac ttt gaa Aap Arg Val Gul Gly Tyr Phe Trp Ile Mer Gly Val Tyr Phe Glu 275  cco caa tac tot ctt ggt aga aag atg ttg aca aaa gto ata goa atg Pro Glu Tyr Ser Leu Gly Acg Lye Met Leu Thr Lya Val Ile Ala Met 270  gct tco att gtt gat gat act tat gat tca tat goa acc tat gat gaa gct tco att gtt gat gat act tat gat tca tat gaa acc tat gat gaa Ala Ser Ile Val Aap Aap Thr Tyr Aap Ser Tyr Ala Thr Tyr Aap Glu 305  ctc att ccc tat aca aat gca att gaa ggtgagattt ttttccttt 2794  Las Ile Pro Tyr Thr Aam Ala Ile Glu 220  cctccaaaaa aaaaaaaagtt tttgagcac ccccaagaat aggggaaaat atatgttttt 2854  aaacgttagg atattcactc caacttgcag ttgctcatat tttaatggtg atagstatga 2717  att aaa tgc atg acc caa ctc ccg aat tca atg aaa at agc tac aag Ile Lya Cys Met Aan Glu Leu Pro Aan Tyr Net Lys Ile Ser Tyr Iye 335  gca cta tta gat gtt tat gaa gaa atg gaa cag ctg ttg gca aat caa Ala Leu Leu Aap Val Tyr Glu Glu Met Glu Glu Leu Leu Ala Aan Glu 395  ggg aga cag tac cga gtt gag tag agg aga gg gtatgtatag 316  ggg aga cag tac gda gtt gag ga atg gg aaa agg gc gtatgtatag 317  tttgcttact ttttatgcc tttaatccata acgtataaaa ttgaaaatt acattagcaa 317  tttgcttact ttttatgcc tttaatccata acgtataaaa ttgaaaatt acattagcaa 317  tttgcttact ttttatgcc tttaatccata acgtataaaa ttgaaaat acatagcaa 317  tttgcttact tttttatgcc tttaatccata acgtataaaa ttgaaaaat acatagcaa 317  tttgcttact ttttatgcc tttaatccata acgtataaaa ttgaaaaat acatagcaa 317  tttgcttact ttttaag ag gaa atg ga ga gt gt gt gc ga gt gc atg gc gaa gtt ata acc cc acc tt 327  428  429  tt aaa atg gc gc agt gac ccc aag atc att aag g	Leu Ser Glu Ile Cys Arg	2496
Trp Trp Lya Asp Leu Asp Phe Thr Asq Lys Leu Pro Phe Ala Asq 260 260 270 260 270 260 270 260 270 260 270 275 270 275 270 275 275 270 275 275 275 275 275 275 275 275 275 275	gtctaatact atttcaattg atcacacgac tgttgctgac attttatgat gctttttta	2556
Asp Arg Val Val Giu Giy Tyr Phe Tup Ile Met Giy Val Tyr Phe Giu 275 275 276 277 278 278 278 279 279 279 270 270 270 270 270 270 270 271 270 270 271 270 271 270 271 271 271 271 271 271 271 271 271 271	Trp Trp Lys Asp Leu Asp Phe Thr Arg Lys Leu Pro Phe Ala Arg	2603
Pro Gln Tyr Ser Leu Giy Arg Lys Met Leu Thr Lys Val Ile Ala Met 300 gct too att gtt gat gat act tat gat too att god acc tat gat gaa Ala Ser Ile Val Aap Aap Thr Tyr Aap Ser Tyr Ala Thr Tyr Aap Glu 305 ctc att coc tat aca aat god att gaa ggtgagattt ttttccttt 2794 Leu Ile Pro Tyr Thr Aam Ala Ile Glu 3220 cctccasaaa aaaaaaaagt ttttgagatc ccccaagaat aggggaaaat atatgtttt 2854 aaacgttagg atattcactc caacttgcag ttgctcatat tttaatggtg atagtagaa 2914 ctaaccaggc taagttttag attcaaatta accctgaaat tgtgtttt agg tgg gat Arg Aap Thr Aap	Asp Arg Val Val Glu Gly Tyr Phe Trp Ile Met Gly Val Tyr Phe Glu	2651
Ala Ser Ile Val Aep Aep Thr Tyr Aep Ser Tyr Ala Thr Tyr Aep Glu 305  ctc att coc tat aca aat gca att gaa ggtgagattt ttttccttt  Leu Ile Pro Tyr Thr Aen Ala Ile Glu 320  225  cctccaaaaa aaaaaaagt ttttgagatc ccccaagaat aggggaaaat atatgtttt  aaacgttagg atatccactc caacttgcag ttgctcatat tttaatggtg atagtatgaa  ctaaccaggc taagtttag attcaaatta accctgaaat tgtgtttt agg tgg gat Arg Trp Aep 330  att aaa tgc atg aac caa ctc ccg aat tac atg aaa ata agc tac aag Ile Lys Cys Met Aen Gln Leu Pro Aen Tyr Met Lys Ile Ser Tyr Lys 335  gca cta tta gat gtt tat gaa gaa atg gaa cag ctg ttg gca aat caa Ala Leu Leu Aep Val Tyr Glu Glu Met Glu Gln Leu Leu Ala Aen Gln 355  ggg aga cag tac cga gtt gag tat gcg aaa aag gcg gtatgtaatg 3173  atacaatagt atgatatgct ttaatcataa acgtataaaa tttgaaaatt acattagcaa 3173  tttgcttact tttttatgcc tttaatcctc ag atg ata cg ctc tgt caa gc 365  gag gag att gag gcc aaa tgg act cac caa aat tat aaa cca acc tt Tyr Leu Leu Glu Ala Lys Trp Thr His Gln Aen Tyr Lys Pro Thr Phe 385  gag gaa ttt aga gat aat gca tat cca acc tct gc tat gcc atg ct 190  gct ata acc gcg tt tgtc gcc atg cgc gaa gt tat acc acc tct gc ata gcc 410  gct ata acc gcg tt tgtc gcc atg gcc gaa gt tat acc gct tac acc 410  425  436  437  437  438  439  439  439  439  431  431  432  432  434  435  436  436  436  436  436  436	Pro Gln Tyr Ser Leu Gly Arg Lys Met Leu Thr Lys Val Ile Ala Met	2699
Leu Ile Pro Tyr Thr Asn Ala Ile Glu 325  cctccaaaaa aaaaaaaagt ttttgagatc ccccaagaat aggggaaaat atatgtttt 2854  aaacgttagg atattcactc caacttgcag ttgctcatat tttaatggtg atagtaatga  ctaaccaggc taagttttag attcaaatta accctgaaat tggtgttt agg tgg gat Arg Trp Asp 330  att aaa tgc atg aac caa ctc ccg aat tac atg aaa ata agc tac aag Ile Lys Cys Met Asn Gln Leu Pro Asn Tyr Met Lys Ile Ser Tyr Lys 335  goa cta tta gat gtt tat gaa gaa atg gaa cag ctg ttg goa aat caa Ala Leu Leu Asp Val Tyr Glu Glu Met Glu Gln Leu Leu Ala Asn Gln 350  ggg aga cag tac cga gtt gag tat gog aaa ag gog gtatgtaatg Gly Arg Gln Tyr Arg Val Glu Tyr Ala Lys Lys Ala 370  atacaatagt atgatatgct ttaatcataa acgtataaaa tttgaaaatt acattagcaa  tttgcttact tttttatgcc tttaatcctc ag atg ata cgt ctt gtt caa gct Met Ile Arg Leu Val Gln Ala 385  gag gag at att aga gat aat gca ttg cca aac aat tat aaa caa acc ttt Tyr Leu Leu Glu Ala Lys Trp Thr His Gln Asn Tyr Lys Pro Thr Phe 385  gag gaa att aga gat aat gca ttg cca acc tt ggc tat gcc atg ct Glu Glu Phe Arg Asp Asn Ala Leu Pro Thr Ser Gly Tyr Ala Met Leu 400  405  gct ata acg gcg ttt gtc ggc atg ggc gaa gtt ata acc acc acc 425  430  ttc aaa tgg gcc gcc agt gac ccc aag atc att aca gcd tcca acc act ttc aac acc gcg gcd ggc gaa gtt cat acc	Ala Ser Ile Val Asp Asp Thr Tyr Asp Ser Tyr Ala Thr Tyr Asp Glu	2747
ctaaccaggc taagttttag attcaaatta accttgaat tttaatggtg atagtatgaa 2914  ctaaccaggc taagttttag attcaaatta accttgaaat tgtgttt agg tgg gat Arg Trp Aap 330  att aaa tgc atg aac caa ctc ccg aat tac atg aaa ata agc atac aag 3019  Ile Lys Cys Met Asn Gln Leu Pro Asn Tyr Met Lys Ile Ser Tyr Lys 335  gca cta tta gat gtt tat gaa gaa atg gaa cag ctg ttg gca aat caa Ala Leu Leu Asp Val Tyr Glu Glu Met Glu Gln Leu Leu Ala Asn Gln 350  ggg aga cag tac cga gtt gag tat gcg aaa aag gcg gtatgtaatg 3113  gly Arg Gln Tyr Arg Val Glu Tyr Ala Lys Lys Ala 365  atacaatagt atgatatgct ttaatcataa acgtataaaa tttgaaaatt acattagcaa 3173  tttgcttact tttttatgcc tttaatcatc ag atg ata cgt ctt gtt caa gct Met Ile Arg Leu Val Gln Ala la 380  tac ctt ttg gag gcc aaa tgg act cat caa aat tat aaa cca acc ttt 3274  Tyr Leu Leu Glu Ala Lys Tyr Thr His Gln Asn Tyr Lys Pro Thr Phe 385  gag gaa ttt aga gat aat gca ttg cca acc tct ggc tat gcc atg ctt gtc alg cat gcc atg ctc Glu Glu Phe Arg Asp Asn Ala Leu Pro Thr Ser Gly Tyr Ala Met Leu 400  gct ata acg gcg ttt gtc ggc atg ggc gaa gtt ata acc cct gag acc Ala Ile Thr Ala Phe Val Gly Met Gly Glu Val Ile Thr Pro Glu Thr 415  ttc aaa tgg gcc gcc agt gac ccc aag atc att aag gct tcc acc att apg ttc aac acc acc ttc gcc Ala Ile Thr Ala Phe Val Gly Met Gly Glu Val Ile Thr Pro Glu Thr 425  ttc aaa tgg gcc gcc agt gac ccc aag atc att aag gct tcc acc att Ada Sr Asp Pro Lys Ile Ile Lys Ala Ser Thr Ile 435  att tgc agg ttc atg gac gat att gtg gac cat ag gtatactata 3464	Leu Ile Pro Tyr Thr Asn Ala Ile Glu	2794
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att aaa tgc atg aac caa ctc ccg aat tac atg aaa ata agc tac aag Ile Lys Cys Met Asn Gln Leu Pro Asn Tyr Met Lys Ile Ser Tyr Lys 335  gca cta tta gat gtt tat gaa gaa atg gaa cag ctg ttg gca aat caa Ala Leu Leu Asp Val Tyr Glu Glu Met Glu Gln Leu Leu Ala Asn Gln 350  ggg aga cag tac cga gtt gag tat gcg aaa aag gcg gtatgtaatg Gly Arg Gln Tyr Arg Val Glu Tyr Ala Lys Lys Ala 365  atacaatagt atgatatgct ttaatcataa acgtataaaa tttgaaaatt acattagcaa  tttgcttact tttttatgcc tttaatcctc ag atg ata cgt ctt gtt caa gct Met Ile Arg Leu Val Gln Ala 380  tac ctt ttg gag gcc aaa tgg act cat caa aat tat aaa cca acc ttt Tyr Leu Leu Glu Ala Lys Trp Thr His Gln Asn Tyr Lys Pro Thr Phe 385  gag gaa ttt aga gat aat gca ttg cca acc tct ggc tat gcc atg ctt Glu Glu Phe Arg Asp Asn Ala Leu Pro Thr Ser Gly Tyr Ala Met Leu 400  gct ata acg gcg ttt gtc ggc gatg ggc gaa gtt ata acc cct gag acc Ala Ile Thr Ala Phe Val Gly Met Gly Glu Val Ile Thr Pro Glu Thr 420  ttc aaa tgg gcc gcc agt gac ccc aag atc att aag gct tcc acc att Phe Lys Trp Ala Ala Ser Asp Pro Lys Ile Ile Lys Ala Ser Thr Ile 445  att tgc agg ttc atg gac gat att gct gaa cat aag gtatactata 3464	aaacgttagg atattcactc caacttgcag ttgctcatat tttaatggtg atagtatgaa	2914
Ile Lys Cys Met Asn Gln Leu Pro Asn Tyr Met Lys Ile Ser Tyr Lys 345  gca cta tta gat gtt tat gaa gaa atg gaa cag ctg ttg gca aat caa Ala Leu Leu Asp Val Tyr Glu Glu Met Glu Gln Leu Leu Ala Asn Gln 350  ggg aga cag tac cga gtt gag tat gcg aaa aag gcg gtatgtaatg Gly Arg Gln Tyr Arg Val Glu Tyr Ala Lys Lys Ala 375  atacaatagt atgatatgct ttaatcataa acgtataaaa tttgaaaatt acattagcaa 3173  tttgcttact tttttatgcc tttaatcctc ag atg atg cct gtt caa gct Met Ile Arg Leu Val Gln Ala 380  tac ctt ttg gag gcc aaa tgg act cat caa aat tat aaa cca acc ttt Tyr Leu Leu Glu Ala Lys Trp Thr His Gln Asn Tyr Lys Pro Thr Phe 385  gag gaa ttt aga gat aat gca ttg cca acc tct ggc tat gcc atg ctt Glu Glu Phe Arg Asp Asn Ala Leu Pro Thr Ser Gly Tyr Ala Met Leu 400  gct ata acg gcg ttt gtc ggc atg ggc gaa gtt ata acc cct gag acc Ala Ile Thr Ala Phe Val Gly Glu Wet Gly Glu Val Ile Thr Pro Glu Thr 415  420  att tgc agg tc atg gac cat gac cct aag act att aag gct tcc acc att Phe Lys Trp Ala Ala Ser Asp Pro Lys Ile Ile Lys Ala Ser Thr Ile 435  att tgc agg tc atg gac gat att gct gaa cat aag gtatactata 3464  lie Cys Arg Phe Met Asp Asp Ile Ala Glu His Lys	Arg Trp Asp	2971
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Gly Arg Gln Tyr Arg Val Glu Tyr Ala Lys Lys Ala 375  atacaatagt atgatatgct ttaatcataa acgtataaaa tttgaaaatt acattagcaa 3173  tttgcttact tttttatgcc tttaatcctc ag atg ata cgt ctt gtt caa gct 3226  Met Ile Arg Leu Val Gln Ala 380  tac ctt ttg gag gcc aaa tgg act cat caa aat tat aaa cca acc ttt 77r Leu Leu Glu Ala Lys Trp Thr His Gln Asn Tyr Lys Pro Thr Phe 385  gag gaa ttt aga gat aat gca ttg cca acc tct ggc tat gcc atg ctt 3322  Glu Glu Phe Arg Asp Asn Ala Leu Pro Thr Ser Gly Tyr Ala Met Leu 400  gct ata acg gcg ttt gtc ggc atg ggc gaa gtt ata acc cct gag acc 410  Ala Ile Thr Ala Phe Val Gly Met Gly Glu Val Ile Thr Pro Glu Thr 425  ttc aaa tgg gcc gcc agt gac ccc aag atc att aag gct tcc acc att 430  ttc aaa tgg gcc gcc agt gac ccc aag atc att aag gct tcc acc att 740  Ala Ile Tyr Ala Ala Ser Asp Pro Lys Ile Ile Lys Ala Ser Thr Ile 435  att tgc agg ttc atg gac gat att gct gaa cat aag gtatactata 3464  Ile Cys Arg Phe Met Asp Asp Ile Ala Glu His Lys	Ala Leu Leu Asp Val Tyr Glu Glu Met Glu Gln Leu Leu Ala Asn Gln	3067
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Met Ile Arg Leu Val Gln Ala 380  tac ctt ttg gag gcc aaa tgg act cat caa aat tat aaa cca acc ttt Tyr Leu Leu Glu Ala Lys Trp Thr His Gln Asn Tyr Lys Pro Thr Phe 385  gag gaa ttt aga gat aat gca ttg cca acc tct ggc tat gcc atg ctt Glu Glu Phe Arg Asp Asn Ala Leu Pro Thr Ser Gly Tyr Ala Met Leu 400  gct ata acg gcg ttt gtc ggc atg ggc gaa gtt ata acc cct gag acc Ala Ile Thr Ala Phe Val Gly Met Gly Glu Val Ile Thr Pro Glu Thr 415  ttc aaa tgg gcc gcc agt gac ccc aag atc att aag gct tcc acc att Phe Lys Trp Ala Ala Ser Asp Pro Lys Ile Ile Lys Ala Ser Thr Ile 435  att tgc agg ttc atg gac gat att gct gaa cat aag gtatactata  3464  Ile Cys Arg Phe Met Asp Asp Ile Ala Glu His Lys	atacaatagt atgatatgct ttaatcataa acgtataaaa tttgaaaatt acattagcaa	3173
Tyr Leu Leu Glu Ala Lys Trp Thr His Gln Asn Tyr Lys Pro Thr Phe 385  gag gaa ttt aga gat aat gca ttg cca acc tct ggc tat gcc atg ctt Glu Glu Phe Arg Asp Asn Ala Leu Pro Thr Ser Gly Tyr Ala Met Leu 400  gct ata acg gcg ttt gtc ggc atg ggc gaa gtt ata acc cct gag acc Ala Ile Thr Ala Phe Val Gly Met Gly Glu Val Ile Thr Pro Glu Thr 415  ttc aaa tgg gcc gcc agt gac ccc aag atc att aag gct tcc acc att Phe Lys Trp Ala Ala Ser Asp Pro Lys Ile Ile Lys Ala Ser Thr Ile 435  att tgc agg ttc atg gac gat att gct gaa cat aag gtatactata Ile Cys Arg Phe Met Asp Asp Ile Ala Glu His Lys  Tyr Ala Tyr Phe 395  3322  3370  3370  3370  3418  3418  3464	Met Ile Arg Leu Val Gln Ala	3226
Glu Glu Phe Arg Asp Asn Ala Leu Pro Thr Ser Gly Tyr Ala Met Leu 400  gct ata acg gcg ttt gtc ggc atg ggc gaa gtt ata acc cct gag acc Ala Ile Thr Ala Phe Val Gly Met Gly Glu Val Ile Thr Pro Glu Thr 415  ttc aaa tgg gcc gcc agt gac ccc aag atc att aag gct tcc acc att Phe Lys Trp Ala Ala Ser Asp Pro Lys Ile Ile Lys Ala Ser Thr Ile 435  att tgc agg ttc atg gac gat att gct gaa cat aag gtatactata Ile Cys Arg Phe Met Asp Asp Ile Ala Glu His Lys  3370  3370  3418  3464	Tyr Leu Leu Glu Ala Lys Trp Thr His Gln Asn Tyr Lys Pro Thr Phe	3274
Ala Ile Thr Ala Phe Val Gly Met Gly Glu Val Ile Thr Pro Glu Thr 415 420 425 430  ttc aaa tgg gcc gcc agt gac ccc aag atc att aag gct tcc acc att Phe Lys Trp Ala Ala Ser Asp Pro Lys Ile Ile Lys Ala Ser Thr Ile 435 440 445  att tgc agg ttc atg gac gat att gct gaa cat aag gtatactata Ile Cys Arg Phe Met Asp Asp Ile Ala Glu His Lys	Glu Glu Phe Arg Asp Asn Ala Leu Pro Thr Ser Gly Tyr Ala Met Leu	3322
Phe Lys Trp Ala Ala Ser Asp Pro Lys Ile Ile Lys Ala Ser Thr Ile 435 440 445  att tgc agg ttc atg gac gat att gct gaa cat aag gtatactata Ile Cys Arg Phe Met Asp Asp Ile Ala Glu His Lys	Ala Ile Thr Ala Phe Val Gly Met Gly Glu Val Ile Thr Pro Glu Thr	3370
Ile Cys Arg Phe Met Asp Asp Ile Ala Glu His Lys	Phe Lys Trp Ala Ala Ser Asp Pro Lys Ile Ile Lys Ala Ser Thr Ile	3418
	Ile Cys Arg Phe Met Asp Asp Ile Ala Glu His Lys	3464

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tat	gggt	gca q			n His			-	-	aA c	_		-	•	c gaa e Glu )	3634
	tac Tyr															3682
	aac Asn															3730
_	aaa Lys 505	_		_	_	_			_		_	_	_			3778
	gct Ala															3826
	gtt Val															3874
	ata Ile			tga	aati	caad	cat t	ggct	taaq	ga ti	taci	tatga	a gat	caaaa	atta	3929
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Ser	Ser	Glu	Asn 20	Arg	Pro	Lys	Ala	Asp 25	Phe	His	Pro	Gly	Ile 30	Trp	Gly	
Asp	Met	Phe 35	Ile	Ile	Cys	Pro	Asp 40	Thr								
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Val	Arg	Lys	Met 20	Ile	Met	Glu	Pro	Val 25	Asp	Asp	Ser	Asn	Gln 30	Lys	Leu	
Pro	Phe	Ile 35	Asp	Ala	Val	Gln	Arg 40	Leu	Gly	Val	Ser	Tyr 45	His	Phe	Glu	
Lys	Glu 50	Ile	Glu	Asp	Glu	Leu 55	Glu	Asn	Ile	Tyr	Arg 60	Asp	Thr	Asn	Asn	
Asn 65	Asp	Ala	Asp	Thr	Asp 70	Leu	Tyr	Thr	Thr	Ala 75	Leu	Arg	Phe	Arg	Leu 80	
Leu	Arg	Glu	His	Gl <b>y</b> 85	Phe	Asp	Ile	Ser	C <b>y</b> s 90							

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<211> LENGTH: 125
<212> TYPE: PRT
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Asp Ala Phe Asn Lys Phe Lys Asp Glu Ala Gly Asn Phe Lys Ala Ser
Leu Thr Ser Asp Val Gln Gly Leu Leu Glu Leu Tyr Glu Ala Ser Tyr
Met Arg Val His Gly Glu Asp Ile Leu Asp Glu Ala Ile Ser Phe Thr 35\,
Thr Ala Gln Leu Thr Leu Ala Leu Pro Thr Leu His His Pro Leu Ser 50 60
Glu Gln Val Gly His Ala Leu Lys Gln Ser Ile Arg Arg Gly Leu Pro 65 70 75 80
Arg Val Glu Ala Arg Asn Phe Ile Ser Ile Tyr Gln Asp Leu Glu Ser 85 \hspace{1.5cm} 90 \hspace{1.5cm} 95 \hspace{1.5cm}
His Asn Lys Ser Leu Leu Gln Phe Ala Lys Ile Asp Phe Asn Leu Leu
Gln Leu Leu His Arg Lys Glu Leu Ser Glu Ile Cys Arg
<210> SEQ ID NO 37
<211> LENGTH: 72
<212> TYPE: PRT
<213> ORGANISM: Gossypium arboreum
<400> SEQUENCE: 37
Trp Trp Lys Asp Leu Asp Phe Thr Arg Lys Leu Pro Phe Ala Arg Asp 1 \phantom{-}10\phantom{0}
Gln Tyr Ser Leu Gly Arg Lys Met Leu Thr Lys Val Ile Ala Met Ala
Ser Ile Val Asp Asp Thr Tyr Asp Ser Tyr Ala Thr Tyr Asp Glu Leu
                          55
Ile Pro Tyr Thr Asn Ala Ile Glu
<210> SEQ ID NO 38
<211> LENGTH: 47
<212> TYPE: PRT
<213> ORGANISM: Gossypium arboreum
<400> SEQUENCE: 38
Arg Trp Asp Ile Lys Cys Met Asn Gln Leu Pro Asn Tyr Met Lys Ile 1 \phantom{\bigg|} 10 \phantom{\bigg|} 15
Ser Tyr Lys Ala Leu Leu Asp Val Tyr Glu Glu Met Glu Gln Leu Leu 20 \hspace{1cm} 25 \hspace{1cm} 30 \hspace{1cm}
Ala Asn Gln Gly Arg Gln Tyr Arg Val Glu Tyr Ala Lys Lys Ala 35 40 45
<210> SEQ ID NO 39
<211> LENGTH: 83
<212> TYPE: PRT
<213> ORGANISM: Gossypium arboreum
<400> SEQUENCE: 39
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Met Ile Arg Leu Val Gln Ala Tyr Leu Leu Glu Ala Lys Trp Thr His 1 5 10 15	
Gln Asn Tyr Lys Pro Thr Phe Glu Glu Phe Arg Asp Asn Ala Leu Pro 20 25 30	
Thr Ser Gly Tyr Ala Met Leu Ala Ile Thr Ala Phe Val Gly Met Gly 35 40 45	
Glu Val Ile Thr Pro Glu Thr Phe Lys Trp Ala Ala Ser Asp Pro Lys 50 55 60	
Ile Ile Lys Ala Ser Thr Ile Ile Cys Arg Phe Met Asp Asp Ile Ala 65 70 75 80	
Glu His Lys	
<210> SEQ ID NO 40 <211> LENGTH: 97 <212> TYPE: PRT <213> ORGANISM: Gossypium arboreum	
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Phe Asn His Arg Arg Glu Asp Asp Cys Ser Ala Ile Glu Cys Tyr Met 1 5 10 15	
Lys Gln Tyr Gly Val Thr Ala Gln Glu Ala Tyr Asn Glu Phe Asn Lys 20 25 30	
His Ile Glu Ser Ser Trp Lys Asp Val Asn Glu Glu Phe Leu Lys Pro 35 40 45	
Thr Glu Met Pro Thr Pro Val Leu Cys Arg Ser Leu Asn Leu Ala Arg 50 55 60	
Val Met Asp Val Leu Tyr Arg Glu Gly Asp Gly Tyr Thr His Val Gly 65 70 75 80	
Lys Ala Ala Lys Gly Gly Ile Thr Ser Leu Leu Ile Asp Pro Ile Gln 85 90 95	
Ile	
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tgaaac atg gca ttg cca tca gct gct atg caa tcc aac cct gaa aag Met Ala Leu Pro Ser Ala Ala Met Gln Ser Asn Pro Glu Lys 1 5 10	108
ctt aac tta ttt cac aga ttg tca agc tta ccc acc act agc ttg gaa Leu Asn Leu Phe His Arg Leu Ser Ser Leu Pro Thr Thr Ser Leu Glu 15 20 25 30	156
tat ggc aat aat cgc ttc cct ttc ttt tcc tca tct gcc aag tca cac Tyr Gly Asn Asn Arg Phe Pro Phe Phe Ser Ser Ser Ala Lys Ser His $35$ $40$ $45$	204
ttt aaa aaa cca act caa gca tgt tta tcc tca aca acc cac caa gaa Phe Lys Lys Pro Thr Gln Ala Cys Leu Ser Ser Thr Thr His Gln Glu 50 55 60	252
gtt cgt cca tta gca tac ttt cct cct act gtc tgg ggc aat cgc ttt Val Arg Pro Leu Ala Tyr Phe Pro Pro Thr Val Trp Gly Asn Arg Phe 65 70 75	300

												con	tin	uea			
-	tcc Ser 80	_					_	-		_	-		-	-		348	
-	att Ile	-	-	_			-	_	-							396	
	gat Asp															444	
	gta Val															492	
	ttc Phe															540	
	act Thr 160															588	
	tct Ser															636	
	gaa Glu															684	
	tcc Ser															732	
	ttc Phe															780	
	ctc Leu 240					_		_			_				_	828	
	gtg Val	_				-					_			-	-	876	
	att Ile															924	
	aat Asn		Val	Gln	Leu		His	Gln	Gln	Glu		Cys	Gln			972	
_	tgg Trp			-	-			-	_	-				-	-	1020	
	aga Arg 320															1068	
	gac Asp															1116	
	tca Ser			-	-			-								1164	
	cat His															1212	
	aag Lys															1260	

					gag Glu											1308
					agg Arg 420											1356
					tgg Trp											1404
					gga Gly											1452
	_			_	gga Gly	-		-						-		1500
		_	_		aat Asn				_		-			-		1548
					gac Asp 500											1596
					tgc C <b>y</b> s											1644
					gtc Val											1692
		-			gaa Glu	-	_	_		_	-					1740
					ata Ile											1788
					ggt Gly 580											1836
					gtt Val						tgaa	actca	aat a	aatto	cctttt	1889
ttca	atttt	gt a	actto	caata	aa gt	ttata	aaato	g acc	ccgto	gcac	tago	ggt	ggt (	gatta	attgta	1949
ttta	aaatt	gc (	ctttt	caaat	tt aa	atata	atgaa	a tca	agaa	attt	tata	ag				1994
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Asn	Asn	Arg 35	Phe	Pro	Phe	Phe	Ser 40	Ser	Ser	Ala	Lys	Ser 45	His	Phe	Lys	
Lys	Pro 50	Thr	Gln	Ala	Cys	Leu 55	Ser	Ser	Thr	Thr	His 60	Gln	Glu	Val	Arg	
Pro																

Leu	Thr	Phe	Asn	Pro 85	Ser	Glu	Phe	Glu	Ser 90	Tyr	Asp	Glu	Arg	Val 95	Ile
Val	Leu	Lys	L <b>y</b> s 100	Lys	Val	Lys	Asp	Ile 105	Leu	Ile	Ser	Ser	Thr 110	Ser	Asp
Ser	Val	Glu 115	Thr	Val	Ile	Leu	Ile 120	Asp	Leu	Leu	Суѕ	Arg 125	Leu	Gly	Val
Ser	Tyr 130	His	Phe	Glu	Asn	Asp 135	Ile	Glu	Glu	Leu	Leu 140	Ser	Lys	Ile	Phe
Asn 145	Ser	Gln	Pro	Asp	Leu 150	Val	Asp	Glu	Lys	Glu 155	Суѕ	Asp	Leu	Tyr	Thr 160
Ala	Ala	Ile	Val	Phe 165	Arg	Val	Phe	Arg	Gln 170	His	Gly	Phe	Lys	Met 175	Ser
Ser	Asp	Val	Phe 180	Ser	Lys	Phe	Lys	Asp 185	Ser	Asp	Gly	Lys	Phe 190	Lys	Glu
Ser	Leu	Arg 195	Gly	Asp	Ala	Lys	Gly 200	Met	Leu	Ser	Leu	Phe 205	Glu	Ala	Ser
	Leu 210					215	_				220				
225	Lys	-	-		230					235					240
_	Arg			245					250					255	
	Arg		260					265	_		_		270		
	Суѕ	275					280				_	285	_	_	
-	Val 290					295				-	300			_	_
305	Lys	_			310			_		315	_		_	_	320
	Ala			325		_			330		-			335	_
-	Ala		340					345	-				350		
	Ile	355					360					365			
	Leu 370					375	Ī	-	-		380	-			
385	Pro				390					395					400
	Glu			405					410					415	
	Tyr		420					425					430		
	Ala	435					440					445			
	Tyr 450					455		_			460				
465	Phe				470					475					480
Leu	Glu	Thr	Asn	Pro 485	Lys	Leu	Ser	Tyr	Ala 490	Ser	Gly	Ala	Phe	Ile 495	Arg

Leu Val Asn Asp Leu Thr Ser His Val Thr Glu Gln Gln Arg Gly His 500 505 510
Val Ala Ser Cys Ile Asp Cys Tyr Met Asn Gln His Gly Val Ser Lys 515 520 525
Asp Glu Ala Val Lys Ile Leu Gln Lys Met Ala Thr Asp Cys Trp Lys 530 535 540
Glu Ile Asn Glu Glu Cys Met Arg Gln Ser Gln Val Ser Val Gly His 545 550 555 560
Leu Met Arg Ile Val Asn Leu Ala Arg Leu Thr Asp Val Ser Tyr Lys 565 570 575
Tyr Gly Asp Gly Tyr Thr Asp Ser Gln Gln Leu Lys Gln Phe Val Lys 580 585 590
Gly Leu Phe Val Asp Pro Ile Ser Ile 595 600
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aga gcc aaa tct gag cgc caa atg atg tgg gtt tgc tcc aga tca ggg 147 Arg Ala Lys Ser Glu Arg Gln Met Met Trp Val Cys Ser Arg Ser Gly 30 35 40
cga acc aga gta aaa atg tcg aga gga agt ggt ggt cct ggt cct gtc 195 Arg Thr Arg Val Lys Met Ser Arg Gly Ser Gly Gly Pro Gly Pro Val 45 50 55
gta atg atg agc agc act ggc act agc aag gtg gtt tcc gag act Val Met Met Ser Ser Ser Thr Gly Thr Ser Lys Val Val Ser Glu Thr 60 65 70
tcc agt acc att gtg gat gat atc cct cga ctc tcc gcc aat tat cat  291  Ser Ser Thr Ile Val Asp Asp Ile Pro Arg Leu Ser Ala Asn Tyr His  75  80  85  90
ggc gat ctg tgg cac cac aat gtt ata caa act ctg gag aca ccg ttt 339 Gly Asp Leu Trp His His Asn Val Ile Gln Thr Leu Glu Thr Pro Phe 95 100 105
cgt gag agt tct act tac caa gaa cgg gca gat gag ctg gtt gtg aaa 387 Arg Glu Ser Ser Thr Tyr Gln Glu Arg Ala Asp Glu Leu Val Val Lys 110 115 120
att aaa gat atg ttc aat gcg ctc gga gac gga gat atc agt ccg tct  125  130  135  435  136  137
gca tac gac act gcg tgg gtg gcg agg ctg gcg acc att tcc tct gat  Ala Tyr Asp Thr Ala Trp Val Ala Arg Leu Ala Thr Ile Ser Ser Asp  140 145 150
gga tct gag aag cca cgg ttt cct cag gcc ctc aac tgg gtt ttc aac 531 Gly Ser Glu Lys Pro Arg Phe Pro Gln Ala Leu Asn Trp Val Phe Asn 155 160 165 170
aac cag ctc cag gat gga tcg tgg ggt atc gaa tcg cac ttt agt tta 579 Asn Gln Leu Gln Asp Gly Ser Trp Gly Ile Glu Ser His Phe Ser Leu

				175					180					185		
-	gat Asp	-	-			-				-		-		-	-	627
	aaa Lys															675
	aat Asn 220															723
	ata Ile															771
	ctt Leu															819
	gcc Ala															867
	atg Met	_			_	_				_	-		_			915
_	att Ile 300	_				_		_				_	_			963
	tcc Ser															1011
	ttt Phe															1059
	tat Tyr															1107
	cat His															1155
	gat Asp 380															1203
_	gac Asp	_		_		_					_			_	_	1251
	ctt Leu															1299
	aaa Lys															1347
	gaa Glu															1395
	cct Pro 460															1443
	ctt Leu															1491
ttc	aaa	gag	att	gag	tac	gtg	gtg	gag	tac	cct	tgg	cac	atg	agt	atc	1539

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Phe	Lys	Glu	Ile	Glu 495	Tyr	Val	Val	Glu	Tyr 500	Pro	Trp	His	Met	Ser 505	Ile	
	-		-	-	-	-			-			-	gac Asp 520			1587
-		_		_				-	_			-	agt Ser			1635
													caa Gln			1683
													gaa Glu			1731
													gtt Val			1779
		-			_		-			_		_	att Ile 600	-		1827
													gct Ala			1875
													gta Val			1923
	-											-	caa Gln		-	1971
													gat Asp			2019
													ccc Pro 680			2067
													gaa Glu			2115
					_				_			-	ata Ile		-	2163
			_	_								_	ggt Gl <b>y</b>			2211
													aat Asn			2259
													aaa Lys 760			2307
													gca Ala			2355
_	_	-				-			-	_	-		aag Lys			2403
													gaa Glu			2451

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aaa cca tcc aat gat atc cca atg ggt tgc aag tcc ttt att ttt aac Lys Pro Ser Asn Asp Ile Pro Met Gly Cys Lys Ser Phe Ile Phe Asn 815 820 825	2499
ctt aga ttg tgt gtc caa atc ttt tac aag ttt ata gat ggg tac gga Leu Arg Leu Cys Val Gln Ile Phe Tyr Lys Phe Ile Asp Gly Tyr Gly 830 835 840	2547
atc gcc aat gag gag att aag gac tat ata aga aaa gtt tat att gat Ile Ala Asn Glu Glu Ile Lys Asp Tyr Ile Arg Lys Val Tyr Ile Asp 845 850 855	2595
cca att caa gta tga tatatcatgt aaaacctctt tttcatgata aattgactta Pro Ile Gln Val 860	2650
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Asn Lys Ala Ile His Asp Pro Thr Asn Cys Arg Ala Lys Ser Glu Arg 20 25 30	
Gln Met Met Trp Val Cys Ser Arg Ser Gly Arg Thr Arg Val Lys Met 35 40 45	
Ser Arg Gly Ser Gly Gly Pro Gly Pro Val Val Met Met Ser Ser Ser 50 55 60	
Thr Gly Thr Ser Lys Val Val Ser Glu Thr Ser Ser Thr Ile Val Asp 65 70 75 80	
Asp Ile Pro Arg Leu Ser Ala Asn Tyr His Gly Asp Leu Trp His His 85 90 95	
Asn Val Ile Gln Thr Leu Glu Thr Pro Phe Arg Glu Ser Ser Thr Tyr 100 105 110	
Gln Glu Arg Ala Asp Glu Leu Val Val Lys Ile Lys Asp Met Phe Asn 115 120 125	
Ala Leu Gly Asp Gly Asp Ile Ser Pro Ser Ala Tyr Asp Thr Ala Trp 130 135 140	
Val Ala Arg Leu Ala Thr Ile Ser Ser Asp Gly Ser Glu Lys Pro Arg 145 150 155 160	
Phe Pro Gln Ala Leu Asn Trp Val Phe Asn Asn Gln Leu Gln Asp Gly 165 170 175	
Ser Trp Gly Ile Glu Ser His Phe Ser Leu Cys Asp Arg Leu Leu Asn 180 185 190	
Thr Thr Asn Ser Val Ile Ala Leu Ser Val Trp Lys Thr Gly His Ser 195 200 205	
Gln Val Gln Gln Gly Ala Glu Phe Ile Ala Glu Asn Leu Arg Leu Leu 210 215 220	
Asn Glu Glu Asp Glu Leu Ser Pro Asp Phe Gln Ile Ile Phe Pro Ala 225 230 235 240	
Leu Leu Gln Lys Ala Lys Ala Leu Gly Ile Asn Leu Pro Tyr Asp Leu 245 250 255	
Pro Phe Ile Lys Tyr Leu Ser Thr Thr Arg Glu Ala Arg Leu Thr Asp 260 265 270	
Val Ser Ala Ala Ala Asp Asn Ile Pro Ala Asn Met Leu Asn Ala Leu 275 280 285	

Glu	Gly 290	Leu	Glu	Glu	Val	Ile 295	Asp	Trp	Asn	Lys	Ile 300	Met	Arg	Phe	Gln
Ser 305	Lys	Asp	Gly	Ser	Phe 310	Leu	Ser	Ser	Pro	Ala 315	Ser	Thr	Ala	Суѕ	Val 320
Leu	Met	Asn	Thr	Gly 325	Asp	Glu	Lys	Суѕ	Phe 330	Thr	Phe	Leu	Asn	Asn 335	Leu
Leu	Asp	Lys	Phe 340	Gly	Gly	Сув	Val	Pro 345	Cys	Met	Tyr	Ser	Ile 350	Asp	Leu
Leu	Glu	Arg 355	Leu	Ser	Leu	Val	Asp 360	Asn	Ile	Glu	His	Leu 365	Gly	Ile	Gly
Arg	His 370	Phe	Lys	Gln	Glu	Ile 375	Lys	Gly	Ala	Leu	Asp 380	Tyr	Val	Tyr	Arg
His 385	Trp	Ser	Glu	Arg	Gly 390	Ile	Gly	Trp	Gly	Arg 395	Asp	Ser	Leu	Val	Pro 400
Asp	Leu	Asn	Thr	Thr 405	Ala	Leu	Gly	Leu	Arg 410	Thr	Leu	Arg	Met	His 415	Gly
Tyr	Asn	Val	Ser 420	Ser	Asp	Val	Leu	Asn 425	Asn	Phe	Lys	Asp	Glu 430	Asn	Gly
Arg	Phe	Phe 435	Ser	Ser	Ala	Gly	Gln 440	Thr	His	Val	Glu	Leu 445	Arg	Ser	Val
Val	Asn 450	Leu	Phe	Arg	Ala	Ser 455	Asp	Leu	Ala	Phe	Pro 460	Asp	Glu	Arg	Ala
Met 465	Asp	Asp	Ala	Arg	L <b>y</b> s 470	Phe	Ala	Glu	Pro	Tyr 475	Leu	Arg	Glu	Ala	Leu 480
Ala	Thr	Lys	Ile	Ser 485	Thr	Asn	Thr	Lys	Leu 490	Phe	Lys	Glu	Ile	Glu 495	Tyr
Val	Val	Glu	Tyr 500	Pro	Trp	His	Met	Ser 505	Ile	Pro	Arg	Leu	Glu 510	Ala	Arg
Ser	Tyr	Ile 515	Asp	Ser	Tyr	Asp	Asp 520	Asn	Tyr	Val	Trp	Gln 525	Arg	Lys	Thr
Leu	<b>Ty</b> r 530	Arg	Met	Pro	Ser	Leu 535	Ser	Asn	Ser	Lys	C <b>y</b> s 540	Leu	Glu	Leu	Ala
L <b>y</b> s 545	Leu	Asp	Phe	Asn	Ile 550	Val	Gln	Ser	Leu	His 555	Gln	Glu	Glu	Leu	L <b>y</b> s 560
Leu	Leu	Thr	Arg	Trp 565	Trp	Lys	Glu	Ser	Gl <b>y</b> 570	Met	Ala	Asp	Ile	Asn 575	Phe
Thr	Arg	His	Arg 580	Val	Ala	Glu	Val	T <b>y</b> r 585	Phe	Ser	Ser	Ala	Thr 590	Phe	Glu
Pro	Glu	T <b>y</b> r 595	Ser	Ala	Thr	Arg	Ile 600	Ala	Phe	Thr	Lys	Ile 605	Gly	Cys	Leu
Gln	Val 610	Leu	Phe	Asp	Asp	Met 615	Ala	Asp	Ile	Phe	Ala 620	Thr	Leu	Asp	Glu
Leu 625	Lys	Ser	Phe	Thr	Glu 630	Gly	Val	Lys	Arg	Trp 635	Asp	Thr	Ser	Leu	Leu 640
His	Glu	Ile	Pro	Glu 645	Cys	Met	Gln	Thr	C <b>y</b> s 650	Phe	Lys	Val	Trp	Phe 655	Lys
Leu	Met	Glu	Glu 660	Val	Asn	Asn	Asp	Val 665	Val	Lys	Val	Gln	Gly 670	Arg	Asp
Met	Leu	Ala 675	His	Ile	Arg	Lys	Pro 680	Trp	Glu	Leu	Tyr	Phe 685	Asn	Cys	Tyr
Val	Gln 690	Glu	Arg	Glu	Trp	Leu 695	Glu	Ala	Gly	Tyr	Ile 700	Pro	Thr	Phe	Glu

Glu 705	Tyr	Leu	Lys	Thr	<b>Ty</b> r 710	Ala	Ile	Ser	Val	Gly 715	Leu	Gly	Pro	Cys	Thr 720					
Leu	Gln	Pro	Ile	Leu 725	Leu	Met	Gly	Glu	Leu 730	Val	Lys	Asp	Asp	Val 735	Val					
Glu	Lys	Val	His 740	Tyr	Pro	Ser	Asn	Met 745	Phe	Glu	Leu	Val	Ser 750	Leu	Ser					
Trp	Arg	Leu 755	Thr	Asn	Asp	Thr	<b>Lys</b> 760		Tyr	Gln	Ala	Glu 765	Lys	Ala	Arg					
Gly	Gln 770		Ala	Ser	Gly	Ile 775		Cys	Tyr	Met	L <b>y</b> s 780	Asp	Asn	Pro	Gly					
Ala 785		Glu	Glu	Asp	Ala 790		Lys	His	Ile	C <b>y</b> s 795		Val	Val	Asp	Arg 800					
	Leu	Lys	Glu	Ala 805	Ser	Phe	Glu	Tyr	Phe		Pro	Ser	Asn	Asp 815	_					
Pro	Met	Gly			Ser	Phe	Ile			Leu	Arg	Leu			Gln					
Ile	Phe		820 Lys	Phe	Ile	Asp		825 Tyr	Gly	Ile	Ala		830 Glu	Glu	Ile					
L <b>y</b> s	_	835 <b>Ty</b> r	Ile	Arg	Lys		840 Tyr	Ile	Asp	Pro		845 Gln	Val							
	850					855					860									
<213 <220 <221	> OR > FE > NA > LO	ATUF ME/F CATI	SM: RE: REY: ON:	CDS (2)	es gr	2347)	)	a_his	ahol	lono		1								
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			ICE:		LION	. 11-6	.ipiic		, abo	rene	synt	nase	9							
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<400 g gg Gl tac Tyr aat Asn	> SE t ta y Ty aga Arg cca Pro gac Asp	gaa Glu gct Ala 35	CGC Arg 20 att Ile	45  tt great Value	tg ca al H: 5 gag Glu gga	at to is Se gtc Val gat Asp gcg Ala	ct cter Leu gga Gly 40	gtt Val 25 gaa Glu	aa toys Se agc Ser tca Ser	ca co er Pi 10 gag Glu atg Met	att Ile att Ile att Ile	at agyr II aaa Lys act Thr 45	tt ga le As gtg Val 30 cca Pro	atg Met tct Ser	er Ser 15 ctt Leu gct Ala	9.	, 5			
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					aaa Lys											577
_		_	_		tct Ser						-		-	-		625
					gtt Val											673
					tgc Cys 230											721
	-		_		agt Ser		_		_					-		769
					gat Asp											817
					atc Ile											865
					tac Tyr											913
	_				ata Ile 310	-	-					-	_			961
					cat His											1009
					aat Asn											1057
		_	_	_	agc Ser	_	_				_	_				1105
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					tat Tyr											1345
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								ata Ile 575	1729			
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-	_		 _	-	-		 _	agc Ser	2017			
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gat Asp 770								taa	2350			

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2410 2424

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Tyr Asp Thr Ala Trp Val Ala Arg Val Pro Ala Ile Asp Gly Ser Ala 50 55 60															Ala
50 55 60  Arg Pro Gln Phe Pro Gln Thr Val Asp Trp Ile Leu Lys Asn Gln Leu 65 70 75 80															
Lys	Asp	Gly	Ser	Trp 85	Gly	Ile	Gln	Ser	His 90	Phe	Leu	Leu	Ser	Asp 95	Arg
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Gly	Asp	Leu 115	Gln	Val	Glu	Gln	Gl <b>y</b> 120	Ile	Glu	Phe	Ile	L <b>y</b> s 125	Ser	Asn	Leu
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Glu 145	Ile	Ile	Phe	Pro	Ser 150	Leu	Leu	Arg	Glu	Ala 155	Gln	Ser	Leu	Arg	Leu 160
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Gln	Glu	Arg	Leu 180	Ala	Lys	Leu	Ser	Arg 185	Glu	Glu	Ile	Tyr	Ala 190	Val	Pro
Ser	Pro	Leu 195	Leu	Tyr	Ser	Leu	Glu 200	Gly	Ile	Gln	Asp	Ile 205	Val	Glu	Trp
Glu	Arg 210	Ile	Met	Glu	Val	Gln 215	Ser	Gln	Asp	Gly	Ser 220	Phe	Leu	Ser	Ser
Pro 225	Ala	Ser	Thr	Ala	C <b>y</b> s 230	Val	Phe	Met	His	Thr 235	Gly	Asp	Ala	Lys	Cys 240
Leu	Glu	Phe	Leu	Asn 245	Ser	Val	Met	Ile	L <b>y</b> s 250	Phe	Gly	Asn	Phe	Val 255	Pro
Суѕ	Leu	Tyr	Pro 260	Val	Asp	Leu	Leu	Glu 265	Arg	Leu	Leu	Ile	Val 270	Asp	Asn
Ile	Val	Arg 275	Leu	Gly	Ile	Tyr	Arg 280	His	Phe	Glu	Lys	Glu 285	Ile	Lys	Glu
Ala	Leu 290	Asp	Tyr	Val	Tyr	Arg 295	His	Trp	Asn	Glu	Arg 300	Gly	Ile	Gly	Trp
Gly 305	Arg	Leu	Asn	Pro	Ile 310	Ala	Asp	Leu	Glu	Thr 315	Thr	Ala	Leu	Gly	Phe 320
Arg	Leu	Leu	Arg	Leu 325	His	Arg	Tyr	Asn	Val 330	Ser	Pro	Ala	Ile	Phe 335	Asp
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Ala	Phe 370	Pro	Gly	Glu	Asn	Ile 375	Leu	Asp	Glu	Ala	L <b>y</b> s 380	Ser	Phe	Ala	Thr
L <b>y</b> s 385	Tyr	Leu	Arg	Glu	Ala 390	Leu	Glu	Lys	Ser	Glu 395	Thr	Ser	Ser	Ala	Trp 400
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Ser	Trp	His	Ala 420	Ser	Val	Pro	Arg	Val 425	Glu	Ala	Lys	Arg	Tyr 430	Cys	Gln
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Glu	Ala 610	Glu	Trp	Leu	Ala	Ala 615	Glu	Tyr	Val	Pro	Thr 620	Leu	Asp	Glu	Tyr
Ile 625	Lys	Asn	Gly	Ile	Thr 630	Ser	Ile	Gly	Gln	Arg 635	Ile	Leu	Leu	Leu	Ser 640
Gly	Val	Leu		Met 645		Gly		Leu			Gln	Glu		Leu 655	
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Ile	Ser	Arg 675	Leu	Ala	Asp	Asp	Thr 680	Lys	Thr	Tyr	Lys	Ala 685	Glu	Lys	Ala
Arg	Gl <b>y</b> 690	Glu	Leu	Ala	Ser	Ser 695	Ile	Glu	Суѕ	Tyr	Met 700	Lys	Asp	His	Pro
Glu 705	Сув	Thr	Glu	Glu	Glu 710	Ala	Leu	Asp	His	Ile 715	Tyr	Ser	Ile	Leu	Glu 720
Pro	Ala	Val	Lys	Glu 725	Leu	Thr	Arg	Glu	Phe 730	Leu	Lys	Pro	Asp	Asp 735	Val
Pro	Phe	Ala	C <b>y</b> s 740	Lys	Lys	Met	Leu	Phe 745	Glu	Glu	Thr	Arg	Val 750	Thr	Met
Val	Ile	Phe 755	Lys	Asp	Gly	Asp	Gly 760	Phe	Gly	Val	Ser	L <b>y</b> s 765	Leu	Glu	Val
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		g ctc ctt caa aag ctt att cag gag 144 u Leu Leu Gln Lys Leu Ile Gln Glu 0 45
		a atg gaa atg gat gat ggc gat cat 192 u Met Glu Met Asp Asp Gl <b>y</b> Asp His 60
		c gtt gac act ttg gaa tgc ctg gga 240 e Val Asp Thr Leu Glu Cys Leu Gly 75 80
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Val Tyr Arg T		a ggt atc ggg gag gga tca aga gat 336 s Gly Ile Gly Glu Gly Ser Arg Asp 105 110
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		c aac ttt act ggt gaa gaa gga aga 480 s Asn Phe Thr Gly Glu Glu Gly Arg 155 160
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Ile Ser Phe I		g atg gaa gag gcc aag gca ttc aca 576 l Met Glu Glu Ala Lys Ala Phe Thr 185 190
		a gct gga cac ggg gat gtg act gac 624 u Ala Gly His Gly Asp Val Thr Asp 0 205
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		g gag gca agg agc ttt ctc gaa ata 720 p Glu Ala Arg Ser Phe Leu Glu Ile 235 240
		c aag tcg aat atc aac caa aaa atg 768 u Lys Ser Asn Ile Asn Gln Lys Met 250 255
ttg aag tta g	gcc aaa ttg gac tto	c aat att ctg caa tgc aaa cat cac 816

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	ctg Leu 290				-	-	-			-					-	912			
	atg Met															960			
	aaa Lys															1008			
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Trp	gat Asp	Leu 355	Ser	Leu	Thr	Åsp	Asp 360	Leu	Pro	Āsp	Tyr	Ile 365	Lys	Ile	Āla	1104			
Phe	Gln 370	Phe	Phe	Phe	Asn	Thr 375	Val	Asn	Glu	Leu	Ile 380	Val	Glu	Ile	Val	1152			
L <b>y</b> s 385	Arg	Gln	Gly	Arg	Asp 390	Met	Thr	Thr	Ile	Val 395	Lys	Asp	Cys	Trp	Lys 400	1200			
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Lys	ggc	Ile 515	Leu	Asn	Arg	Ser	Leu 520	Glu	Glu	Phe	Asn	Trp 525	Glu	Phe	Met	1632			
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Tyr Gly Ala Pro Ala Tyr Tyr Glu Leu Leu Gln Lys Leu Ile Gln Glu 35 40 45													
Ile Lys His Leu Leu Thr Glu Met Glu Met Asp Asp Gly Asp His 50 55 60													
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Phe Gln Phe Phe Phe Asn Thr Val	usn Glu Leu Ile Val Glu Ile Val 380	
Lys Arg Gln Gly Arg Asp Met Thr	hr Ile Val Lys Asp Cys Trp Lys 395 400	
Arg Tyr Ile Glu Ser Tyr Leu Gln 405	lu Ala Glu Trp Ile Ala Thr Gly 410 415	
His Ile Pro Thr Phe Asn Glu Tyr		
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Leu Pro Asp Asn Ile Leu Glu Gln 450 455		
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Lys Gly Ile Leu Asn Arg Ser Leu	<u>-</u>	
515 520  Lys Gln Asp Ser Val Pro Met Cys		
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gac gat cgc ata cag tct ctc aac Asp Asp Arg Ile Gln Ser Leu Asn 35		
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agt gac atg gac gat agc tgc aat Ser Asp Met Asp Asp Ser Cys Asn 65 70		

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			-		aaa Lys			-					-	-		336	
					gga Gly											384	
					ttg Leu											432	
	-				gtc Val	_				_	-	_			_	480	
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					atg Met											576	
					gtg Val											624	
					tta Leu											672	
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					gag Glu 245											768	
					aag Lys											816	
					aat Asn											864	
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					tac Tyr											960	
					ttt Phe 325											1008	
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					atc Ile											1104	
					ctc Leu											1152	
					aat Asn											1200	

_													con	tin	uea		
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	y					gcc Ala 405											1248
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				_		tac Tyr	_									_	1344
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	е.					caa Gln											1440
	.e	-	_	_		agg Arg 485		-	-	_		_	-		_		1488
						gat Asp											1536
						gta Val											1584
						ctg Leu											1632
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Th	ır	Glu	Ser	Ser 20	Ile	Thr	Ser	Asn	Arg 25	His	Gly	Asn	Met	Trp 30	Glu	Asp	
As	p.	Arg	Ile 35	Gln	Ser	Leu	Asn	Ser 40	Pro	Tyr	Gly	Ala	Pro 45	Ala	Tyr	Gln	
G1		Arg 50	Ser	Glu	Lys	Leu	Ile 55	Glu	Glu	Ile	Lys	Leu 60	Leu	Phe	Leu	Ser	
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Gln	Pro	Glu	Ile 100	Lys	Leu	Ala	Leu	Asp 105	Tyr	Val	Tyr	Arg	Cys 110	Trp	Asn
Glu	Arg	Gly 115	Ile	Gly	Glu	Gly	Ser 120	Arg	Asp	Ser	Leu	Lys 125	Lys	Asp	Leu
Asn	Ala 130	Thr	Ala	Leu	Gly	Phe 135	Arg	Ala	Leu	Arg	Leu 140	His	Arg	Tyr	Asn
Val 145	Ser	Ser	Gly	Val	Leu 150	Glu	Asn	Phe	Arg	Asp 155	Asp	Asn	Gly	Gln	Phe 160
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His	Val	Arg	C <b>y</b> s 180	Met	Leu	Ser	Leu	Ser 185	Arg	Ala	Ser	Asn	Ile 190	Leu	Phe
Pro	Gly	Glu 195	Lys	Val	Met	Glu	Glu 200	Ala	Lys	Ala	Phe	Thr 205	Thr	Asn	Tyr
Leu	L <b>y</b> s 210	Lys	Val	Leu	Ala	Gly 215	Arg	Glu	Ala	Thr	His 220	Val	Asp	Glu	Ser
Leu 225	Leu	Gly	Glu	Val	L <b>y</b> s 230	Tyr	Ala	Leu	Glu	Phe 235	Pro	Trp	His	Cys	Ser 240
Val	Gln	Arg	Trp	Glu 245	Ala	Arg	Ser	Phe	Ile 250	Glu	Ile	Phe	Gly	Gln 255	Ile
Asp	Ser	Glu	Leu 260	Lys	Ser	Asn	Leu	Ser 265	Lys	Lys	Met	Leu	Glu 270	Leu	Ala
Lys	Leu	<b>Asp</b> 275	Phe	Asn	Ile	Leu	Gln 280	Суѕ	Thr	His	Gln	Lys 285	Glu	Leu	Gln
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<b>Ty</b> r 305	Arg	Lys	Суѕ	Tyr	Val 310	Glu	Phe	Tyr	Phe	Trp 315	Met	Ala	Ala	Ala	Ile 320
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Leu 385		Thr			Glu 390				Glu			Lys	Ala		Gly 400
Gln	Asp	Met	Ala	Ala 405	Tyr	Ile	Arg	Lys	Asn 410	Ala	Trp	Glu	Arg	<b>Ty</b> r 415	Leu
Glu	Ala	Tyr	Leu 420	Gln	Asp	Ala	Glu	Trp 425	Ile	Ala	Thr	Gly	His 430	Val	Pro
Thr	Phe	Asp 435	Glu	Tyr	Leu	Asn	Asn 440	Gly	Thr	Pro	Asn	Thr 445	Gly	Met	Сув
Val	Leu 450	Asn	Leu	Ile	Pro	Leu 455	Leu	Leu	Met	Gly	Glu 460	His	Leu	Pro	Ile
Asp 465	Ile	Leu	Glu	Gln	Ile 470	Phe	Leu	Pro	Ser	Arg 475	Phe	His	His	Leu	Ile 480
Glu	Leu	Ala	Ser	Arg 485	Leu	Val	Asp	Asp	Ala 490	Arg	Asp	Phe	Gln	Ala 495	Glu
Lys	Asp	His	Gly 500	Asp	Leu	Ser	Сув	Ile 505	Glu	Сув	Tyr	Leu	L <b>y</b> s 510	Asp	His
Pro	Glu	Ser	Thr	Val	Glu	Asp	Ala	Leu	Asn	His	Val	Asn	Gly	Leu	Leu

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Gly	Asn 530	Cys	Leu	Leu	Glu	Met 535	Asn	Trp	Lys	Phe	Leu 540	Lys	Lys	Gln	Asp	
Ser 545	Val	Pro	Leu	Ser	C <b>y</b> s 550	Lys	Lys	Tyr	Ser	Phe 555	His	Val	Leu	Ala	Arg 560	
Ser	Ile	Gln	Phe	Met 565		Asn	Gln	Gly	Asp 570	Gly	Phe	Ser	Ile	Ser 575	Asn	
Lys	Val	Ile	L <b>y</b> s 580	Asp	Gln	Val	Gln	<b>Lys</b> 585	Val	Leu	Ile	Val	Pro 590	Val	Pro	
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													gga Gl <b>y</b>			100
													gtt Val			148
-							_		_	_		_	act Thr	_	-	196
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						Gln		Asp	Asn	Asp	Asn		ctt Leu			340
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Pro 280	Lys	Tyr	Ser	Arg	Ala 285 gac	Arg	Lys	Met	Met	Thr 290	Lys	Val	Leu	Asn	Leu 295	964
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gat	tca	ata	315 caa	cca	Asp tat	atg	aga	320 cct	gct	tat	caa	gct	325 ctt	cta	gac	1060
att	tac	330 agt	gaa	atg	gaa	caa	335 gtg	ttg	tcc	aaa	gaa	340 ggt	aaa	ctg	gac	1108
cgt	345 gta	tac	tat	gca	Glu aaa Lys	350 aat	gag	atg	aaa	aag	355 ttg	gtg	aga	gcc	tat	1156
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			tgc		gtc Val			gaa					cac			1300
					aat Asn											1348
	-	-	-	_	aac Asn 445	-		-			-	-	-		-	1396
					tca Ser											1444
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					cga Arg											1588

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Leu Tyr Lys Glu Lys Asp Thr Tyr Ser Thr Ala Lys Gly Lys Leu Lys 520 525 530 535  aac atg att aat cca ata cta att gaa tct gtc aaa ata taa Asn Met Ile Asn Pro Ile Leu Ile Glu Ser Val Lys Ile 540 545  atataatgct gaaattgcac cttcatcatc caactattca cagcaaaata aggcatataa	1678 1738 1798
Asn Met Ile Asn Pro Ile Leu Ile Glu Ser Val Lys Ile 540 545  atataatgct gaaattgcac cttcatcatc caactattca cagcaaaata aggcatataa	1738 1798
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	1858
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Asp Ala Met Gln Arg Leu Gly Val Ala Tyr His Phe Asp Asn Glu Ile 65 70 75 80	
Glu Thr Ser Ile Gln Asn Ile Phe Asp Ala Ser Ser Lys Gln Asn Asp 85 90 95	
Asn Asp Asn Asn Leu Tyr Val Val Ser Leu Arg Phe Arg Leu Val Arg	
Gln Gln Gly His Tyr Met Ser Ser Asp Val Phe Lys Gln Phe Thr Asn 115 120 125	
Gln Asp Gly Lys Phe Lys Glu Thr Leu Thr Asn Asp Val Gln Gly Leu 130 135 140	
Leu Ser Leu Tyr Glu Ala Ser His Leu Arg Val Arg Asn Glu Glu Ile 145 150 155 160	
Leu Glu Glu Ala Leu Thr Phe Thr Thr His Leu Glu Ser Ile Val 165 170 175	
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Ala Leu Thr Gln Pro Ile Arg Met Thr Leu Pro Arg Met Gly Ala Arg 195 200 205	
Lys Tyr Ile Ser Ile Tyr Glu Asn Asn Asp Ala His His His Leu Leu 210 215 220	
Leu Lys Phe Ala Lys Leu Asp Phe Asn Met Leu Gln Lys Phe His Gln 225 230 235 240	
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Asn Lys Tyr Pro Tyr Ala Arg Asp Arg Leu Val Glu Cys Tyr Phe Trp	

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Ala 305	Tyr	Ala	Thr	Phe	Asp 310	Glu	Leu	Val	Thr	Phe 315	Asn	Asp	Ala	Ile	Gln 320					
Arg	Trp	Asp	Ala	Asn 325	Ala	Ile	Asp	Ser	Ile 330	Gln	Pro	Tyr	Met	Arg 335	Pro					
Ala	Tyr	Gln	Ala 340	Leu	Leu	Asp	Ile	Tyr 345	Ser	Glu	Met	Glu	Gln 350	Val	Leu					
Ser	Lys	Glu 355	Gly	Lys	Leu	Asp	Arg 360	Val	Tyr	Tyr	Ala	<b>Lys</b> 365	Asn	Glu	Met					
Lys	L <b>y</b> s 370	Leu	Val	Arg	Ala	<b>Ty</b> r 375	Phe	Lys	Glu	Thr	Gln 380	Trp	Leu	Asn	Asp					
C <b>y</b> s 385	Asp	His	Ile	Pro	Lys 390	Tyr	Glu	Glu	Gln	Val 395	Glu	Asn	Ala	Ile	Val 400					
Ser	Ala	Gly	Tyr	Met 405	Met	Ile	Ser	Thr	Thr 410	Cys	Leu	Val	Gly	Ile 415	Glu					
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Gly	His 450	Glu	Asp	Glu	Gln	Glu 455	Arg	Gly	His	Val	Ala 460	Ser	Leu	Ile	Glu					
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Phe	Leu	Lys	Glu	Val 485	Thr	Asn	Ala	Trp	Lys 490	Asp	Ile	Asn	Lys	Gln 495	Phe					
Ser	Arg	Pro	Thr 500	Glu	Val	Pro	Met	Phe 505	Val	Leu	Glu	Arg	Val 510	Leu	Asn					
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				aga Arg									676
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				ctt Leu									772
				gat Asp 255									820
				gcc Ala					_	-	-		868
				aaa Lys									916
				cca Pro									964
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caa gtg tgc tat ttg gca cta tac aac ttc gtt tct gag cgg gct tac Gln Val Cys Tyr Leu Ala Leu Tyr Asn Phe Val Ser Glu Arg Ala Tyr 380 385 390	1204
gat att cta aaa gat caa cat ttc aac agc atc cca tat tta cag aga Asp Ile Leu Lys Asp Gln His Phe Asn Ser Ile Pro Tyr Leu Gln Arg 395 400 405	1252
tcg tgg gta agt ttg gtt gaa gga tat ctt aag gag gca tac tgg tac Ser Trp Val Ser Leu Val Glu Gly Tyr Leu Lys Glu Ala Tyr Trp Tyr 410 415 420 425	1300
tac aat ggc tat aaa cca agc ttg gaa gaa tat ctc aac aac gcc aag Tyr Asn Gly Tyr Lys Pro Ser Leu Glu Glu Tyr Leu Asn Asn Ala Lys 430 435 440	1348
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Asn Phe Glu Arg Lys Pro Ser Lys Ala Trp Ser Thr Ser Cys Thr Ala  $20 \\ 25 \\ 30$ 

Pro	Ala	Ala 35	Arg	Leu	Arg	Ala	Ser 40	Ser	Ser	Leu	Gln	Gln 45	Glu	Lys	Pro
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Asn 65	Tyr	Ile	Gln	Ser	Leu 70	Asn	Thr	Pro	Tyr	<b>Lys</b> 75	Glu	Gln	Arg	His	Phe 80
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Lys	Met	Glu	Ala 100	Ile	Gln	Gln	Leu	Glu 105	Leu	Ile	Asp	Asp	Leu 110	Gln	Tyr
Leu	Gly	Leu 115	Ser	Tyr	Phe	Phe	Gln 120	Asp	Glu	Ile	Lys	Gln 125	Ile	Leu	Ser
Ser	Ile 130	His	Asn	Glu	Pro	Arg 135	Tyr	Phe	His	Asn	Asn 140	Asp	Leu	Tyr	Phe
Thr 145	Ala	Leu	Gly	Phe	Arg 150	Ile	Leu	Arg	Gln	His 155	Gly	Phe	Asn	Val	Ser 160
Glu	Asp	Val	Phe	Asp 165	Cys	Phe	Lys	Ile	Glu 170	Lys	Cys	Ser	Asp	Phe 175	Asn
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Ser	Phe	Leu 195	Leu	Arg	Glu	Gly	Glu 200	Asp	Thr	Leu	Glu	Leu 205	Ala	Arg	Arg
Phe	Ser 210	Thr	Arg	Ser	Leu	Arg 215	Glu	Lys	Phe	Asp	Glu 220	Gly	Gly	Asp	Glu
Ile 225	Asp	Glu	Asp	Leu	Ser 230	Ser	Trp	Ile	Arg	His 235	Ser	Leu	Asp	Leu	Pro 240
Leu	His	Trp	Arg	Val 245	Gln	Gly	Leu	Glu	Ala 250	Arg	Trp	Phe	Leu	Asp 255	Ala
Tyr	Ala	Arg	Arg 260	Pro	Asp	Met	Asn	Pro 265	Leu	Ile	Phe	Lys	Leu 270	Ala	Lys
Leu	Asn	Phe 275	Asn	Ile	Wal					_		a1	T	T	Asp
Ile		2,0			vai	Gln	Ala 280	Thr	Tyr	Gln	Glu	285	Leu	ьуѕ	
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Val 305	290	Arg	_	-	Asn	Ser 295	280 Ser	Cys	Leu	Ala	Glu 300	285 Lys	Leu	_	
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305 Glu Thr	290 Arg Pro	Arg Asp His	Arg Gln Thr 340	Ile Tyr 325	Asn Val 310 Ser Ile	Ser 295 Glu Tyr Asp	280 Ser Cys Gln Asp	Cys Phe Arg Val 345	Leu Phe Lys 330	Ala Trp 315 Met	Glu 300 Ala Ala Val	285 Lys Ile Ala Tyr	Leu Ala Val Gly 350	Pro Ala Ile 335	Phe 320 Ile
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Ala																
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Phe	Thr	Asp	Asp	Leu	Val 550		Ala	Ala	Ala	Asn 555		Ala	Arg	Ala	Ala 560	
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	p G												gta Val			719
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	r S												gag Glu			1199
	1 E												ttg Leu			1247
	-				-		_	-					aga Arg	-		1295
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													ccg Pro			1391
	r I												ctg Leu			1439

got cty gaa aat gty gat got tit gac aan tyg got tit aan aay aat 1607 Alai Leu Glu Aan Val Aap Ala Phe Aap 129 Trp Ala Phe Lys Lya Aan Alai Cau Glu Aan Val Aap Ala Phe Aap 129 Trp Ala Phe Lys Lya Aan Ado 485 Ado 1485 Ado																		 	 
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did by a Tyr Leu Giu Leu Ala Lye Leu Aap Phe Aan Lys Val Gin Ser 545         545         255         1757         172	-		Trp					Val		_	_		Tyr		_		1631		
Ile His Gin Thr Giu Leu Gin Asp Leu Arg Arg Trp Trp Lys Ser Ser 575  ggt ttc acg gat ctg aat ttc act cgt gag cgt gtg acg gaa ata tat 1775  gly Phe Thr Asp Leu Asn Phe Thr Arg Giu Arg Val Thr Giu Ile Tyr 580  ttc tca ccg gca tcc ttt atc ttt gag ccc gag ttt tct aag tgc aga 1823  Phe Ser Pro Ala Ser Phe Ile Phe Giu Pro Giu Phe Ser Lys Cys Arg 605  gag gtt tat aca aaa act tcc aat ttc act gtg att ttt aga gat ctt 1871  Glu Val Tyr Thr Lys Thr Ser Asn Phe Thr Val Ile Leu Asp Asp Leu 615  aaa aga gcc cat gga tct tta gac gat ctt aag tgt tc aca ca aca acg 1919  gag gt act ttt gac gat ctt aga gat ctt ang hasp Leu Ser Cys Arg 625  gaa aga tyg gat cta tca cta gtg gac caa atg cca caa caa acg 1919  gtc aaa aga tyg gat cta tca cta gtg gac caa atg cca caa caa acg 1916  aaa ata tgt ttt gtg ggt tct tac act act act act gta gat ata bys Ile Ala Lys 665  aaa aga tyg gag agg cg gg g	-	Lys			_		Ala		_	_		Asn	_		_		1679		
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Phe Ser Pro Ala Ser Phe IIe Phe Glu Pro Glu Phe Ser Lys Cye Arg 595 595 595 600 600 605 605 605 605 605 600 600 60					Leu					Glu					Ile		1775		
dilu Val Tyr Thr Lys Thr Ser Asn Phe Thr Val Ile Leu Asp Asp Leu       610       620       1919         tat gac gac cat gga tct tta gac gat ctt aag ttg tc aca gaa tca       1919       1919         Tyr Asp Ala His Gly Ser Leu Asp Asp Leu Lys Leu Phe Thr Glu Ser 625       630       1919         gtc aaa aga tgg gat cta cta cta gtg gac caa atg cca caa atg cca caa atg caa ata tgt ttt gtg ggt ttc tac aat act ttt aat gat ata gca aaa Lys Ile Cys Phe Val Gly Phe Tyr Asn Thr Phe Asn Asp Ile Ala Lys 660       2015         gaa gga cgt gag agg caa ggg cgc gat gtg cta gtg cta gtg cta att caa aat caa caa atg caa gga cgc gaa ggc gad atg gtg cta gtg trac att caa aat caa act caa atg caa gga gd day and caa ctt gaa gct tac acg aaa gaa gca gaa tgg tct val atg caa atg caa gat caa atg caa gat tag caa atg caa atg caa atg caa atg caa atg caa atg caa gat caa atg caa atg caa atg caa gat caa atg caa gat caa ata tg caa caa ata tag caa caa caa taa gaa ata caa ata gaa atg caa gat caa atg caa gat caa ata caa ata gaa atg caa atg caa gat caa atg caa gat caa atg caa gat gaa gaa gaa gaa gaa gaa gaa gaa g				Āla					Glu					Lys			1823		
Tyr Asp Ala His Gly Ser Leu Asp Asp Leu Lys Leu Phe Thr Glu Ser 630  Gas aga aga tgg gat cta tca cta gtg gac caa atg cca caa caa atg Val Lys Arg Trp Asp Leu Ser Leu Val Asp Gln Met Pro Gln Gln Met 640  Gas aga ata tgt ttt gtg ggt ttc tac aat act ttt aat gat ata gca aaa Lys Ile Cys Phe Val Gly Phe Tyr Asn Thr Phe Asn Asp Ile Ala Lys 660  Gaa gga cgt gag agg caa ggg cgc gat gtg cta ggc tac att caa aat Glu Gly Arg Gln Ag Gln Gly Arg Asp Val Leu Gly Tyr Ile Gln Asn 675  gat tgg aaa gtc caa ctt gaa gct tac acg aaa gaa gca gaa tgg ttc Val Trp Lys Val Gln Leu Glu Ala Tyr Thr Lys Glu Ala Glu Trp Ser 690  Gaa gct aaa tat gtg cca tcc ttc aat gaa tac ata gag aat gcg agt Glu Ala Lys Tyr Val Pro Ser Phe Asn Glu Tyr Ile Gln Asn Ala Ser 705  gtg tca ata gca ttg gga aca gtc gtt ctc att agg act tac at gag act gcg agt Glu Ala Lys Tyr Val Pro Ser Phe Asn Glu Tyr Ile Glu Asn Ala Ser 705  gtg tca ata gca ttg gga aca gtc gtt ctc att agt gcc tct ttc act Val Ser Ile Ala Leu Gly Thr Val Val Leu Ile Ser Ala Leu Phe Thr 775  ggg gag gtt ctt aca gat gaa gta ctc tcc aaa att gat cgc gaa tct Gly Glu Val Leu Thr Asp Glu Val Leu Ser Lys Ile Asp Arg Glu Ser 776  aga ttt ctc caa ctc atg ggc tta aca ggg cgt ttg tg gt aat gac Arg Phe Leu Gln Leu Met Gly Leu Thr 760  caa tgt tat atg aag gac cat cct aaa gar gag gt gg tg ct tct gcc ata Lys Try Cln Ala Glu Arg Cly Gln Gly Glu Val Ala Ser Ala Ile 770  caa tgt tat atg aag gac cat cct aaa att ctc taaa gar gaa gcc gaa gcc Arg ttat atg aag gac cat cct aaa gar gag gtg gct tct gcc ata Lys Try Cln Ala Glu Arg Cly Gln Gly Glu Val Ala Ser Ala Ile 770  caa tgt tat atg aag gac cat cct aaa att ctc taaa gar aga gcc gar gcd cta Arg ttat atg aag gac cat cct aaa att ctc taaa gar aga gcc Arg ttat at aga gac cat cct caa aar att ctc gaa gaa aga ggt caa gar gar gcd ctc Arg tyr Cln Ala Glu Arg Cly Gln Gly Glu Val Ala Ser Ala Ile 770  caa tgt tat atg aag gac cat cct caa aa att ctc taaa gar gaa gcc Arg ttat atg aag gac cat cct ctc aaa att ctc taaa gar gaa gcc Arg tyr Cln Ala Glu Arg Gly Gln Gly Glu Val Ala Ser Ala Ile 770  c		-	${\tt Tyr}$					Asn			_		Leu	-	-		1871		
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gaa Glu		_	_		_					_			-		-	2543
aca Thr																2591
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Ser	Ser	Thr			Lys	_					_		_	_	Asp	
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Glu	Lys	Arg 115	Ile	Glu	Thr	Leu	Ile 120	Ser	Glu	Ile	Lys	Asn 125	Met	Phe	Arg	
Cys	Met 130	Gly	Tyr	Gly	Glu	Thr 135	Asn	Pro	Ser	Ala	Tyr 140	Asp	Thr	Ala	Trp	
Val 145	Ala	Arg	Ile	Pro	Ala 150	Val	Asp	Gly	Ser	Asp 155	Asn	Pro	His	Phe	Pro 160	
Glu	Thr	Val	Glu	Trp 165	Ile	Leu	Gln	Asn	Gln 170	Leu	Lys	Asp	Gly	Ser 175	Trp	
Gly	Glu	Gly	Phe	Tyr	Phe	Leu	Ala	Tyr 185	Asp	Arg	Ile	Leu	Ala 190	Thr	Leu	
Ala	Сув	Ile 195	Ile	Thr	Leu	Thr	Leu 200	Trp	Arg	Thr	Gly	Glu 205	Thr	Gln	Val	
Gln	L <b>y</b> s 210	Gly	Ile	Glu	Phe	Phe 215	Arg	Thr	Gln	Ala	Gly 220	Lys	Met	Glu	Asp	

Glu 225	Ala	Asp	Ser	His	Arg 230	Pro	Ser	Gly	Phe	Glu 235	Ile	Val	Phe	Pro	Ala 240
Met	Leu	Lys	Glu	Ala 245	Lys	Ile	Leu	Gly	Leu 250	Asp	Leu	Pro	Tyr	<b>A</b> sp 255	Leu
Pro	Phe	Leu	L <b>y</b> s 260	Gln	Ile	Ile	Glu	<b>Lys</b> 265	Arg	Glu	Ala	Lys	Leu 270	Lys	Arg
Ile	Pro	Thr 275	Asp	Val	Leu	Tyr	Ala 280	Leu	Pro	Thr	Thr	Leu 285	Leu	Tyr	Ser
Leu	Glu 290	Gly	Leu	Gln	Glu	Ile 295	Val	Asp	Trp	Gln	Lys 300	Ile	Met	Lys	Leu
Gln 305	Ser	Lys	Asp	Gly	Ser 310	Phe	Leu	Ser	Ser	Pro 315	Ala	Ser	Thr	Ala	Ala 320
Val	Phe	Met	Arg	Thr 325	Gly	Asn	Lys	Lys	Cys 330	Leu	Asp	Phe	Leu	Asn 335	Phe
Val	Leu	Lys	Lys 340	Phe	Gly	Asn	His	Val 345	Pro	Суѕ	His	Tyr	Pro 350	Leu	Asp
Leu	Phe	Glu 355	Arg	Leu	Trp	Ala	Val 360	Asp	Thr	Val	Glu	Arg 365	Leu	Gly	Ile
Asp	Arg 370	His	Phe	Lys	Glu	Glu 375	Ile	Lys	Glu	Ala	Leu 380	Asp	Tyr	Val	Tyr
Ser 385	His	Trp	Asp	Glu	Arg 390	Gly	Ile	Gly	Trp	Ala 395	Arg	Glu	Asn	Pro	Val 400
Pro	Asp	Ile	Asp	Asp 405	Thr	Ala	Met	Gly	Leu 410	Arg	Ile	Leu	Arg	Leu 415	His
Gly	Tyr	Asn	Val 420	Ser	Ser	Asp	Val	Leu 425	Lys	Thr	Phe	Arg	Asp 430	Glu	Asn
Gly	Glu	Phe 435	Phe	Cys	Phe	Leu	Gly 440	Gln	Thr	Gln	Arg	Gly 445	Val	Thr	Asp
Met	Leu 450	Asn	Val	Asn	Arg	С <b>у</b> в 455	Ser	His	Val	Ser	Phe 460	Pro	Gly	Glu	Thr
Ile 465	Met	Glu	Glu	Ala	L <b>y</b> s 470	Leu	Сув	Thr	Glu	Arg 475	Tyr	Leu	Arg	Asn	Ala 480
Leu	Glu	Asn	Val	Asp 485	Ala	Phe	Asp	Lys	Trp 490	Ala	Phe	Lys	Lys	Asn 495	Ile
Arg	Gly	Glu	Val 500	Glu	Tyr	Ala	Leu	L <b>y</b> s 505	Tyr	Pro	Trp	His	<b>Lys</b> 510	Ser	Met
Pro	Arg		Glu		Arg	Ser		Ile						Asp	Asp
Val	Trp 530	Leu	Gly	Lys	Thr	Val 535	Tyr	Met	Met	Pro	<b>Ty</b> r 540	Ile	Ser	Asn	Glu
L <b>y</b> s 545	Tyr	Leu	Glu	Leu	Ala 550	Lys	Leu	Asp	Phe	Asn 555	Lys	Val	Gln	Ser	Ile 560
His	Gln	Thr	Glu	Leu 565	Gln	Asp	Leu	Arg	Arg 570	Trp	Trp	Lys	Ser	Ser 575	Gly
Phe	Thr	Asp	Leu 580	Asn	Phe	Thr	Arg	Glu 585	Arg	Val	Thr	Glu	Ile 590	Tyr	Phe
Ser	Pro	Ala 595	Ser	Phe	Ile	Phe	Glu 600	Pro	Glu	Phe	Ser	L <b>y</b> s 605	Сув	Arg	Glu
Val	<b>Tyr</b> 610	Thr	Lys	Thr	Ser	Asn 615	Phe	Thr	Val	Ile	Leu 620	Asp	Asp	Leu	Tyr

Lys	Arg	Trp	Asp	Leu 645	Ser	Leu	Val	Asp	Gln 650	Met	Pro	Gln	Gln	Met 655	Lys		
Ile	Сув	Phe	Val 660	Gly	Phe	Tyr	Asn	Thr 665	Phe	Asn	Asp	Ile	Ala 670	Lys	Glu		
Gly	Arg	Glu 675	Arg	Gln	Gly	Arg	Asp 680	Val	Leu	Gly	Tyr	Ile 685	Gln	Asn	Val		
Trp	L <b>y</b> s 690	Val	Gln	Leu	Glu	Ala 695	Tyr	Thr	Lys	Glu	Ala 700	Glu	Trp	Ser	Glu		
Ala 705	Lys	Tyr	Val	Pro	Ser 710	Phe	Asn	Glu	Tyr	Ile 715	Glu	Asn	Ala	Ser	Val 720		
Ser	Ile	Ala	Leu	Gl <b>y</b> 725	Thr	Val	Val	Leu	Ile 730	Ser	Ala	Leu	Phe	Thr 735	Gly		
Glu	Val	Leu	Thr 740	Asp	Glu	Val	Leu	Ser 745	Lys	Ile	Asp	Arg	Glu 750	Ser	Arg		
Phe	Leu	Gln 755	Leu	Met	Gly	Leu	Thr 760	Gly	Arg	Leu	Val	Asn 765	Asp	Thr	Lys		
Thr	<b>Ty</b> r 770	Gln	Ala	Glu	Arg	Gly 775	Gln	Gly	Glu	Val	Ala 780	Ser	Ala	Ile	Gln		
C <b>y</b> s 785	Tyr	Met	Lys	Asp	His 790	Pro	Lys	Ile	Ser	Glu 795	Glu	Glu	Ala	Leu	Gln 800		
His	Val	Tyr	Ser	Val 805	Met	Glu	Asn	Ala	Leu 810	Glu	Glu	Leu	Asn	Arg 815	Glu		
Phe	Val	Asn	Asn 820	Lys	Ile	Pro	Asp	Ile 825	Tyr	Lys	Arg	Leu	Val 830	Phe	Glu		
Thr	Ala	Arg 835	Ile	Met	Gln	Leu	Phe 840	Tyr	Met	Gln	Gly	Asp 845	Gly	Leu	Thr		
Leu	Ser 850	His	Asp	Met	Glu	Ile 855	Lys	Glu	His	Val	L <b>y</b> s 860	Asn	Суѕ	Leu	Phe		
Gln 865	Pro	Val	Ala														
<213 <213 <213 <220 <223 <223	)> FE l> NA ?> LO	NGTH PE: RGANI ATUR ME/F	H: 20 DNA [SM: RE: KEY: [ON:	089 Abie	) (	(1983	3)	-limo	onene	e s <b>y</b> r	nthas	se					
<400	)> SE	QUE	ICE:	57													
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aggo	cagga	aat o	Me				eu S					eu G			cc aaa co Lys	111	
				aaa Lys												159	
	_			aca Thr	-	-				_	_	-		-	-	207	
				atc Ile 50												255	
				ggt Gly												303	

_												COII	tin	ueu			
	c ctg n Leu		_	_	_										,,,	351	
	a tct y Ser 95	Ser		_		-	-			-	-		-	-		399	
	g atg 1 Met )															447	
	t gat n Asp															495	
	g ata y Ile	-	_							_	-	-		-		543	
	t tac l Tyr															591	
	t act r Thr 175	Phe														639	
-	a ctg g Leu )								-		_	-				687	
	t gaa o Glu															735	
	c act e Thr															783	
	ggg Gly															831	
	g aaa 1 Lys 255	Lys	_			_		_	_							879	
	a gaa e Glu )		_		-					_		_		_	_	927	
-	a gca ı Ala	Arg	Asn	Tyr		Glu	Val	Tyr	Glu	Gln	-	Gly	Tyr	Glu	-	975	
	a aac 1 Asn		_			-		_	_	_					-	1023	
	a ttg s Leu								_			_				1071	
	t atc Ile 335	Ser														1119	
	a cgg r Arg															1167	
	g ttg t Leu															1215	
	t ctt s Leu	_		_		-	-			_					_	1263	

aac gaa ctc caa ctt ttt acg gat gca att aag aga tgg gat ttg tca Asn Glu Leu Gln Leu Phe Thr Asp Ala Ile Lys Arg Trp Asp Leu Ser 400 405 410	1311
acg aca agg tgg ctt cca gaa tat atg aaa gga gtg tac atg gac ttg Thr Thr Arg Trp Leu Pro Glu Tyr Met Lys Gly Val Tyr Met Asp Leu 415 420 425	1359
tat caa tgc att aat gaa atg gtg gaa gag gct gag aag act caa ggc Tyr Gln Cys Ile Asn Glu Met Val Glu Glu Ala Glu Lys Thr Gln Gly 430 435 440 445	1407
cga gat atg ctc aac tat att caa aat gct tgg gaa gcc cta ttt gat Arg Asp Met Leu Asn Tyr Ile Gln Asn Ala Trp Glu Ala Leu Phe Asp 450 455 460	1455
acc ttt atg caa gaa gca aag tgg atc tcc agc agt tat ctc cca acg Thr Phe Met Gln Glu Ala Lys Trp Ile Ser Ser Ser Tyr Leu Pro Thr $465$ $470$ $475$	1503
ttt gag gag tac ttg aag aat gca aaa gtt agt tct ggt tct cgc ata Phe Glu Glu Tyr Leu Lys Asn Ala Lys Val Ser Ser Gly Ser Arg Ile 480 485 490	1551
gcc aca tta caa ccc att ctc act ttg gat gta cca ctt cct gat tac Ala Thr Leu Gln Pro Ile Leu Thr Leu Asp Val Pro Leu Pro Asp Tyr 495 500 505	1599
ata ctg caa gaa att gat tat cca tcc aga ttc aat gag tta gct tcg Ile Leu Gln Glu Ile Asp Tyr Pro Ser Arg Phe Asn Glu Leu Ala Ser 510 520 525	1647
tcc atc ctt cga cta cga ggt gac acg cgc tgc tac aag gcg gat agg Ser Ile Leu Arg Leu Arg Gly Asp Thr Arg Cys Tyr Lys Ala Asp Arg 530 535 540	1695
gcc cgt gga gaa gaa gct tca gct ata tcg tgt tat atg aaa gac cat Ala Arg Gly Glu Glu Ala Ser Ala Ile Ser Cys Tyr Met Lys Asp His 545 550 555	1743
cct gga tca ata gag gaa gat gct ctc aat cat atc aac gcc atg atc Pro Gly Ser Ile Glu Glu Asp Ala Leu Asn His Ile Asn Ala Met Ile 560 565 570	1791
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Leu Lys Ser Leu Ile Ser Ser Ser Asn Val Gln Lys Ala Leu Cys Ile 20 25 30	
20 23 30	

Ser	Thr	Ala 35	Val	Pro	Thr	Leu	Arg 40	Met	Arg	Arg	Arg	Gln 45	Lys	Ala	Leu
Val	Ile 50	Asn	Met	Lys	Leu	Thr 55	Thr	Val	Ser	His	Arg 60	Asp	Asp	Asn	Gly
Gly 65	Gly	Val	Leu	Gln	Arg 70	Arg	Ile	Ala	Asp	His 75	His	Pro	Asn	Leu	Trp 80
Glu	Asp	Asp	Phe	Ile 85	Gln	Ser	Leu	Ser	Ser 90	Pro	Tyr	Gly	Gly	Ser 95	Ser
Tyr	Ser	Glu	Arg 100	Ala	Glu	Thr	Val	Val 105	Glu	Glu	Val	Lys	Glu 110	Met	Phe
Asn	Ser	Ile 115	Pro	Asn	Asn	Arg	Glu 120	Leu	Phe	Gly	Ser	Gln 125	Asn	Asp	Leu
Leu	Thr 130	Arg	Leu	Trp	Met	Val 135	Asp	Ser	Ile	Glu	Arg 140	Leu	Gly	Ile	Asp
Arg 145	His	Phe	Gln	Asn	Glu 150	Ile	Arg	Val	Ala	Leu 155	Asp	Tyr	Val	Tyr	Ser 160
Tyr	Trp	Lys	Glu	Lys 165	Glu	Gly	Ile	Gly	C <b>y</b> s 170	Gly	Arg	Asp	Ser	Thr 175	Phe
Pro	Asp	Leu	Asn 180	Ser	Thr	Ala	Leu	Ala 185	Leu	Arg	Thr	Leu	Arg 190	Leu	His
Gly	Tyr	Asn 195	Val	Ser	Ser	Asp	Val 200	Leu	Glu	Tyr	Phe	L <b>y</b> s 205	Asp	Glu	Lys
Gly	His 210	Phe	Ala	Суѕ	Pro	Ala 215	Ile	Leu	Thr	Glu	Gly 220	Gln	Ile	Thr	Arg
Ser 225	Val	Leu	Asn	Leu	Tyr 230	Arg	Ala	Ser	Leu	Val 235	Ala	Phe	Pro	Gly	Glu 240
Lys	Val	Met	Glu	Glu 245	Ala	Glu	Ile	Phe	Ser 250	Ala	Ser	Tyr	Leu	<b>Lys</b> 255	Lys
Val	Leu	Gln	L <b>y</b> s 260	Ile	Pro	Val	Ser	Asn 265	Leu	Ser	Gly	Glu	Ile 270	Glu	Tyr
Val	Leu	Glu 275	Tyr	Gly	Trp	His	Thr 280	Asn	Leu	Pro	Arg	Leu 285	Glu	Ala	Arg
Asn	<b>Ty</b> r 290	Ile	Glu	Val	Tyr	Glu 295	Gln	Ser	Gly	Tyr	Glu 300	Ser	Leu	Asn	Glu
Met 305	Pro	Tyr	Met	Asn	Met 310	Lys	Lys	Leu	Leu	Gln 315	Leu	Ala	Lys	Leu	Glu 320
Phe	Asn	Ile		His 325	Ser	Leu		Leu	-		Leu	Gln		Ile 335	
Arg	Trp	Trp	Lys 340	Glu	Ser	Gly	Ser	Ser 345	Gln	Leu	Thr	Phe	Thr 350	Arg	His
Arg	His	Val 355	Glu	Tyr	Tyr	Thr	Met 360	Ala	Ser	Cys	Ile	Ser 365	Met	Leu	Pro
Lys	His 370	Ser	Ala	Phe	Arg	Met 375	Glu	Phe	Val	Lys	Val 380	Cys	His	Leu	Val
Thr 385	Val	Leu	Asp	Asp	Ile 390	Tyr	Asp	Thr	Phe	Gly 395	Thr	Met	Asn	Glu	Leu 400
Gln	Leu	Phe	Thr	Asp 405	Ala	Ile	Lys	Arg	Trp 410	Asp	Leu	Ser	Thr	Thr 415	Arg
Trp	Leu	Pro	Glu 420	Tyr	Met	Lys	Gly	Val 425	Tyr	Met	Asp	Leu	Tyr 430	Gln	Сув
Ile	Asn	Glu 435	Met	Val	Glu	Glu	Ala 440	Glu	Lys	Thr	Gln	Gly 445	Arg	Asp	Met
Leu	Asn	Tyr	Ile	Gln	Asn	Ala	Trp	Glu	Ala	Leu	Phe	Asp	Thr	Phe	Met

	450					455					460				
Gln 465	Glu	Ala	Lys	Trp	Ile 470	Ser	Ser	Ser	Tyr	Leu 475	Pro	Thr	Phe	Glu	Glu 480
Tyr	Leu	Lys	Asn	Ala 485	Lys	Val	Ser	Ser	Gl <b>y</b> 490	Ser	Arg	Ile	Ala	Thr 495	Leu
Gln	Pro	Ile	Leu 500	Thr	Leu	Asp	Val	Pro 505	Leu	Pro	Asp	Tyr	Ile 510	Leu	Gln
Glu	Ile	Asp 515	Tyr	Pro	Ser	Arg	Phe 520	Asn	Glu	Leu	Ala	Ser 525	Ser	Ile	Leu
Arg	Leu 530	Arg	Gly	Asp	Thr	Arg 535	Сув	Tyr	Lys	Ala	Asp 540	Arg	Ala	Arg	Gly
Glu 545	Glu	Ala	Ser	Ala	Ile 550	Ser	Cys	Tyr	Met	L <b>y</b> s 555	Asp	His	Pro	Gly	Ser 560
Ile	Glu	Glu	Asp	Ala 565	Leu	Asn	His	Ile	Asn 570	Ala	Met	Ile	Ser	<b>A</b> sp 575	Ala
Ile	Arg	Glu	Leu 580	Asn	Trp	Glu	Leu	Leu 585	Arg	Pro	Asp	Ser	L <b>y</b> s 590	Ser	Pro
Ile	Ser	Ser 595	Lys	Lys	His	Ala	Phe 600	Asp	Ile	Thr	Arg	Ala 605	Phe	His	His
Val	Tyr 610	Lys	Tyr	Arg	Asp	Gl <b>y</b> 615	Tyr	Thr	Val	Ser	Asn 620	Asn	Glu	Thr	Lys
Asn 625	Leu	Val	Met	Lys	Thr 630	Val	Leu	Glu	Pro	Leu 635	Ala	Leu			

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#### We claim the following:

1. An isolated nucleic acid encoding a terpene synthase having a region with 20% or greater sequence identity to residues 265 to 535 of SEQ ID NO: 2, said synthase comprising nine  $\alpha$ -carbons having interatomic distances in Angstroms between said a-carbons that are  $\pm 2.3$  Angstroms of the following interatomic distances:

α- Carbon	1	2	3	4	5	6	7	8	9
1	0.0	8.4	13.7	12.7	11.9	10.2	13.1	9.4	12.8
2	8.4	0.0	11.3	8.7	10.2	7.2	14.8	15.1	17.4
3	13.7	11.3	0.0	3.8	5.4	9.3	6.6	13.9	13.7
4	12.7	8.7	3.8	0.0	3.8	6.0	9.2	15.4	16.1
5	11.9	10.2	5.4	3.8	0.0	5.0	7.8	14.6	15.5
6	10.2	7.2	9.3	6.0	5.0	0.0	12.0	16.1	18.0
7	13.1	14.8	6.6	9.2	7.8	12.0	0.0	10.2	9.5
8	9.4	15.1	13.9	15.4	14.6	16.1	10.2	0.0	3.8
9	12.8	17.4	13.7	16.1	15.5	18.0	9.5	3.8	0.0

the center point of each said  $\alpha$ -carbon positioned within a sphere having a radius of 2.3 Angstroms, the center points of each said sphere having the following structural coordinates:

				_
α-Carbon Number	X Position	Y Position	Z Position	60
1	120.203	38.695	43.506	
2	114.058	43.884	41.015	
3	106.807	36.336	45.151	
4	107.629	38.010	41.804	
5	109.375	34.842	40.617	65
6	111.944	37.854	37.602	

-continued

α-Carbon Number	X Position	Y Position	Z Position
7	110.233	31.098	47.361
8	118.846	34.443	51.796
9	116.461	32.848	54.290,

each said  $\alpha$ -carbon having an associated R-group, said synthase having an ordered arrangement of said R-groups other than the following ordered arrangements of R-groups:

TABLE 9

]	Α '	1 W	2	3	4	_				
1	_	W				5	6	7	8	9
	В		I	Т	Т	Y	L	С	Т	Y
		W	I	S	T	Y	L	C	T	Y
	C 1	W	I	C	G	Y	L	C	L	Y
]	, C	W	I	S	G	Y	L	C	L	Y
]	E '	W	L	Α	G	Y	I	Α	L	Y
55	F '	W	L	T	V	H	L	G	V	Y
~ (	G '	W	L	Α	G	Y	I	Α	L	Y
]	H	W	I	V	G	N	L	F	L	Y
	I '	W	I	T	Α	G	L	S	C	Y
	J '		V	S	C	I	M	G	S	Y
1	K	F	F	I	T	Α	T	G	T	Y
60	L '	W	N	I	S	G	M	L	M	Y
00 I	M '	W	V	S	S	Y	L	G	L	Y
1	N	F	F	T	L	A	L	G	S	Y
(	, c	W	N	S	G	P	L	L	M	Y
	P '	W	N	G	G	I	L	L	I	Y
(	Q .	Y	L	V	T	M	T	G	T	Y
		W	I	I	S	A	I	L	I	Y
65	s '	W	F	S	S	V	I	L	I	Y
,	Г	W	I	V	Α	S	I	L	I	Y

TABLE 9-continued

		O	rdered A	Arrange	ments c	of α-Ca	rbons 1	<b>_</b> 9	
	1	2	3	4	5	6	7	8	9
U	W	N	I	s	S	I	F	M	Y
V	L	Α	I	G	Q	L	S	I	F
$\mathbf{w}$	S	S	I	Α	L	V	G	F	Y
X	L	С	С	G	Η	S	L	G	Y
Y	S	F	S	S	V	I	L	V	Y
Z	$\mathbf{W}$	Α	S	G	M	L	G	I	Y
AA	Α	N	L	T	S	T	C	L	Y
$^{\mathrm{BB}}$	L	C	S	Α	$\mathbf{Y}$	V	L	L	Y
CC	W	Α	T	G	M	L	S	M	Y
DD	M	C	S	S	G	I	L	V	Y
EE	S	G	V	G	L	C	W	F	Y
FF	S	G	Α	L	G	V	G	F	Y
GG	S	G	F	Α	L	I	G	F	Y
$_{ m HH}$	Α	G	F	Α	L	I	G	F	Y
II	W	V	T	G	L	V	I	S	Y
JJ	W	Α	S	G	M	L	G	I	Y
KK	W	I	S	T	$\mathbf{Y}$	L	С	T	Y
LL	W	I	T	T	Y	L	С	T	Y
MM	W	N	I	S	G	M	L	M	Y
NN	Α	Α	I	G	Q	L	S	I	F
OO	Α	I	V	Α	S	I	L	I	Y.

- 2. The nucleic acid of claim 1, wherein said synthase has  $^{25}$  25% or greater sequence identity to residues 265 to 535 of SEQ ID NO:2.
- 3. The nucleic acid of claim 1, wherein said synthase has 35% or greater sequence identity to residues 265 to 535 of SEQ ID NO:2.
- 4. The nucleic acid of claim 1, wherein said synthase catalyses the formation of a terpenoid product from a monoterpene substrate.
- 5. The nucleic acid of claim 1, wherein said synthase catalyses the formation of a terpenoid product from a  $_{35}$  sesquiterpene substrate.
- **6.** The nucleic acid of claim **1,** wherein said synthase catalyses the formation of a terpenoid product from a diterpene substrate.
- 7. The nucleic acid of claim 4, herein said product is a cyclic terpenoid hydrocarbon.

- 8. The nucleic acid of claim 5, wherein said product is a cyclic terpenoid hydrocarbon.
- **9**. The nucleic acid of claim **6**, wherein said product is a cyclic terpenoid hydrocarbon.
- 10. A cultured host cell containing the nucleic acid of claim 1.
- 11. The host cell of claim 10, wherein said cell is a prokaryotic cell.
- 12. The host cell of claim 10, wherein said cell is a eukaryotic cell.
- 13. The host cell of claim 12, wherein said cell is an insect cell.
- 14. The host cell of claim 12, wherein said cell is a plant cell.
- 15. The host cell of claim 14, wherein said host cell is an Angiosperm cell.
- 16. The host cell of claim 14, wherein said host cell is an Gymnosperm cell.
- 17. The host cell of claim 14 wherein said host cell is selected from the group consisting of: a cell from a Graminaceae plant, a cell from a Legumineae plant, a cell from a Solanaceae plant, a cell from a Brassicaeae plant and a cell from a Conifereae plant.
- 18. A transgenic animal cell culture, wherein cells in said cell culture comprise the nucleic acid of claim 1.
- 19. The nucleic acid of claim 1, wherein said synthase has 40% or greater sequence identity to residues 265 to 535 of SEQ ID NO:2.
- **20**. The nucleic acid of claim 1, wherein said synthase has 50% or greater sequence identity to residues 265 to 535 of SEQ ID NO:2.
- 21. The nucleic acid of claim 1, wherein said synthase has 60% or greater sequence identity to residues 265 to 535 of SEQ ID NO:2.
- 22. The nucleic acid of claim 1, wherein said synthase has 70% or greater sequence identity to residues 265 to 535 of SEQ ID NO:2.
- 23. The nucleic acid of claim 1, wherein said synthase has 80% or greater sequence identity to residues 265 to 535 of SEQ ID NO:2.

\* \* \* \* \*