

1964

# The Y-glutamyl bond in chick embryo collagen

William Brewster Pratt  
*Yale University*

Follow this and additional works at: <http://elischolar.library.yale.edu/ymtdl>

---

## Recommended Citation

Pratt, William Brewster, "The Y-glutamyl bond in chick embryo collagen" (1964). *Yale Medicine Thesis Digital Library*. 3042.  
<http://elischolar.library.yale.edu/ymtdl/3042>

This Open Access Thesis is brought to you for free and open access by the School of Medicine at EliScholar – A Digital Platform for Scholarly Publishing at Yale. It has been accepted for inclusion in Yale Medicine Thesis Digital Library by an authorized administrator of EliScholar – A Digital Platform for Scholarly Publishing at Yale. For more information, please contact [elischolar@yale.edu](mailto:elischolar@yale.edu).

T113  
+Y12  
2617

YALE UNIVERSITY LIBRARY



3 9002 06679 0867

THE GAMMA-GLUTAMYL BOND IN  
CHICK EMBRYO COLLAGEN



WILLIAM PRATT


1964

MUDD  
LIBRARY  
Medical

YALE



MEDICAL LIBRARY



Digitized by the Internet Archive  
in 2017 with funding from  
The National Endowment for the Humanities and the Arcadia Fund

<https://archive.org/details/yglutamylbondinc00prat>







The  $\gamma$ -Glutamyl Bond in Chick Embryo Collagen

By

William Brewster Pratt

B.A., Dartmouth College, 1960

A thesis Presented to the Faculty and Officers of  
the Yale University School of Medicine in partial  
fulfillment of the requirements for the degree of  
Doctor of Medicine

April, 1964

Department of Biochemistry  
Yale University School of Medicine



YALE MEDICAL LIBRARY  
JUL 1964  
LIPDIN

T113

Y12

2617

### Acknowledgement

The author would like to thank Dr. Lewis Lukens for his patient guidance and encouragement in both the experimentation and in the writing of the thesis.



Table of Contents.

Introduction.....	1
Materials.....	16
Methods.....	19
Results.....	32
Discussion.....	50
Summary.....	56
Bibliography.....	58



## Introduction

Quastel, Stewart, and Tunicliffe in 1923 (1) were the first to present evidence for the existence of gamma-glutamyl peptide linkages in a naturally occurring substance, glutathione, which had been isolated from yeast and demonstrated in various animal tissues by Hopkins in 1921 (2). Using the observation of Dakin (3) that alpha-amino acids, oxidized with hydrogen peroxide in the presence of a trace of iron salt, yield carboxylic acids with one carbon less than the original compound, Quastel et al. were able to oxidize impure preparations of glutathione with hydrogen peroxide and after acid hydrolysis to identify succinic acid among the products. This evidence suggested that the glutamic acid in glutathione was joined to the tripeptide through its gamma-carboxyl group leaving its alpha amino carboxylic grouping open to attack by the oxidizing agent. Kendall et al. (4) were able to demonstrate that oxidation of crystalline glutathione with hypobromite or Chloramine T did not disrupt the peptide bonds but did form products that yielded succinic acid on hydrolysis. The final proof for the presence of gamma-linkage in glutathione was offered by Harrington and Mead (5) who synthesized gamma-glutamylcysteinylglycine and showed that its



properties were identical with those of the crystalline peptide.

In 1937 Ivanovics and Bruckner (6) isolated and purified the capsular substance of Bacillus anthracis and Bacillus subtilis. On hydrolysis of the material they found only glutamic acid, but they were not able to say that the protein was made up solely of this amino acid. They pointed out that, if the capsular substance contained only glutamic acid, then two isomeric forms are possible depending on whether the alpha-carboxyl or the gamma-carboxyl or both are involved in the peptide linkages (Fig. 1).

Bovarnick (7) was able by comparison of total nitrogen to glutamic acid nitrogen to show that the capsular substance of Bacillus subtilis consisted solely of glutamic acid units. Working on the assumption that racemization of the glutamic acid units in alkaline solution would not occur if a free carboxyl was situated adjacent to the asymmetric carbon atom (gamma-glutamyl form), Bovarnick allowed purified capsular substance to stand for ten days in alkaline solution. There was no change in optical rotation of the solution over this period - amino nitrogen showed only a small increase - from 10 to 20% - demonstrating that significant hydrolysis which would prevent racemization.



... ..

... ..

... ..

... ..

... ..

... ..

... ..

... ..

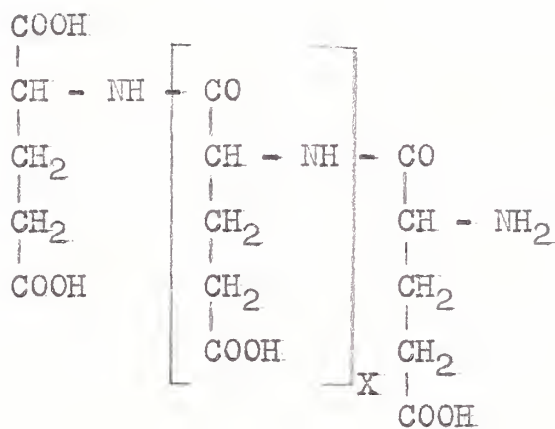
... ..

... ..

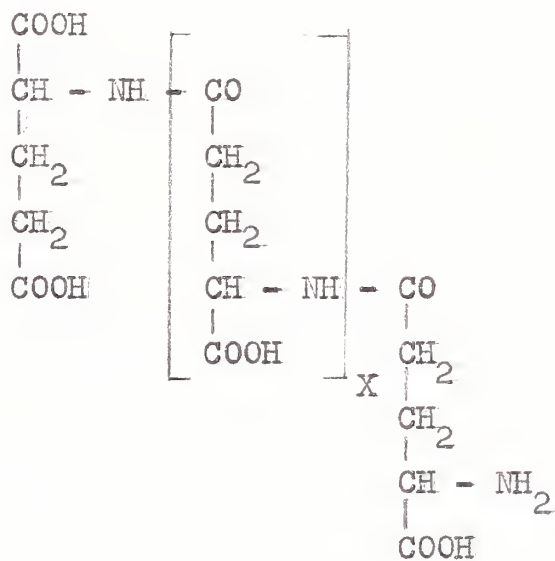
... ..

... ..

(3)



Alpha-linkage



Gamma-linkage

Fig. 1. Structure of the alpha- and gamma-peptide linkages.



had not occurred. Gamma-peptide linkage of the glutamic acid was indicated on this basis. In support of this conclusion, a biuret test, which is specific for alpha-linked units, was negative.

Further evidence in support of a homogenous composition for the capsular substance was provided when Hanby and Rydon (8) converted the glutamic acid obtained from hydrolysis of Bacillus anthracis hapten to pyrrolidone-carboxylic acid and isolated this product quantitatively by chromatography on silica gel, thus verifying that the capsular substance was composed solely of glutamic acid units. The ready fission of the capsular substance by acid indicated to these investigators that the glutamic acid residues were linked, in part at least, by linkages which were more susceptible to acid hydrolysis than alpha-peptide bonds. They noticed that during Van Slyke amino-nitrogen assays the apparent amino nitrogen content of the capsular substance increased (the apparent molecular weight decreased) with increasing reaction time, a finding which indicated to the authors that in addition to the free amino group, the capsular substance also contained gamma-peptide groupings which liberated their nitrogen more slowly under Van Slyke conditions. One sample submitted to this procedure (molecular

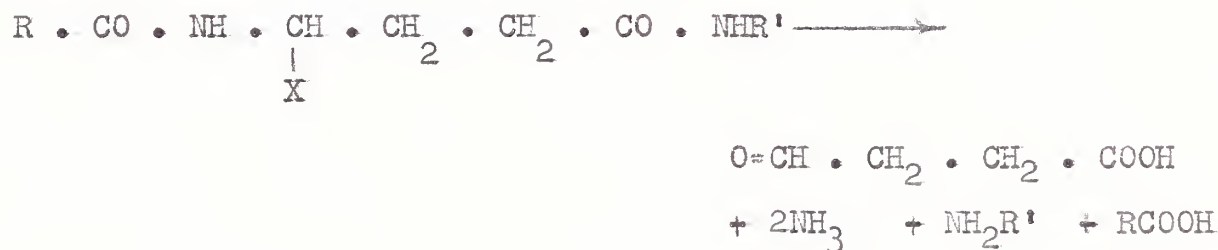


(5)



Alpha-glutamyl peptide

Alpha,gamma-diaminobutyric acid



Gamma-glutamyl peptide

Succinic semialdehyde

Fig. 2. Conversion of the alpha- and gamma-glutamyl peptides to the polyhydrazide followed by Curtius degradation. Unaltered polypeptide, X=COOH; polymethyl ester, X=COOCH<sub>3</sub>; polyhydrazide, X=CO · NH · NH<sub>2</sub>; product of Curtius degradation, X=NH<sub>3</sub>



weight 5000) showed only a slight increase in amino-nitrogen with reaction time and they felt that this represented a final degradation product composed solely of alpha-peptide linkage, the gamma-linkages having been hydrolyzed by acid during the preparation.

Kovacs and Bruckner (9) (Fig. 2.) converted the poly-D-glutamic acid of Bacillus subtilis into the polymethyl ester and then into the polyhydrazide. After Curtius degradation followed by acid hydrolysis only succinic semialdehyde (the gamma-linkage rearrangement product) was found in the hydrolysate - there was no alpha,gamma-diaminobutyric acid. This indicated that in native poly-D-glutamic acid gamma-glutamyl bonds predominated. Later, Kovacs, Bruckner, and Kovacs (10) subjected alpha-L-polyglutamic acid hydrazide to the Curtius procedure and on hydrolysis recovered only alpha,gamma-diaminobutyric acid. Bruckner, Kovacs, and Nagy (11) then converted methyl poly-D-glutamate (prepared from the capsule of Bacillus subtilis) into the polyamide which was submitted to Hofmann degradation. After hydrolysis of the rearranged product only succinic semialdehyde was recovered.

Waley (12) synthesized Poly-(gamma-L-glutamyl)-L-glutamic acid. Many of its properties (eg. solubility in water; ionization constant, infra-red spectra, and reactivity to ninhydrin) were similar to those of Bacillus subtilis and



The first part of the report deals with the general situation of the country and the progress of the work done during the year. It is followed by a detailed account of the various projects and schemes undertaken, and a summary of the results achieved. The report concludes with a statement of the financial position and a list of the members of the committee.

The committee has the honor to acknowledge the assistance rendered by the various departments of the Government, and the co-operation of the public in the execution of the work. It is a pleasure to state that the progress made during the year has been satisfactory, and that the various projects have been carried out in accordance with the programme laid down in the budget.

The committee has also the honor to state that the financial position of the country is sound, and that the revenue has been collected in accordance with the estimates. It is a pleasure to state that the various departments of the Government have worked in a harmonious and efficient manner, and that the public has co-operated in the execution of the work.

The committee has the honor to state that the progress made during the year has been satisfactory, and that the various projects have been carried out in accordance with the programme laid down in the budget. It is a pleasure to state that the financial position of the country is sound, and that the revenue has been collected in accordance with the estimates.

The committee has the honor to state that the progress made during the year has been satisfactory, and that the various projects have been carried out in accordance with the programme laid down in the budget. It is a pleasure to state that the financial position of the country is sound, and that the revenue has been collected in accordance with the estimates.

The committee has the honor to state that the progress made during the year has been satisfactory, and that the various projects have been carried out in accordance with the programme laid down in the budget. It is a pleasure to state that the financial position of the country is sound, and that the revenue has been collected in accordance with the estimates.

The committee has the honor to state that the progress made during the year has been satisfactory, and that the various projects have been carried out in accordance with the programme laid down in the budget. It is a pleasure to state that the financial position of the country is sound, and that the revenue has been collected in accordance with the estimates.

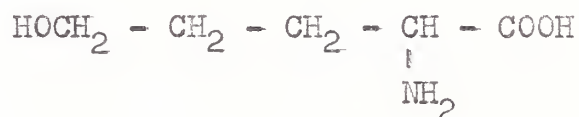
The committee has the honor to state that the progress made during the year has been satisfactory, and that the various projects have been carried out in accordance with the programme laid down in the budget. It is a pleasure to state that the financial position of the country is sound, and that the revenue has been collected in accordance with the estimates.

The committee has the honor to state that the progress made during the year has been satisfactory, and that the various projects have been carried out in accordance with the programme laid down in the budget. It is a pleasure to state that the financial position of the country is sound, and that the revenue has been collected in accordance with the estimates.

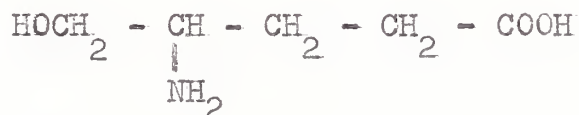
The committee has the honor to state that the progress made during the year has been satisfactory, and that the various projects have been carried out in accordance with the programme laid down in the budget. It is a pleasure to state that the financial position of the country is sound, and that the revenue has been collected in accordance with the estimates.

Bacillus licheniformis and differed from those of a synthetic alpha-linked polypeptide.

Chibnall, Rees, and Richards (13) esterified several samples of poly-glutamic acid from Bacillus subtilis, reduced them with lithium borohydride and analyzed the products given on subsequent acid hydrolysis. Under these conditions an alpha-linked glutamyl unit would be expected to yield delta-hydroxy-alpha-aminovaleric acid:



and a gamma-linked glutamyl unit would yield delta-hydroxy-gamma-aminovaleric acid:



Only the latter product was found, thus affirming the solely gamma-linked nature of the capsular substance. A synthetic poly-alpha-glutamic acid methyl ester treated in the same manner yielded 77.8% of total N as delta-hydroxy-alpha-aminovaleric acid and none as the gamma-linked reduction product.

Since the work on capsular substance was carried out, gamma-glutamyl linkages have been found in several natural

The first part of the document discusses the importance of maintaining accurate records of all transactions. It emphasizes that every entry should be supported by a valid receipt or invoice. This ensures transparency and allows for easy verification of the data.

Furthermore, it is noted that the records should be kept in a secure and accessible format. Regular backups are recommended to prevent data loss in the event of a system failure or disaster.

--- -- --

In addition, the document outlines the procedures for handling discrepancies. If there is a mismatch between the recorded amounts and the actual cash flow, it is crucial to investigate the cause immediately. This may involve reviewing the source documents and contacting the relevant parties for clarification.

--- -- --

The final section of the document provides a summary of the key points discussed. It reiterates the importance of consistency and accuracy in all financial reporting. By following these guidelines, the organization can ensure that its financial data is reliable and trustworthy.

It is also noted that these procedures should be reviewed periodically to ensure they remain up-to-date with current regulations and best practices.

The document concludes with a statement of intent to implement these procedures effectively. It expresses confidence that these measures will lead to improved financial management and reporting.

substances, a most interesting example being the presence of these bonds in collagen as ascertained by Gallop, Seifter et al (14).

These investigators noted that the conditions necessary to bring about Hofmann and Curtius rearrangements (references 9-11) when applied to proteins on a small scale, require drastic conditions and are often non-quantitative. In view of this, they studied the application of the Lossen rearrangement of the dinitrophenyl derivatives of hydroxamic acids to analysis of carboxyl groups in several compounds including a commercial base processed gelatin obtained from pig skin. Gallop et al. felt that the relative mildness of the conditions employed endowed the reactions with greater specificity. They esterified gelatin with methanol and acetic anhydride, formed the hydroxamic acid derivative by adding the methyl ester to neutral aqueous hydroxylamine, reacted the hydroxamate with FDNB<sup>1</sup> at pH 7.0 and promoted Lossen rearrangement of the hydroxamate-DNP derivative by heating at 100° C. for 2 min. in 0.1 N NaOH. After acid hydrolysis the rearrangement products, alpha,gamma-diaminobutyric acid in the case of alpha-linkage

1. The abbreviation used is: FDNB, fluorodinitrobenzene.

1. The first part of the document discusses the importance of maintaining accurate records of all transactions and activities. It emphasizes that this is essential for ensuring transparency and accountability in the organization's operations.

2. The second part of the document outlines the various methods and tools used to collect and analyze data. It highlights the need for consistent data collection procedures and the use of advanced analytical techniques to derive meaningful insights from the data.

3. The third part of the document focuses on the role of technology in data management and analysis. It discusses how modern software solutions can streamline data collection, storage, and processing, thereby improving efficiency and reducing the risk of errors.

4. The fourth part of the document addresses the challenges associated with data security and privacy. It stresses the importance of implementing robust security measures to protect sensitive information and ensure compliance with relevant regulations.

5. The fifth part of the document provides a summary of the key findings and recommendations. It concludes that a comprehensive data management strategy is crucial for the organization's success and suggests several actionable steps to improve data practices.

and succinic semialdehyde from gamma-linkage, were isolated (Fig. 3.). The presence of succinic semialdehyde was ascertained by forming the 2,4-dinitrophenylhydrazone and chromatographing against an authentic sample. Elution of the phenylhydrazone from the paper resulted in variable recoveries thereby limiting the procedure to a qualitative evaluation.

The authors note that the conditions of the procedure may promote interconversion of the alpha- and gamma-glutamyl peptide bonds. The esterification procedure was carried out in the presence of a dehydrating agent which could promote imide formation in the gelatin. Further, the gamma-glutamyl esters in the protein when undergoing attack by hydroxylamine under mild alkaline conditions might be converted to imide intermediates. In either case the imide intermediates could be cleaved with the formation of gamma instead of alpha peptide bonds (Fig. 4.). Thus the appearance of succinic semialdehyde in the hydrolysate of the rearranged gelatin preparation provided strong but not absolute evidence of gamma-linkage.

From their analysis of the hydrolysates of the pre- and post-rearranged gelatin, the authors observed that there was no serious discrepancy between the number of residues









(11)

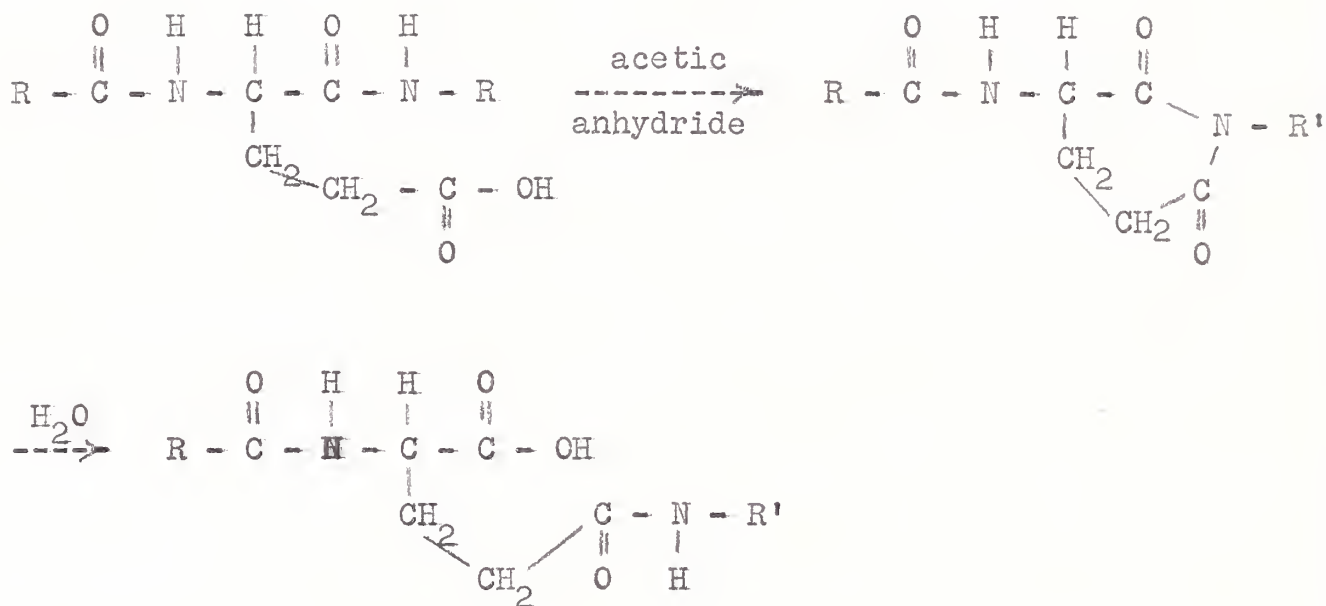


Fig. 4. Method of possible interconversion of alpha- and gamma-glutamyl peptide bonds in the presence of a dehydrating agent.

The first part of the document discusses the importance of maintaining accurate records of all transactions. It emphasizes that every entry should be supported by a valid receipt or invoice. This ensures transparency and allows for easy verification of the data.

In addition, it is crucial to review the records regularly to identify any discrepancies or errors. This proactive approach helps in resolving issues before they become significant problems. The document also mentions the use of digital tools to streamline the record-keeping process and reduce the risk of human error.

Finally, the document concludes by stating that consistent and accurate record-keeping is essential for the long-term success and stability of any organization. It serves as a foundation for informed decision-making and financial planning.

of aspartic acid which disappeared during Lossen rearrangement (13 residues per 1000) and the number of molecules of diaminopropionic acid which appeared (8.8 residues per 1000). However, the alpha,gamma-diaminobutyric acid recovered accounted for only 21% of the glutamic residues lost during rearrangement. They reasoned that, in the absence of theoretical contraindications, if interconversion of alpha- and gamma-bonds took place, then it should occur to an equal extent with both alpha-glutamyl and alpha-aspartyl linkages. On the basis of this reasoning they felt that some gamma-glutamyl peptide links existed in the original collagen.

Franzblau (15), in order to avoid conditions which could promote intermediate imide formation and possible alpha to gamma interchange and also to prevent hydroxylaminolysis of the intramolecular ester cross linkages (Gallop, Seifter, and Meilman (16)), performed a direct conversion by hydroxylamine hydrochloride of free carboxyl groups to hydroxamic acids by carrying out the reaction in aqueous, mildly acidic medium containing 1-cyclohexyl-3-(2-morpholinyl-(4)-ethyl)-carbodiimide metho-p-toluene sulfonate, a water soluble carbodiimide described by Sheehan and Hlavka (17). Franzblau treated the hydroxamate derivative in the same manner



Table I\*

Products identified after Lossen rearrangement of the di-nitrophenyl hydroxamate derivative of gelatin derived from ichthocol as compared with a suitable control

(Expressed as residues per 1000 residues)

<u>Compound</u>	<u>Control gelatin</u>	<u>DNP-Hydroxamate Gelatin after Rearrangement</u>	<u>Difference</u>
Glutamic acid	71.4	46.0	-25.4
alpha,gamma-Diaminobutyric acid	0	4.3	+4.3
Succinic semialdehyde	0	22.0	+22.0
Ammonia	42.0	78.0	+36.0
Aspartic acid	45.8	39.6	-6.2
alpha,beta-Di-aminopropionic acid	0	5.0	+5.0
Dinitrophenol**	0	31.0	+31.0

\* Franzblau, op. cit., p. 76.

\*\* Each mole of dinitrophenylhydroxamate which undergoes rearrangement yields one mole of dinitrophenol.



as Gallop et al. except that the dinitrophenyl derivative of the hydroxamate was formed at pH 8.0 instead of pH 7.0. In this work the succinic semialdehyde was quantitated along with the other rearrangement products and with the amino acids obtained after acid hydrolysis of the rearranged gelatin (Table I). It is evident from Table I that per 1000 residues, 25 residues of glutamic acid were lost with rearrangement and 4.3 residues of alpha,gamma-diaminobutyric acid were recovered. If we are to assume that the rest of the unrecovered glutamic acid, (about 20 residues), was present as gamma-linked glutamic acid, then 40 residues of ammonia and 20 residues of succinic semialdehyde should be found per 1000 residues. In fact 36 molecules of ammonia beyond those present in the control gelatin and 22 molecules of succinic semialdehyde were recovered. Therefore Franzblau concluded that at least 20 residues of glutamic acid per 1000 residues of total amino acids in collagen are in gamma-glutamyl linkage. The failure of roughly half of the glutamate residues to be converted to hydroxamates does not affect Franzblau's quantitative statement; because he demonstrated in synthetic polypeptide experiments that the unconverted glutamic acid residues were in alpha-linkage. This conclusion was further supported by the fact that this





investigator had demonstrated the absence of alpha,gamma-carboxyl interchange when either alpha- or gamma-polyglutamic acid was subjected to an identical procedure. If a similar interpretation is applied to the aspartic acid residues then these must necessarily be alpha-linked.

In addition to the above results, obtained with gelatin prepared from ichthyocol, Franzblau also carried out the same procedure with gelatin derived from calf-skin collagen. In this case, however, no quantitative estimation of the amount of succinic semialdehyde was carried out, although its presence was verified by chromatography.

In conjunction with some investigations being conducted in this laboratory in which a chick embryo system is being used to study the mechanism of formation of hydroxyproline from proline in the process of collagen synthesis, it was felt that it would be interesting to employ the same system in an investigation of the gamma-glutamyl bond. This dissertation will be concerned with the identification and measurement of gamma-linked glutamic acid in chick embryo collagen and with the establishment of an in vivo preparation in which its formation may be studied.



Materials

Bovine Achilles tendon collagen was purchased from the Worthington Biochemical Corp., Freehold, New Jersey. It had been purified by the method of Einbinder and Schubert (18).

Medium A was prepared according to Littlefield and Keller (19) and contained 0.25 M sucrose, 0.025 M KCl, 0.005 M MgCl<sub>2</sub>, and 0.05 M Tris buffer, pH 7.6.

Sephadex-G-25 was obtained from Pharmacia, Uppsala, Sweden.

Dowex AG 50W-X8, 200-400 mesh, Hydrogen form, was purchased from the California Corporation for Biochemical Research, Los Angeles, California.

Silica gel G (according to Stahl) was purchased from Brinkmann Instruments Inc., Great Neck, New York.

1-cyclohexyl-3-(2-morpholinyl-(4)-ethyl)-carbodiimide metho-p-toluene sulfonate was purchased from the Aldrich Chemical Co., Milwaukee, Wisconsin. This compound will be referred to as WSC throughout the remainder of this paper. This is a water-soluble carbodiimide which Franzblau (15) found could promote, in an aqueous, slightly acidic medium, the direct conversion by hydroxylamine hydrochloride of free carboxyl to hydroxamic acid groups. When applied to the gelatin from ichthyocol, Franzblau found that this method resulted in the conversion of 40 to 50% of the

CHAPTER

THE HISTORY OF THE UNITED STATES OF AMERICA

FROM 1789 TO 1861

BY

JOHN B. HENNINGSEN

NEW YORK: THE CENTURY CO., 1908

Copyright, 1908, by The Century Company

Printed in the United States of America

Published by The Century Company, 300 N. 4th St., New York, N. Y.

Entered as Second-Class Matter, October 3, 1879, under No. 233, Postoffice at New York, N. Y., and Mailed at Special Rate of Postage provided for in Postoffice Department Publication Circular No. 263, approved July 16, 1879, authorized on July 16, 1879, and on July 16, 1896, and on July 16, 1906, and on July 16, 1912, and on July 16, 1918, and on July 16, 1924, and on July 16, 1930, and on July 16, 1936, and on July 16, 1942, and on July 16, 1948, and on July 16, 1954, and on July 16, 1960, and on July 16, 1966, and on July 16, 1972, and on July 16, 1978, and on July 16, 1984, and on July 16, 1990, and on July 16, 1996, and on July 16, 2002, and on July 16, 2008, and on July 16, 2014, and on July 16, 2020.

Acceptance for mailing at special rate of postage provided for in Postoffice Department Publication Circular No. 263, approved July 16, 1879, authorized on July 16, 1879, and on July 16, 1896, and on July 16, 1906, and on July 16, 1912, and on July 16, 1918, and on July 16, 1924, and on July 16, 1930, and on July 16, 1936, and on July 16, 1942, and on July 16, 1948, and on July 16, 1954, and on July 16, 1960, and on July 16, 1966, and on July 16, 1972, and on July 16, 1978, and on July 16, 1984, and on July 16, 1990, and on July 16, 1996, and on July 16, 2002, and on July 16, 2008, and on July 16, 2014, and on July 16, 2020.

free carboxyl groups to hydroxamic acids in 2 hours. A separate experiment conducted for 19 hours resulted in approximately 65% conversion to hydroxamic acid groups.

Acethydroxamic acid was prepared by a modification of the synthesis of benzohydroxamic acid according to Blatt (20). Separate solutions of 46.7 g. of hydroxylamine hydrochloride in 240 ml of methanol and 56.1 g. of KOH in 140 ml. of methanol were prepared at the boiling point of the solvent. Both solutions were cooled to 30-40° C., and the alkali solution was added to the hydroxylamine solution in an ice bath. This mixture was allowed to stand in an ice bath for 5 min., and the KCl which formed was filtered off on a Buchner funnel. 41 g. of ethyl acetate were added with stirring and the mixture was allowed to stand at room temperature for 6 hours. This solution was evaporated and an oil appeared which was neutralized to pH 7.0 with HCl. The neutralized oil was dissolved in hot ethyl acetate. The hot solution was allowed to cool slowly to 4° C. The acethydroxamic acid readily crystallized out on cooling. It was then filtered and stored in a dessicator in vacuo. The purity of the compound was verified by a melting point determination.

Succinic semialdehyde-2,4-dinitrophenylhydrazone was

The first part of the document discusses the importance of maintaining accurate records of all transactions. It emphasizes that every entry should be supported by a valid receipt or invoice. This not only helps in tracking expenses but also serves as a legal safeguard in case of an audit.

Furthermore, the document highlights the need for regular reconciliation of bank statements with the company's ledger. This process ensures that there are no discrepancies between the two records, which is crucial for maintaining the integrity of the financial data.

In addition, it is advised to keep all financial documents in a secure and organized manner. This can be achieved by using a dedicated filing system or a secure digital storage solution. Regular backups of these records are also recommended to prevent data loss.

The document also touches upon the importance of staying up-to-date with the latest tax regulations and accounting standards. This ensures that the company's financial reporting remains compliant and accurate.

Overall, the document provides a comprehensive guide to effective financial record-keeping, covering everything from initial data entry to final reporting and archiving. It is a valuable resource for any business owner or accountant looking to streamline their financial processes.

prepared by the method of Hendler and Anfinson (21). 147 mg. of glutamic acid were dissolved in 10 ml. of warm 0.1 N NaOH. Nitrogen was bubbled through the solution for a few minutes and 290 mg. of chloramine-T (sodio-p-toluenesulfochloramine) were added. The mixture was placed in a water bath at 50° C. for 15 min., cooled in ice, centrifuged and the insoluble p-toluenesulfonamide was discarded. To the supernatant solution containing succinic semialdehyde were added 28 ml. of 0.8% 2,4-dinitrophenylhydrazine in 2 N HCl. The suspension was filtered, and dried in vacuo. Purification of the compound was carried out by dissolving it in alcohol at 35° C., adding water at room temperature, and cooling in a dry ice-acetone mixture. The succinic semialdehyde-2,4-dinitrophenylhydrazone crystallized out on cooling; it was centrifuged, the supernatant was decanted, and the compound dried in vacuo. The M.P. of the prepared sample was 197-199° C.; the reported authentic M.P. is 199-201° C.



The first part of the document discusses the importance of maintaining accurate records of all transactions. It emphasizes that every entry should be supported by a valid receipt or invoice. This ensures transparency and allows for easy verification of the data.

In the second section, the author outlines the various methods used to collect and analyze the data. This includes both primary and secondary data collection techniques. The primary data was gathered through direct observation and interviews, while secondary data was obtained from existing reports and databases.

The third part of the document details the statistical analysis performed on the collected data. It describes the use of descriptive statistics to summarize the data and inferential statistics to test hypotheses. The results of these analyses are presented in a clear and concise manner, highlighting the key findings of the study.

Finally, the document concludes with a summary of the findings and their implications. It discusses the limitations of the study and suggests areas for future research. The author expresses confidence in the reliability of the data and the validity of the conclusions drawn from the analysis.

Methods

Preparation of cell-free enzyme fractions - 15 7-day old White Leghorn chick embryos weighing 0.7 g. each were blended in three fourths of their volume of Medium A containing 10 umoles/ml of mercaptoethanol in a Servall Omni-Mix homogenizer for 15 seconds. The resulting suspension was spun for 10 minutes at 12,500 r.p.m. at 3° C. in the 40 rotor of the Spinco Model L centrifuge. The supernatant from this centrifugation was centrifuged as above for 20 min. at 15,000 r.p.m. The supernatant from this second centrifugation constituted the S-15 fraction. The S-15 fraction was centrifuged for 90 minutes at 40,000 r.p.m. (105 X g). The sediment from this S-105 fraction was homogenized in 15% of the volume of the S-105 fraction of Medium A with added mercaptoethanol as per above and this suspension constituted the microsomal fraction. To prepare the pH 5 enzyme system, the S-105 supernatant was brought to pH 5.2 with 1 N acetic acid, centrifuged for 15 minutes at 20,000 r.p.m., and the sediment was taken up in 15% of the S-105 volume of Medium A with added mercaptoethanol.

Preparation of whole-cell incubations - The chick eggs were shelled, the embryos were cut up with scissors, and minced in a loose-fitting glass homogenizer at slow speed.



with either 2.5 or 3.0 ml of cold Krebs Ringer's phosphate solution per gram of tissue. Although a microscopic examination of the resultant suspension was not made, it was felt that this procedure yielded a predominantly whole-cell preparation.

Incubations were terminated by adding sufficient cold 10% or 20% TCA<sup>1</sup> to make a final concentration of 5%. In the case of the whole-cell preparations, the incubations were mixed thoroughly in a Waring blender (10 - 20 seconds) after precipitation of the protein.

Extraction of the collagen as gelatin - Collagen was separated from cold TCA-soluble material by washing the first precipitate 4 times with cold 5% TCA. The collagen was then solubilized in the gelatin form by heating in 5% TCA for 70 minutes at 90° C. after the method of Peterkofsky and Udenfriend (22). By this method these authors were consistently able to remove approximately 85% of the hydroxyproline from the precipitate. The hot-TCA extracts were extracted 4 times with anhydrous ether in order to remove the trichloroacetic acid.

Purification of hot TCA-insoluble protein and determination of specific activity - 4ml of 0.4N NaOH and

1. The abbreviation used is: TCA, trichloroacetic acid.

The first part of the document discusses the importance of maintaining accurate records of all transactions. It emphasizes that proper record-keeping is essential for the success of any business and for the protection of the interests of all parties involved. The text also mentions the need for regular audits and the importance of having a clear system in place for handling financial data.

In addition, the document highlights the role of technology in modern business operations. It suggests that investing in reliable software and hardware can significantly improve efficiency and reduce the risk of errors. The text also touches upon the importance of data security and the need to implement robust measures to protect sensitive information from unauthorized access.

Furthermore, the document discusses the importance of clear communication and collaboration between different departments within an organization. It notes that effective communication is key to ensuring that everyone is working towards the same goals and that any potential issues are identified and resolved promptly. The text also mentions the need for regular meetings and reports to keep everyone informed of the company's progress.

Finally, the document concludes by reiterating the importance of a strong financial foundation for any business. It stresses that a solid understanding of the company's financial health is crucial for making informed decisions and for ensuring long-term sustainability. The text also suggests that businesses should always be prepared for unexpected challenges and have a contingency plan in place.

The document ends with a call to action, encouraging all business owners and managers to take the time to review and implement the best practices outlined in the text. It expresses confidence that by following these guidelines, businesses can achieve greater success and stability in the future.

0.5 ml of a solution of unlabeled glutamate (10 mg/ml) were added to the hot TCA-insoluble proteins and the solutions were heated at 60° for 2 minutes. 1.0 ml of 50% TCA was added to precipitate the proteins and the vessels were cooled in the refrigerator for 15 minutes. The vessels were centrifuged and the precipitated protein was washed two times with 95% ethanol and once with ether-ethanol solution (1:3 by volume). The protein was then heated to 70° C. for 2 minutes, washed once with anhydrous ether and dried. The dried purified protein was then dissolved in 1.0 ml of anhydrous formic acid and the activities of weighed amounts were determined.

Separation of the hot TCA-soluble fraction into a small molecular weight and a macromolecular fraction -  
The hot-TCA extracts were evaporated to a volume of 2.0 ml and placed on the top of columns of Sephadex G-25, 16 cm X 1.5 cm, containing 9.3 g dry weight of gel with a water regain of 2.4 g H<sub>2</sub>O/g dry gel. The columns were eluted with 100 ml 0.2 M ammonium formate. 2 ml fractions were collected and alkaline hydrolysis according to the method of Hirs, Stein, and Moore (23) was carried out on each of the first 20 fractions eluted. 1.0 ml of 2.5 N NaOH was added to 0.13 ml aliquots of each 2.0 ml fraction to be hydrolyzed. The tubes were placed

The first part of the document discusses the importance of maintaining accurate records of all transactions. It emphasizes that proper record-keeping is essential for the success of any business and for the protection of the interests of all parties involved. The document then outlines the various methods and procedures for recording transactions, including the use of journals, ledgers, and other accounting systems. It also discusses the importance of regular audits and the role of the auditor in ensuring the accuracy and integrity of the financial records. The document concludes by reiterating the importance of transparency and accountability in all financial dealings.



in a 90° C. bath for 2.5 hours, during which time the contents evaporated to a volume of about 0.2 to 0.4 ml. The hydrolyzed fractions were then analyzed by a modified ninhydrin method. After 1.0 ml of 30% (by volume) acetic acid was added to each of the cooled tubes, in order to bring the solutions to about pH 5, 0.5 ml of ninhydrin reagent (prepared according to the method of Moore and Stein (24)) was added and the tubes were covered and placed in a boiling water bath for 15 minutes. After cooling, 2.5 ml of 1:1 (v/v) ethanol-water diluent were added to each tube and the absorbancy was read at 570 mμ against an ammonium formate standard. By this method the fractions containing the small molecular compounds were able to be identified by their respective ninhydrin reactive peaks. The macromolecular components came off the columns between about 11.0 ml and 16.0 ml. The smaller molecules started to be eluted at between 21.0 and 24.0 ml. The respective pooled macromolecular and small molecular fractions were lyophilized. The residue was dissolved in 1.0 ml of distilled water, and 0.1 ml and 0.2 ml samples were plated and counted.

All radioactivity was measured with a Nuclear-Chicago Geiger Counter, Model # 181 B.

Unless otherwise mentioned, acid hydrolysis was carried





out by making the solution of sample 6 N with respect to HCl and heating in a pressure cooker at approximately 15 lbs./sq. in., for three hours.

Preparation of the polyhydroxamic acid derivative of gelatin - This was carried out according to the method employed by Franzblau (15). 350 mg of hydroxylamine hydrochloride and 846 mg WSC were added per each 5.0 ml of aqueous gelatin solution (after evaporation of the gelatin solutions to convenient volumes). The resultant solutions were allowed to stand at room temperature, with occasional stirring, for 2.5 hours in the case of the bovine collagen and 2.0 hours in the chick embryo experiments. After standing, the protein was precipitated with cold acetone (care must be taken here since as much as one third of the collagen may not be precipitated), washed with ether and dried in vacuo. The dry hydroxamate derivative was then weighed and dissolved in varying volumes of warm distilled water as stated in the record of the individual experiments. The hydroxamic acid was either assayed by the procedure described at the end of the methods section of this paper or its presence was verified by placing a drop of the aqueous hydroxamate solution on a small quantity of ferric chloride in a test tube and observing the evolution of a wine-red color (the red reaction is specific for hydroxamic acid

The first part of the document is a preface, which is written in a very simple and direct style. It explains the purpose of the document and the author's intentions. The preface is followed by a list of chapters, which are arranged in a logical order. The chapters are:

- Chapter I: Introduction
- Chapter II: The first part of the document
- Chapter III: The second part of the document
- Chapter IV: The third part of the document
- Chapter V: The fourth part of the document
- Chapter VI: The fifth part of the document
- Chapter VII: The sixth part of the document
- Chapter VIII: The seventh part of the document
- Chapter IX: The eighth part of the document
- Chapter X: The ninth part of the document
- Chapter XI: The tenth part of the document
- Chapter XII: The eleventh part of the document
- Chapter XIII: The twelfth part of the document
- Chapter XIV: The thirteenth part of the document
- Chapter XV: The fourteenth part of the document
- Chapter XVI: The fifteenth part of the document
- Chapter XVII: The sixteenth part of the document
- Chapter XVIII: The seventeenth part of the document
- Chapter XIX: The eighteenth part of the document
- Chapter XX: The nineteenth part of the document
- Chapter XXI: The twentieth part of the document
- Chapter XXII: The twenty-first part of the document
- Chapter XXIII: The twenty-second part of the document
- Chapter XXIV: The twenty-third part of the document
- Chapter XXV: The twenty-fourth part of the document
- Chapter XXVI: The twenty-fifth part of the document
- Chapter XXVII: The twenty-sixth part of the document
- Chapter XXVIII: The twenty-seventh part of the document
- Chapter XXIX: The twenty-eighth part of the document
- Chapter XXX: The twenty-ninth part of the document
- Chapter XXXI: The thirtieth part of the document
- Chapter XXXII: The thirty-first part of the document
- Chapter XXXIII: The thirty-second part of the document
- Chapter XXXIV: The thirty-third part of the document
- Chapter XXXV: The thirty-fourth part of the document
- Chapter XXXVI: The thirty-fifth part of the document
- Chapter XXXVII: The thirty-sixth part of the document
- Chapter XXXVIII: The thirty-seventh part of the document
- Chapter XXXIX: The thirty-eighth part of the document
- Chapter XL: The thirty-ninth part of the document
- Chapter XLI: The fortieth part of the document
- Chapter XLII: The forty-first part of the document
- Chapter XLIII: The forty-second part of the document
- Chapter XLIV: The forty-third part of the document
- Chapter XLV: The forty-fourth part of the document
- Chapter XLVI: The forty-fifth part of the document
- Chapter XLVII: The forty-sixth part of the document
- Chapter XLVIII: The forty-seventh part of the document
- Chapter XLIX: The forty-eighth part of the document
- Chapter L: The forty-ninth part of the document
- Chapter LI: The fiftieth part of the document
- Chapter LII: The fifty-first part of the document
- Chapter LIII: The fifty-second part of the document
- Chapter LIV: The fifty-third part of the document
- Chapter LV: The fifty-fourth part of the document
- Chapter LVI: The fifty-fifth part of the document
- Chapter LVII: The fifty-sixth part of the document
- Chapter LVIII: The fifty-seventh part of the document
- Chapter LIX: The fifty-eighth part of the document
- Chapter LX: The fifty-ninth part of the document
- Chapter LXI: The sixtieth part of the document
- Chapter LXII: The sixty-first part of the document
- Chapter LXIII: The sixty-second part of the document
- Chapter LXIV: The sixty-third part of the document
- Chapter LXV: The sixty-fourth part of the document
- Chapter LXVI: The sixty-fifth part of the document
- Chapter LXVII: The sixty-sixth part of the document
- Chapter LXVIII: The sixty-seventh part of the document
- Chapter LXIX: The sixty-eighth part of the document
- Chapter LXX: The sixty-ninth part of the document
- Chapter LXXI: The seventieth part of the document
- Chapter LXXII: The seventy-first part of the document
- Chapter LXXIII: The seventy-second part of the document
- Chapter LXXIV: The seventy-third part of the document
- Chapter LXXV: The seventy-fourth part of the document
- Chapter LXXVI: The seventy-fifth part of the document
- Chapter LXXVII: The seventy-sixth part of the document
- Chapter LXXVIII: The seventy-seventh part of the document
- Chapter LXXIX: The seventy-eighth part of the document
- Chapter LXXX: The seventy-ninth part of the document
- Chapter LXXXI: The eightieth part of the document
- Chapter LXXXII: The eighty-first part of the document
- Chapter LXXXIII: The eighty-second part of the document
- Chapter LXXXIV: The eighty-third part of the document
- Chapter LXXXV: The eighty-fourth part of the document
- Chapter LXXXVI: The eighty-fifth part of the document
- Chapter LXXXVII: The eighty-sixth part of the document
- Chapter LXXXVIII: The eighty-seventh part of the document
- Chapter LXXXIX: The eighty-eighth part of the document
- Chapter LXXXX: The eighty-ninth part of the document
- Chapter LXXXXI: The ninetieth part of the document
- Chapter LXXXXII: The ninety-first part of the document
- Chapter LXXXXIII: The ninety-second part of the document
- Chapter LXXXXIV: The ninety-third part of the document
- Chapter LXXXXV: The ninety-fourth part of the document
- Chapter LXXXXVI: The ninety-fifth part of the document
- Chapter LXXXXVII: The ninety-sixth part of the document
- Chapter LXXXXVIII: The ninety-seventh part of the document
- Chapter LXXXXIX: The ninety-eighth part of the document
- Chapter LXXXXX: The ninety-ninth part of the document
- Chapter LXXXXXI: The hundredth part of the document

groups since hydroxylamine itself does not yield color).

Dinitrophenylation and Lossen rearrangement of the gelatin polyhydroxamate derivative - The method employed was that of Seifter, Gallop et al (25) as modified by Franzblau (15). The pH of the aqueous hydroxamate solution was adjusted to 8.0 with NaOH and an equal volume of 1% FDNB (v/v) in ethanol was added with constant stirring by means of a magnetic bar. The pH of the mixture was measured with a pH meter and the reaction was maintained at pH 8.0 by constant titration with NaOH, until the pH was stable (approximately 5.0 minutes). The solution containing the hydroxamate-DNP derivative was extracted twice with anhydrous ether and once with petroleum ether in order to extract the excess fluorodinitrobenzene. The extracted solution was made 0.1 N with respect to NaOH and heated for 2 minutes at 100° C. to promote Lossen rearrangement.

Formation and extraction of the 2,4-dinitrophenylhydrazones derivatives - To either one half or all of the hydrolysate an equal volume of 0.8% 2,4-dinitrophenylhydrazine in 2 N HCl was added, and the resulting solution containing the 2,4-dinitrophenylhydrazones was evaporated to a convenient volume in a flash evaporator (the temperature of the evaporating bath was kept below

The first part of the document discusses the importance of maintaining accurate records of all transactions. It emphasizes that every entry should be supported by a valid receipt or invoice. This ensures transparency and allows for easy verification of the data.

In the second section, the author outlines the various methods used to collect and analyze the data. This includes both manual data entry and the use of specialized software tools. The goal is to ensure that the data is both accurate and easy to interpret.

The third part of the document provides a detailed breakdown of the results. It shows that there has been a significant increase in sales over the period covered by the report. This is attributed to several factors, including improved marketing strategies and better customer service.

Finally, the document concludes with a series of recommendations for future actions. It suggests that the company should continue to invest in its marketing efforts and focus on building long-term relationships with its customers. This will help to ensure continued growth and success in the future.

35° C.). The concentrated solution was extracted 2 or 3 times with ethyl acetate and the extracts were combined. The ethyl acetate solution was extracted with 0.1 M borate buffer, pH 9.0; until at least 2 extractions had been performed with the resulting borate solution being basic as tested by pH paper. The borate solutions were pooled and acidified with 2 N HCl and extracted into ethyl acetate. Where mentioned, this final ethyl acetate extract was further purified by electrophoresing a known volume on Whatman 3 MM paper with a sample of the standard succinic semialdehyde-2,4-dinitrophenylhydrazone at pH 8.6 in sodium barbital buffer (5 g sodium barbital, 3.25g sodium acetate trihydrate, and 34.2 ml of 0.1 N HCl brought to 1 liter with distilled water) at 1200 v across 59 cm for approximately 2 hours. The band corresponding to the marker was completely eluted with a known volume of 0.1 M borate buffer (pH 9.0).

Thin layer chromatography - It was found that paper chromatography as employed by both Gallop (14) and Franzblau (15) was totally inadequate in this work, as both the bovine and the avian experiments yielded several 2,4-dinitrophenylhydrazone derivatives which acted similarly on electrophoresis and one of these also migrated very close to the unknown on chromatography. After trying without success several methods of paper chromatography with a





great variety of solvents recommended for separation of 2,4-dinitrophenylhydrazone derivatives, it was found that a high degree of resolution with excellent separation of the components was attained by thin layer chromatography with silica gel. The 2,4-dinitrophenylhydrazones were developed by spraying with alcoholic KOH (3.5 ml 15 N KOH to 20 ml with ethanol). Glass plates were spread with Silica gel G, 0.5 mm in thickness, by conventional methods. These were allowed to stand at room temperature for a few minutes and then dried in an oven at 110° C.. The samples to be chromatographed were extracted from the borate buffer into the ethyl acetate as described above and a measured volume of the ethyl acetate was placed on the silica covered glass plates in a connected series of very small drops thus forming a line. Although it may not be necessary, the chromatograms were run in the dark. The solvent used for separation was tertiary amyl alcohol-ethanol-water (5:1:4, top layer). The band corresponding to succinic semialdehyde-2,4-dinitrophenylhydrazone was scraped off the plate into a test tube and eluted into a known volume of 0.1 M borate buffer, pH 9.0.<sup>1</sup> Part

1. It is interesting to note that in order to extract the succinic semialdehyde-2,4-dinitrophenylhydrazone from the silica gel with ethyl acetate, a drop of HCl must be added, a fact that suggests that, for an unknown reason, the carboxyl group is converted from the acid to the salt form during the process of chromatography. In any case, complete elution is readily attained with the borate buffer.



1. The first part of the document discusses the importance of maintaining accurate records of all transactions and activities. It emphasizes the need for transparency and accountability in financial reporting.

2. The second part of the document outlines the various methods and techniques used to collect and analyze data. It includes a detailed description of the sampling process and the statistical tools employed to interpret the results.

3. The third part of the document presents the findings of the study. It provides a comprehensive overview of the data collected and the conclusions drawn from the analysis. The results indicate that there is a significant correlation between the variables studied.

4. The fourth part of the document discusses the implications of the findings and offers recommendations for future research. It suggests that further studies should be conducted to explore the underlying causes of the observed trends and to develop effective strategies to address them.

5. The final part of the document provides a summary of the key points and reiterates the importance of the research. It concludes by stating that the findings have important implications for the field and that the research has contributed to the understanding of the subject matter.

of this solution was assayed and part extracted into ethyl acetate and chromatographed with a sample of the authentic succinic semialdehyde-2,4-dinitrophenylhydrazone (the authentic sample had also been purified by chromatography in the tertiary amyl alcohol-ethanol-water system, eluted into borate solution, and extracted with ethyl acetate prior to its use as a standard marker) in the tertiary amyl alcohol-ethanol-water (5:1:4) solvent and also in a solvent of normal butanol saturated with water. In placing the experimental solution on the origin of the chromatogram, a line 3 cm long was made. The authentic marker was then placed in a similar series of dots forming a line 3 cm in length such that 1.5 cm of the marker was superimposed on the terminal 1.5 cm of the experimental line and 1.5 cm was free. Failure of the experimental sample to separate from the standard compound in two solvent systems was accepted as proof of identity.

Desalting and isolation of amino acids - The aqueous phase remaining after extraction of the hydrolysate with ethyl acetate was evaporated to a volume of two ml and placed on top of a 20 ml column containing Dowex 50-8X ( $H^+$ ). The column was rinsed with 80 to 100 ml of distilled water and the amino acids were eluted with 50 ml of 2 N  $NH_4OH$ . The eluate was evaporated to 5.0 ml and known volumes

The first part of the document discusses the importance of maintaining accurate records of all transactions. It emphasizes that every entry should be supported by a valid receipt or invoice. This not only helps in tracking expenses but also serves as a legal proof in case of an audit. The document further outlines the steps to be followed when recording transactions, including the need to categorize them correctly and ensure that the debits and credits are balanced. It also mentions the importance of regular reconciliation to identify any discrepancies early on. The second part of the document provides a detailed guide on how to prepare a budget. It suggests that a budget should be based on realistic assumptions and should include provisions for unexpected expenses. The document also discusses the importance of monitoring the budget regularly and making adjustments as needed. Finally, the document concludes by stating that good financial management is essential for the long-term success of any business or organization.

were placed on Whatman 3 MM paper and electrophoresed in pH 6.0 pyridine acetate buffer (100 ml pyridine and 10 ml glacial acetic acid brought to 1 liter with distilled water) at 2000 volts across 47 cm for 1 hour 15 minutes. Alpha,gamma-diaminobutyric acid was identified by its similar migration to a marker of the commercially prepared compound and by its characteristic red-brown staining reaction when the experimental marker section of the paper was sprayed with 0.5% ninhydrin in acetone. The area identified as alpha,gamma-diaminobutyric acid and those of the other amino acids desired were cut out and eluted from the electrophorogram with known volumes of water. These solutions were then assayed, and, when pertinent, their radioactivity was assessed.

Assay procedures - Hydroxamic acid assay was carried out according to the ferric perchlorate method of Seifter, Gallop et al (25). The solution (1 ml) containing 0.2 to 2.0  $\mu$ moles of hydroxamic acid was mixed with 2 ml of ferric perchlorate reagent (prepared by dissolving 0.8 g of pure iron wire in 10 ml of warm 60 % perchloric acid and bringing the solution to 100 ml with ethanol) diluted 1:1 with distilled water. After 5 minutes it was read at 505 m $\mu$  against a reagent blank. A standard curve was simultaneously prepared from a standard (2  $\mu$ moles/ml)

The first part of the document is a general introduction to the project. It describes the objectives and the scope of the work. The second part is a detailed description of the methodology used in the study. This includes a discussion of the data collection methods, the analysis techniques, and the results of the study. The third part is a conclusion and a list of references.

The methodology section is particularly important as it details the steps taken to ensure the reliability and validity of the research. It covers the selection of participants, the design of the study, and the procedures for data collection and analysis. The results section presents the findings of the study, which are discussed in the context of the research objectives and the existing literature.

The conclusion summarizes the key findings and provides a final assessment of the study's contributions. The references list the sources used in the research, providing a clear path for further exploration of the topic.

solution of acethydroxamic acid. This method measures only hydroxamic acids and gives no color from hydroxylamine. Optical density readings were performed with a Zeiss Spectrophotometer.

Glutamic acid and alpha,gamma-diaminobutyric acid were measured by a modified ninhydrin procedure as follows: Preparation of Ninhydrin reagent; 40 mg of  $\text{SnCl}_2$  were dissolved in 25.0 ml pH 5.0 acetate buffer and 0.667g of ninhydrin were dissolved in 25.0 ml of methyl cellosolve (monomethyl ether of ethylene glycol). Just before the assay 8.3 ml of the  $\text{SnCl}_2$  solution were mixed with all of the ninhydrin solution. Assay: to 0.5 ml samples, 0.5 ml pH 5.0 acetate buffer and 1.0 ml ninhydrin reagent were added in test tubes which had been rinsed several times with distilled water and dried. The solutions were mixed well, the test tubes were capped with glass marbles and they were placed in a covered boiling water bath for 15 minutes. At the end of this period they were cooled to room temperature in ice water and the volume was brought to 5.0 ml with 95% ethanol. The solutions were mixed well and the O.D. was read at 570 m $\mu$  against a reagent blank. Standard curves (0.05-0.50  $\mu$ moles) of glutamic acid and alpha,gamma-diaminobutyric acid were carried out each time the assay was done.

... ..  
... ..  
... ..

... ..  
... ..

... ..  
... ..

... ..  
... ..

... ..  
... ..

... ..  
... ..



Hydroxyproline was measured according to the method of Newman & Logan (26) as modified by Leach (27) at 1/10 volume. To the sample brought to 0.1 ml with water was added 0.1 ml of 2.5 N NaOH. The tubes were placed in a 40° C. water bath, and when the contents of the tubes reached 40° C. (3-5 minutes), 0.1 ml of 6% H<sub>2</sub>O<sub>2</sub> was added and the solution was mixed. After another 10 minutes in the bath, the solutions were cooled and 0.4 ml of 3 N H<sub>2</sub>SO<sub>4</sub> and 0.2 ml of 5% p-dimethylaminobenzaldehyde in n-propanol were added. The tubes were placed in a 70° C. bath for 20 minutes, cooled and the O.D. was read at 555 mμ. Three different concentrations of each unknown solution were assayed and a standard curve (0.01 - 0.05 μmoles) was constructed with each assay.

Proline was measured by the method of Chinard (28). To 1.0 ml of the sample to be assayed, 1.0 ml of glacial acetic acid and 1.0 ml of ninhydrin reagent were added (each ml of reagent contained 0.4 ml of 6 N H<sub>3</sub>PO<sub>4</sub> and 0.6 ml of glacial acetic acid; 25 mg of ninhydrin were added per ml of this acid mixture and the mixture was heated to about 70° C. to insure solution of the ninhydrin). The tubes were capped and heated at 100° C. for 1 hour, after which time they were cooled to room temperature, and brought to a volume of 5.0 ml with glacial acetic acid. The O.D. was read at 515 mμ against a reagent blank. A standard



THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

THE UNIVERSITY OF CHICAGO PRESS

(31)

curve (0.02-0.2  $\mu$ moles) was constructed and 3 different concentrations of each unknown solution were assayed.

Succinic semialdehyde was assayed according to the procedure described by Waelsch (29). The solution to be assayed was brought to 5.0 ml in borate buffer pH 9.0, and mixed with 2 ml of alcoholic KOH (3.5 ml of 15 N KOH made to 20 ml with absolute ethanol). After mixing the tube was placed in a water bath at 25<sup>o</sup> C. Exactly 2 minutes after addition of the alkali the absorbancy was measured at 420 m $\mu$  against a blank containing 5 ml of borate buffer and 2 ml of alcoholic KOH. A standard curve was simultaneously prepared from a standard solution (0.1  $\mu$ moles/ml) of the authentic compound in borate buffer.



Results

I. Incorporation of C<sup>14</sup>-glutamic acid into the collagen-containing hot trichloroacetic acid-extractable fraction by a cell-free system from 7-day chick embryos.

Cell-free enzyme fractions were prepared from 15 7-day old White Leghorn chick embryos and incubated as described in Table II. The incubations were terminated by precipitating the protein with cold TCA. The precipitated protein was then washed, 20  $\mu$ moles of unlabeled glutamic acid carrier were added to each vessel, and the protein was extracted for 80 minutes with hot TCA. The hot TCA-extracts were extracted with ether, and their radioactivity was measured. That protein which was not soluble in hot TCA was purified and a determination of its specific activity was made. The gelatin-containing hot TCA-extracts were separated into small molecular weight and macromolecular fractions, and the radioactivity of each was determined (Table III).

From the data presented in Table II it is evident that the cell-free 7-day old chick embryo incubation using a pH 5 precipitated enzyme system in conjunction with the microsomal fraction, incorporated C<sup>14</sup>-glutamic acid into the collagen-containing hot TCA-extractable protein fraction. The amount of incorporation, however, was small. The spec-

MEMORANDUM

TO : SAC, NEW YORK

FROM : SAC, NEW YORK

SUBJECT: [Illegible]

[The remainder of the memorandum text is illegible due to extreme blurriness.]

Table II

Incorporation of C<sup>14</sup>-glutamic acid into the collagen-containing hot TCA-extractable fraction by a cell-free system from 7-day chick embryos and specific activity of the purified non-collagenous protein.

All incubations contained in a final volume of 6.0 ml: 0.30 ml Tris buffer 1.0 M, pH 7.7; 0.12 ml ATP, 50  $\mu$ moles/ml; 0.12 ml magnesium acetate 0.3 M; 0.63 ml KCl, 1.0 M; 0.12 ml GTP, 2.5  $\mu$ moles/ml; 0.24 ml glutathione, 200  $\mu$ moles/ml; 0.33 ml sodium phosphoenolpyruvate, 100  $\mu$ moles/ml; 0.12 ml pyruvate kinase, 1 mg/ml; 0.12 ml ascorbic acid, 50  $\mu$ moles/ml; 0.15 ml cysteine, 2  $\mu$ moles/ml; 0.30 ml amino acid mixture; containing-L-proline, argenine, lysine, threonine, serine, leucine, phenylalanine, methionine, valine, asparagine, aspartic acid, isoleucine, histidine, tryptophan, alanine, tyrosine, and glycine at a concentration of 1.0  $\mu$ mole/ml. 0.50 ml of C<sup>14</sup>-glutamic acid (28 $\mu$ c/ $\mu$ mole, 2.24 X 10<sup>6</sup> cpm/ml) was added to vessels 1 and 2 before incubation and to vessel 3 after precipitation of the protein.

The incubation was carried out at 38° C. for 90 minutes.

<u>Vessel #</u>	<u>Contents</u>			<u>Total activity of collagenous protein</u>	<u>Specific activity of non-collagenous protein</u>
	<u>pH 5 fraction</u>	<u>Microsome fraction</u>	<u>S-15 fraction</u>	<u>(cpm)</u>	<u>(cpm/mg)</u>
1 (pH 5)	1.20ml	1.80ml	-	1290	100
2 (S-15)	-	1.80ml	1.20ml	781	36
3 (pH 5 control)	1.20ml	1.80ml	-	693	32

CHAPTER 10

1. The first part of the chapter discusses the importance of the...  
 2. The second part of the chapter discusses the importance of the...  
 3. The third part of the chapter discusses the importance of the...

4. The fourth part of the chapter discusses the importance of the...  
 5. The fifth part of the chapter discusses the importance of the...  
 6. The sixth part of the chapter discusses the importance of the...  
 7. The seventh part of the chapter discusses the importance of the...  
 8. The eighth part of the chapter discusses the importance of the...  
 9. The ninth part of the chapter discusses the importance of the...  
 10. The tenth part of the chapter discusses the importance of the...

11. The eleventh part of the chapter discusses the importance of the...  
 12. The twelfth part of the chapter discusses the importance of the...  
 13. The thirteenth part of the chapter discusses the importance of the...  
 14. The fourteenth part of the chapter discusses the importance of the...  
 15. The fifteenth part of the chapter discusses the importance of the...

16. The sixteenth part of the chapter discusses the importance of the...  
 17. The seventeenth part of the chapter discusses the importance of the...  
 18. The eighteenth part of the chapter discusses the importance of the...  
 19. The nineteenth part of the chapter discusses the importance of the...  
 20. The twentieth part of the chapter discusses the importance of the...

21. The twenty-first part of the chapter discusses the importance of the...  
 22. The twenty-second part of the chapter discusses the importance of the...  
 23. The twenty-third part of the chapter discusses the importance of the...  
 24. The twenty-fourth part of the chapter discusses the importance of the...  
 25. The twenty-fifth part of the chapter discusses the importance of the...

Table III

The radioactivity in the macromolecular and small molecular fractions separated by means of a Sephadex G-25 column.

<u>Fraction</u>	<u>Macromolecular activity</u> <u>(total cpm)</u>	<u>Small molecular activity</u> <u>(total cpm)</u>
pH 5 & Microsomes	59	219
S-15 & Microsomes	104	118
pH 5 control	Background	119





ific activities of the non-collagenous protein fraction verify the fact that the extent of incorporation was not impressive. Due to the low incorporation of the system, it was decided to employ whole-cell systems from minces of embryos in the future.

Only a fraction (about 20%) of the total hot TCA extract radioactivity was recovered after Sephadex fractionation (Table III). Much of the activity recovered was present in the small molecular fractions, but there was some incorporation into macromolecular protein (eg. collagen). In a larger incubation employing  $C^{14}$ -glutamic acid of a higher specific activity it is expected that a greater incorporation into the macromolecular fraction would take place.

II. Evidence for the presence of gamma-glutamyl bonds in a commercially prepared collagen obtained from bovine Achilles tendon.

For the purpose of testing the rearrangement procedure and the methods of recovery of the rearrangement products, it was decided to submit some commercially prepared collagen to the various manipulations that will be employed with the chick embryo gelatin.

2.0 g of bovine Achilles tendon collagen were suspended in 50 ml. of distilled water and autoclaved at  $125^{\circ}$  C. for



3 hours. As some tissue remained at the end of this procedure, 20 additional ml of distilled water were added to the above solution and it was heated again under the same conditions. The solution was cooled, centrifuged, and the gelatin-containing supernatant was decanted. One half of the gelatin solution was hydrolyzed and served as the unrearranged control. The other half of the gelatin solution was evaporated to 5.0 ml and the polyhydroxamate derivative was formed. The dry gelatin hydroxamate weighed 949.2 mg and was dissolved in 62.5 ml of warm distilled water. There were 781  $\mu$ moles of hydroxamic acid in the solution submitted to Lossen rearrangement. The polyhydroxamate-DNP derivative was formed, the rearrangement was promoted by heating at alkaline pH, and the solution was hydrolyzed.

The hydrolysates of both the rearranged and the unrearranged gelatins were divided into two equal parts. One half of each hydrolysate was evaporated to 10 ml, 1.0 ml of this volume was electrophoresed at pH 6.0 without previous desalting, and glutamic acid, alpha, gamma-diaminobutyric acid and hydroxyproline were eluted and assayed. An equal volume of 0.8% 2,4-dinitrophenylhydrazine in 2 N HCl was added to the other half of each hydrolysate and the 2,4-dinitrophenylhydrazone derivatives were extracted



Table IV

Analysis of some of the products in the Lossen rearranged dinitrophenylhydroxamate derivative of gelatin derived from bovine Achilles tendon as compared with an unrearranged control gelatin.

(Expressed as total number of  $\mu$ moles recovered)

<u>Compound</u>	<u>Rearranged</u>	<u>Control</u>	<u>Rearranged values adjusted to hydroxyproline recovered from Control</u>	<u>Difference</u>
Hydroxyproline	467	676	676	-
Glutamic acid	284	612	411	-201
alpha,gamma- diaminobutyric acid	13.1	0	19	+19
Succinic semi- aldehyde	15.9	4.4	23	+19



with ethyl acetate. A portion of the final ethyl acetate solution after extraction with borate buffer was chromatographed and the succinic semialdehyde-2,4-dinitrophenylhydrazone was identified and assayed. In this case the above compound was eluted from the silica gel in 2.0 ml of ethyl acetate to which had been added a drop of 2 N HCl.

It may be seen from Table IV that some succinic semialdehyde was present in the hydrolyzed gelatin that had not been rearranged. The loss of glutamic acid in the rearranged as compared to the unrearranged control is not accounted for by the appearance of an equivalent quantity of glutamic acid rearrangement products in the rearranged gelatin. Because of this discrepancy no quantitative statements concerning the amount of gamma-linked glutamic acid present in the bovine Achilles tendon collagen can be made. It is evident that Lossen rearrangement of glutamic acid units took place, as witnessed by the appearance of alpha,gamma-diaminobutyric acid (the alpha-linked rearrangement product) in the rearranged gelatin and not in the unrearranged control. The rearranged gelatin also yielded 5.2 times as much succinic semialdehyde (the gamma-linked rearrangement product) as was endogenously present in the control.





III. The application of the Lossen rearrangement to the hot TCA-extractable protein derived from 14 day old chick embryos after incubation with  $C^{14}$ -glutamate.

It was decided to do an isotopic experiment with a 14 day old chick whole-cell preparation in order to see if the gamma-linked rearrangement product could be isolated from this age embryo, and also to see if the preparation would incorporate radioactivity into gamma-linked glutamyl units. As this experiment was designed as a quick pilot experiment to indicate where future investigation should be focused, neither a rearrangement control nor a radioactivity incorporation control was included.

A whole-cell incubation was prepared by homogenizing 20 14 day old White Leghorn chick embryos weighing approximately 9.9 g each with 2.5 ml of Krebs Ringer's phosphate solution per gram of tissue. To the resulting suspension were added 1.2 ml of a solution of  $C^{14}$ -glutamic acid ( $3.8 \times 10^6$  cpm/ml, 205  $\mu$ curies/ $\mu$ mole) and 0.9 ml of a solution of  $C^{14}$ -glutamic acid ( $3 \times 10^6$  cpm/ml, 170  $\mu$ curies/ $\mu$ mole), the total activity being equivalent to 11  $\mu$ curies. This whole-cell suspension was incubated at  $37^\circ$  C. for 1.5 hours with constant agitation. After precipitation, the protein was washed with cold TCA and extracted into hot TCA. The gelatin-containing solution was dialyzed against distilled water at  $4^\circ$  C. for 3 hours with one change in

The first part of the document discusses the importance of maintaining accurate records of all transactions. It emphasizes that every entry should be supported by a valid receipt or invoice. This ensures transparency and allows for easy verification of the data.

In the second section, the author outlines the various methods used to collect and analyze the data. This includes both manual data entry and the use of specialized software tools. The goal is to ensure that the data is both accurate and easy to interpret.

The third part of the document provides a detailed breakdown of the results. It shows that there is a significant correlation between the variables being studied. This finding is supported by statistical analysis and is consistent with previous research in the field.

Finally, the document concludes with a series of recommendations for future research. It suggests that further studies should be conducted to explore the underlying causes of the observed trends. This will help to refine the current model and provide a more comprehensive understanding of the phenomenon being studied.

the external phase midway in the dialysis. The total radioactivity of the hot TCA extract decreased from 13,000 cpm before dialysis to 8,500 cpm after dialysis.

The dialyzed solution was evaporated to 10 ml and the polyhydroxamate derivative was prepared as before. The dry hydroxamate weighing 1.0590 g was dissolved in 20 ml of warm distilled water (ferric chloride test was positive for hydroxamic acid), and the polyhydroxamate-DNP derivative was formed. A fine yellow gelatin-hydroxamate-DNP precipitate formed while the FDNB was being added to the hydroxamate solution. Franzblau noted that this occasionally happened when the pH was not well titrated, and that this precipitation limited the extent of the dinitrophenylation of the hydroxamate. During the extraction of the excess FDNB with anhydrous ether, a laboratory accident occurred which resulted in the loss of much of the gelatin-DNP derivative. An attempt to extract the recovered material with petroleum ether was abandoned because a difficult emulsion formed. NaOH was added to the solution and Lossen rearrangement was promoted by heating - the gelatin went back into solution on heating. The rearranged solution was evaporated to a volume of 20 ml, made 6 N with respect to HCl and hydrolyzed at 110<sup>o</sup> C. for 16 hours in vacuo.



Table V

A partial analysis of the pertinent products obtained from the Lossen rearrangement of the hydroxamate-DNP derivative of gelatin of 14 day old chick embryos.

<u>Compound</u>	<u>Total yield</u> ( $\mu$ moles)	<u>Total activity</u> (cpm)
Glutamic acid	not assayed	195
Hydroxyproline	138.5	not counted
alpha,gamma-diamino butyric acid	not assayed	109
Succinic semialdehyde	3.3	0

Y 1000

-----  
-----  
-----

-----	-----	-----
-----	-----	-----
-----	-----	-----
-----	-----	-----
-----	-----	-----

The hydrolysate was filtered and the 2,4-dinitrophenylhydrazone derivative was prepared from the entire hydrolysate. The resulting solution was evaporated to a convenient volume for extraction, and the 2,4-dinitrophenylhydrazones were extracted in ethyl acetate. This solution was re-extracted and chromatographed in the usual manner. The presence of succinic semialdehyde was verified in both the tertiary amyl alcohol - ethanol - water and in the water saturated butanol solvents. The compound was assayed and its radioactivity was determined. A known volume of hydrolysate remaining after ~~ex~~traction was electrophoresed at pH 6.0, the hydroxyproline was eluted and assayed, and the glutamic acid was eluted and its radioactivity determined. The resolution in the area of the electrophorogram containing the dibasic amino acids was poor, probably because of the presence of a large amount of salt. Therefore the remainder of the sample which had not been electrophoresed was desalted and then electrophoresed at pH 6.0. A small amount of alpha,gamma-diaminobutyric acid was identified, eluted, and its radioactivity determined.

In this experiment succinic semialdehyde was isolated and identified. However, in the absence of an unrearranged control, one cannot state, at this juncture, that the gamma-





linkage must necessarily exist in gelatin derived from the chick embryo.

It is interesting to note from Table V that there was radioactivity present in the alpha,gamma-diaminobutyric acid and that none existed in the succinic semialdehyde. If the counts in alpha,gamma-diaminobutyric acid represent incorporation and if this succinic semialdehyde originated as a result of Lossen rearrangement of gamma-glutamyl bonds in gelatin, there appear to be three possible explanations for the absence of radioactivity from the succinic semialdehyde. First, this succinic semialdehyde might come from collagen already present in the embryos and not from newly synthesized collagen. However, this system is known to synthesize collagen actively, as Lukens (30) has found that a 14-day whole-cell system from chick embryos will incorporate labeled proline into hydroxyproline. As hydroxyproline, for all practical purposes in this instance, is only present in collagen, this incorporation serves as a proof of collagen synthesis. The second explanation that must be considered is that there is a delay between the incorporation of C<sup>14</sup>-glutamate into alpha-linkage and its incorporation into gamma-linkage, such that a measurable amount of activity is not incorporated into the gamma



rearrangement product during the period of the incubation. This possibility will be discussed later. The third explanation is that the diaminobutyric acid may have been derived from non-collagenous protein, and the collagen may not be highly enough labeled.

The low activity presented in Table V for glutamic acid in relation to that of alpha,gamma-diaminobutyric acid may be due, in part, to the fact that all of the glutamic acid was plated for counting and that the solution was not desalted before electrophoresis, as was the alpha,gamma-diaminobutyric acid sample. Both of these factors would tend to increase self-absorption. As most of the gelatin-hydroxamate-DNP derivative was lost, these values represent only a small amount of the total activity of the original sample.

In any event, it may be stated that the 14-day chick embryo whole-cell system apparently incorporated  $C^{14}$ -glutamic acid into hot TCA-extractable protein, and that a small portion of this activity was recoverable, after Lossen rearrangement, as the alpha-linkage rearrangement product.

**IV. The presence of the gamma-glutamyl peptide linkage in the hot TCA-extractable protein of the 14-day chick embryo.**

It was now decided to establish more rigorously whether or not the collagen-containing fraction of 14 day old chick embryos contained glutamic acid in the gamma-glutamyl linkage.

The first part of the document discusses the importance of maintaining accurate records of all transactions. It emphasizes that every entry should be supported by a valid receipt or invoice. This ensures transparency and allows for easy verification of the data.

Furthermore, it is noted that the records should be kept in a secure and accessible format. Regular backups are recommended to prevent data loss in the event of a system failure or disaster. The document also mentions the need for periodic audits to ensure the integrity and accuracy of the information stored.

In addition, the text highlights the role of technology in streamlining record-keeping processes. Modern accounting software can automate many tasks, reducing the risk of human error and saving valuable time. However, it is stressed that users must be properly trained and that the software is regularly updated to address any security vulnerabilities.

Overall, the document serves as a comprehensive guide for anyone responsible for financial record-keeping. It provides clear instructions and best practices to ensure that all records are accurate, complete, and secure. By following these guidelines, organizations can maintain a high level of financial transparency and accountability.

A whole-cell incubation was prepared from 20 14-day White Leghorn chick embryos weighing 13.8 g each with 3 ml of Krebs Ringer's phosphate solution per gram of tissue. The resulting suspension was incubated at 37° C. for two hours. The protein was precipitated, washed and extracted twice with 5% TCA. The two hot TCA extracts were combined and extracted with ether - during the ether extraction a white precipitate formed which was filtered out. The gelatin rich solution was dialyzed against distilled water at 4° C. for three hours with the external phase being changed every hour. This dialyzed solution was evaporated to 20 ml and redialyzed for 11 hours with one change of the external phase. The solution was then evaporated to 12 ml and the polyhydroxamate derivative was made. The hydroxamate derivative weighing 1.6157 g was dissolved in 106 ml of warm water and the solution was divided into two equal portions - one to be rearranged and one to remain as the unrearranged control. There were 463  $\mu$ moles of hydroxamic acid groups in the 52 ml submitted to Lossen rearrangement.

The control hydroxamate was hydrolyzed as usual after bubbling nitrogen through the sample for five minutes. The other half of the gelatin hydroxamate was subjected to

The first part of the document discusses the importance of maintaining accurate records of all transactions. It emphasizes that every entry should be supported by a valid receipt or invoice. This ensures transparency and allows for easy verification of the data.

In the second section, the author outlines the various methods used to collect and analyze the data. This includes both manual data entry and the use of specialized software tools. The goal is to ensure that the data is both accurate and easy to interpret.

The third part of the document provides a detailed breakdown of the results. It shows that there has been a significant increase in sales over the period covered by the report. This is attributed to several factors, including improved marketing strategies and better customer service.

Finally, the document concludes with a series of recommendations for future actions. These include continuing to invest in marketing, maintaining high standards of customer service, and regularly reviewing financial performance to identify areas for improvement.

the Lossen rearrangement according to the usual procedure and was then hydrolyzed in the same manner as the control. The 2,4-dinitrophenylhydrazone derivative of each hydrolysate was made, extracted, electrophoresed and chromatographed. Succinic semialdehyde-2,4-dinitrophenylhydrazone was identified in the rearranged gelatin but not in the control. The identity of the compound was verified in two solvent systems, and it was assayed. After removal of the 2,4-dinitrophenylhydrazones, the aqueous phase was evaporated to 2.0 ml and desalted. After desalting, duplicate electrophoreses, elutions and assays were performed in order to demonstrate that the results obtained were not the result of differences in the degree of elution from the electrophorograms but represented real differences in the amino acid compositions of the control and rearranged gelatins.

In this experiment the presence of gamma-linked glutamic acid in the gelatin-rich hot TCA-extractable protein from the 14-day chick embryo was affirmed. From Table VI it is evident that there was a great deal less glutamic acid recovered from the rearranged gelatin than was recovered from the control. The fact that the amount of proline remained relatively constant, as indeed





Table VI

Analysis of the products obtained on hydrolysis of the  
Lossen rearranged gelatin from the 14-day chick embryo  
as compared with a suitable unrearranged control.

<u>Compound</u>	<u>Total <math>\mu</math>moles recovered</u>						<u>Difference*</u>
	<u>Control</u>			<u>Rearranged</u>			
	<u>Gelatin</u>		<u>Average</u>	<u>Gelatin</u>		<u>Average</u>	
<u>Duplicates**</u>		<u>Duplicates</u>					
Hydroxyproline	235	240	238	228	240	234	N.S.***
Proline	480	485	483	492	504	498	+15
Glutamic acid	413	435	424	295	275	285	-139
Alpha, gamma- diaminobutyric acid	0	0	0	8.34	9.34	8.84	+8.84
Succinic semialdehyde		0	0	12.21****		12.21	+12.21
<u><math>\mu</math>moles Hydroxyproline</u>	-	-	-	18.7	19.7	19.2	-
<u><math>\mu</math>moles Succinic semialdehyde</u>							

\* Difference between control average and rearranged average.

\*\* Duplicates = Duplicate elutions from duplicate electrophoreses.

\*\*\* N.S. = Not significant.

\*\*\*\*A single isolation and assay was done with succinic semialdehyde.



it should for it is not involved in the rearrangement procedure, indicates that the decrease in glutamic acid units is real. The appearance of 21  $\mu$ moles of glutamic acid rearrangement products still leaves 118  $\mu$ moles unaccounted for out of the 139  $\mu$ moles of the glutamic acid that disappeared as a result of the rearrangement. As the alpha,gamma-diaminobutyric acid was isolated from the same electrophorograms in the same way as the other amino acids that were analyzed, and as there is no evidence that this compound is unstable, it would not seem as though the discrepancy can be explained in terms of any large amounts of alpha,gamma-diaminobutyric acid being unaccounted for. It seems likely that the failure to account for the missing residues is due to the incomplete recovery of succinic semialdehyde as the phenylhydrazone. If the discrepancy lies in the unrecovered succinic semialdehyde-2,4-dinitrophenylhydrazone, then it probably lies in the loss of succinic semialdehyde during hydrolysis, the incomplete formation of the hydrazone derivative, or in incomplete extraction procedures. Precautions against photic degradation or heat breakdown of the hydrazone derivative were observed during the experimental procedure. Because of the aforementioned discrepancy, an absolute statement of the number of gamma-linkages in gelatin cannot be made. However, one



may conclude that there is at least one glutamic acid residue in gamma-peptide linkage for every 19 hydroxyproline residues in the gelatin-containing fraction derived from the 14-day chick embryo.

It might be worth repeating at this point that these investigations were focused on the establishment of an in vivo preparation in which the gamma-glutamyl bond formation could be studied. Therefore it is not quantitative recovery of succinic semialdehyde that is important but rather the attainment of a high degree of purity for this compound.



Discussion

Two of the questions that are raised by the presence of gamma-glutamyl linkages are that of how and when in the process of protein synthesis, as we conceive of it today, the bonds are formed. It is hoped that the work reviewed in the experimental section of this paper will, with greater refinement, lead to the establishment of a whole-cell system which may be used as a tool with which to study the enzyme or enzymes responsible for the formation of this gamma linkage.

Figure 5. is a partial representation of the mechanism of protein synthesis as it is conceived of today. There are several stages in the synthetic process where the formation of the gamma-linkage may take place. First it may take place at the level of the formation of the aminoacyl-s-RNA complex. It is known from Ochoa (31) that the aminoacyladenylate-enzyme complex interacts with the corresponding transfer RNA whereby the amino acid is esterified through its carboxyl group to a hydroxyl residue on the terminal adenosine moiety of transfer RNA.

In the formation of the gamma-glutamyl bond, it may be that the glutamic acid attaches to the s-RNA by its gamma-carboxyl group instead of by its alpha-carboxyl





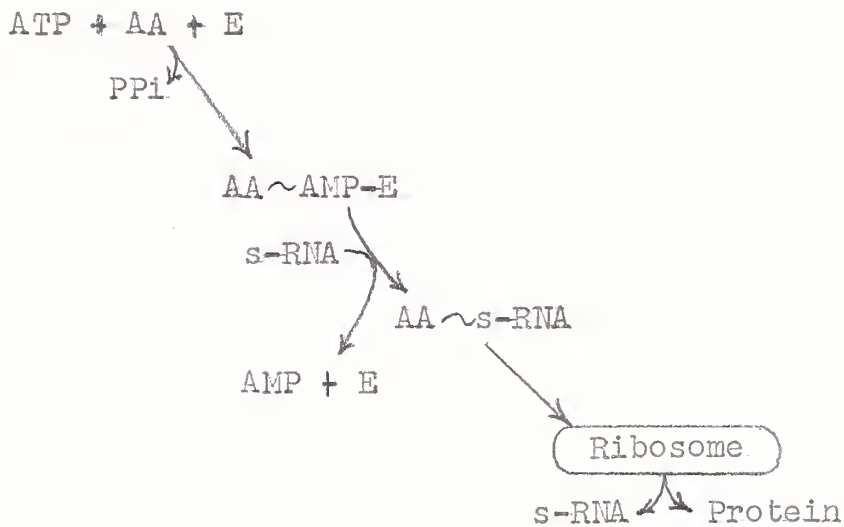


Fig. 5. Diagram of Protein Synthesis. E = enzyme, AA = amino acid, P<sub>i</sub> = inorganic pyrophosphate, s-RNA = transfer RNA.







group. In the formation of the usual peptide linkage there is a specific distance between the carboxyl group-s-RNA bond and the amino group which is entering into peptide linkage with the emerging protein. If we are to assume the formation of a glutamyl-gamma-carboxyl-s-RNA, then we must assume that the increased amino group to carboxyl group distance is accounted for either by folding of the amino acid or by bending of the s-RNA. If one is to assume that it is at this level that the gamma-linkage is formed, then it may be necessary to assume that there is a specific s-RNA for the glutamic acid entering gamma-linkage with a different code triplet or triplet order from the s-RNA of the glutamic acid entering into alpha-linkage.

A second possibility is that the gamma-linkage may be formed by an intramolecular rearrangement of the alpha-linked amino acid. Such a rearrangement could take place before or after polymerization of collagen subunits into tropocollagen. The assumption in this case is that glutamic acid would have been incorporated into alpha-peptide linkage, an imide might then be formed and finally the alpha-carboxyl-nitrogen bond would be hydrolyzed leaving the gamma-carboxyl group in peptide linkage. This would be the same type of reaction that was discussed in the introduction in relation to the possibility of



artificial formation of the gamma-linkage during the synthesis of the hydroxamic acid derivative of gelatin by the method of Gallop, Seifter et al (14).

If the formation of alpha- and gamma-linkages occurred at the same rate, as they presumably would if the gamma-bond is defined at the aminoacyl-s-RNA level, then one would expect the incorporation into gamma-linkage to show a similar variation with time as the incorporation into alpha-linkage. The same result would be expected if an intramolecular rearrangement took place in a short enough time after the incorporation of glutamic acid into the alpha-linkage, such that the delay period could not be determined. If, however, intramolecular rearrangement occurs and there is a significant delay between incorporation into alpha-linked glutamic acid and its rearrangement into the gamma-linked form, then one might expect that the alpha-linked glutamyl units would contain radioactivity before activity appeared in the gamma-linked units. This is a possible explanation for the presence of activity in the alpha,gamma-diaminobutyric acid and not in the succinic semialdehyde in experiment III.

Further experimentation will be focused on the study of the rate of incorporation of radioactivity into the





alpha- and gamma-linkages, in order to gain some insight into the nature of the enzyme system responsible for the gamma-bond formation. If it is shown that the bond is created by an intramolecular rearrangement, then I believe that this would be the first instance in which a peptide bond was demonstrated to be created in this manner.



Summary

- 1) It was found that a cell-free system from 7-day old chick embryos would incorporate  $C^{14}$ -glutamic acid into the gelatin-rich hot-TCA-extractable protein. After separation of macromolecular from small molecular components, some of the activity was recovered in the macromolecular fraction. The amount of incorporation in this system was small.
- 2) Commercial bovine Achilles tendon collagen was converted to the polyhydroxamic acid-DNP derivative and submitted to Lossen rearrangement. Analysis of the hydrolysis products revealed the presence of succinic semialdehyde (the gamma-glutamyl-linkage rearrangement product) in both the rearranged gelatin and in an unrearranged control. The rearranged gelatin yielded 5.3 times as much succinic semialdehyde as it would be calculated to yield on the basis of the endogenous amount recovered from the unrearranged control. This collagen appears to contain at least 19  $\mu$ moles of gamma-glutamyl residues per 612 total glutamyl residues or per 676 residues of hydroxyproline. As discussed in the text, this is a minimal figure.
- 3) A whole-cell system of 20 14-day old chick embryos was incubated with  $C^{14}$ -glutamic acid. After conversion



to the polyhydroxamate-DNP derivative, the hot-TCA-extractable protein was submitted to Lossen rearrangement. Succinic semialdehyde was recovered and found to contain no radioactivity, whereas the alpha,gamma-diaminobutyric acid (alpha-linkage rearrangement product) which was recovered contained a small amount of radioactivity.

4) The collagen-containing hot-TCA-extractable protein was extracted from 20 14-day embryos. After formation of the polyhydroxamate derivative, one half of the solution was rearranged and the other was used as an unrearranged control. Analysis of the hydrolysis products revealed that both alpha- and gamma-linkage rearrangement products were present in the rearranged gelatin hydrolysate but not in that of the control. Quantitative analysis revealed that there was at least one gamma-linked glutamic acid residue for every nineteen hydroxyproline residues in the chick embryo collagen-containing hot TCA-extract.

5) A discussion of the possible mechanisms of gamma-bond formation was presented.



Bibliography

1. Quastel, J.H., Stewart, C.P., and Tunnicliffe, H.E.,  
Biochem. J. 17, 586 (1923).
2. Dakin, H.D., J. Biol. Chem. I, 171 (1905).
3. Hopkins, F.G., Biochem. J. 15, 286 (1921).
4. Kendall, E.C., Mason, H.L., and McKenzie, B.F., J.  
Biol. Chem. 87, 55 (1930).
5. Harrington, C.R., and Mead, T.H., Biochem. J. 29,  
1602 (1935).
6. Ivanovics, G., and Bruckner, V., Zeitschrift für  
Immunitätsforschung und Experimentelle Therapie. 90,  
304 (1937).
7. Bovarnick, M., J. Biol. Chem. 145, 415 (1942).
8. Hanby, W.E., and Rydon, H.N., Biochem. J., 40, 297 (1946).
9. Kovacs, J., and Bruckner, V., J. Chem. Soc. 4255 (1952).
10. Kovacs, J., Bruckner, V., and Kovacs, K., J. Chem.  
Soc., 145 (1953).
11. Bruckner, V., Kovacs, J., and Nagy, H., J. Chem. Soc.,  
149 (1953).
12. Waley, S.G., J. Chem. Soc., 517 (1955).
13. Chibnall, A.C., Rees, M.W., and Richards, F.M., Biochem.  
J. 68, 246 (1958).



Handwritten title or header at the top of the page.

First main section of handwritten text, consisting of several lines of cursive script.

Second main section of handwritten text, continuing the narrative or list.

Third main section of handwritten text, possibly concluding the document.

Final handwritten notes or a signature at the bottom of the page.

14. Gallop, P.M., Seifter, S., Lukin, M., and Meilman, E., J. Biol. Chem. 235, 2619 (1960).
15. Franzblau, C., "Unusual Linkages in Collagen", Ph.D. Thesis, Sue Golding Graduate Division of Medical Sciences, Albert Einstein College of Medicine, Yeshiva University, Obtained from University Microfilms, Inc., Ann Arbor, Michigan.
16. Gallop, P.M., Seifter, S., and Meilman, E., Nature 183, 1659 (1959).
17. Sheehan, J.C., and Hlavaka, J.J., J. Org. Chem. 21, 439 (1956).
18. Einbänder, J., and Schubert, M., J. Biol. Chem. 188, 335 (1951).
19. Littlefield, J.W., and Keller, E.B., J. Biol. Chem. 224, 13 (1957).
20. Blatt, A.H., in "Organic Synthesis" 7, 67 (1927).
21. Hendler, R.W., and Anfinson, C.B., J. Biol. Chem. 209, 55 (1954).
22. Peterkofsky, B., and Udenfriend, S., Biochem. and Biophys. Res. Comm. 6, 184 (1961).
23. Hirs, C.H.W., Moore, S., and Stein, W.H., J. Biol. Chem. 219, 623 (1956).
24. Moore, S., and Stein, W.H., J. Biol. Chem. 176, 367 (1948).

*[The page contains extremely faint, illegible text, likely bleed-through from the reverse side of the document. The text is arranged in several paragraphs and is difficult to decipher.]*

25. Seifter, S., Gallop, P.M., Michaels, S., and Meilman, E., J. Biol. Chem. 235, 2613 (1960).
26. Neuman, R.E., and Logan, M.A., J. Biol. Chem. 184, 299 (1950).
27. Leach, A.A., Biochem. J. 74, 70 (1960).
28. Chinard, F.P., J. Biol. Chem. 199, 91 (1952).
29. Waelsch, H. in "Methods in Enzymology"; vol. III (S.P. Colowick and N.O. Kaplan, ed.) p. 573, Academic Press, New York (1957).
30. Lukens, L., Personal Communication of unpublished material.
31. Ochoa, S., in "Informational Macromolecules", (H.J. Vogel, V. Bryson, and J.O. Lampen, ed.) p. 4, Academic Press, New York (1963).

1. Introduction

2. Methodology

3. Results

4. Discussion

5. Conclusion

6. References

7. Appendix

8. Bibliography

9. Index

10. Glossary

11. Acknowledgements

12. Author Biographies

13. Contact Information

14. Disclaimer

15. Copyright

16. Privacy Policy

17. Terms of Service

18. About Us

19. Press Release

20. Media Kit

21. Investor Relations

22. Sustainability

23. Diversity & Inclusion

24. Community Engagement

25. Risk Management

26. Governance

27. Compliance

28. Security

29. Data Protection

30. Environmental Impact

31. Social Responsibility

32. Human Resources

33. Finance

34. Operations

35. Technology

36. Marketing

37. Sales

38. Customer Support

39. Quality Assurance

40. Project Management

41. Procurement

42. Logistics

43. Manufacturing

44. Distribution

45. Retail

46. Wholesale

47. E-commerce

48. Mobile

49. Cloud

50. AI/ML

51. Blockchain

52. IoT

53. AR/VR

54. Big Data

55. Analytics

56. CRM

57. ERP

58. HRM

59. SCM

60. Supply Chain

61. Inventory Management

62. Warehouse Management

63. Transportation Management

64. Fleet Management

65. Risk Assessment

66. Insurance

67. Legal

68. Tax

69. Accounting

70. Finance

71. Investment

72. Banking

73. Finance

74. Insurance

75. Real Estate

76. Healthcare

77. Education

78. Government

79. Non-Profit

80. Media

81. Entertainment

82. Sports

83. Hospitality

84. Retail

85. Food & Beverage

86. Fashion

87. Beauty

88. Pharmaceuticals

89. Biotech

90. Aerospace

91. Defense

92. Energy

93. Utilities

94. Telecommunications

95. Media

96. Entertainment

97. Sports

98. Hospitality

99. Retail

100. Food & Beverage

101. Fashion

102. Beauty

103. Pharmaceuticals

104. Biotech

105. Aerospace

106. Defense

107. Energy

108. Utilities

109. Telecommunications

110. Media

111. Entertainment

112. Sports

113. Hospitality

114. Retail

115. Food & Beverage

116. Fashion

117. Beauty

118. Pharmaceuticals

119. Biotech

120. Aerospace

121. Defense

122. Energy

123. Utilities

124. Telecommunications

125. Media

126. Entertainment

127. Sports

128. Hospitality

129. Retail

130. Food & Beverage

131. Fashion

132. Beauty

133. Pharmaceuticals

134. Biotech

135. Aerospace

136. Defense

137. Energy

138. Utilities

139. Telecommunications

140. Media

141. Entertainment

142. Sports

143. Hospitality

144. Retail

145. Food & Beverage

146. Fashion

147. Beauty

148. Pharmaceuticals

149. Biotech

150. Aerospace

151. Defense

152. Energy

153. Utilities

154. Telecommunications

155. Media

156. Entertainment

157. Sports

158. Hospitality

159. Retail

160. Food & Beverage

161. Fashion

162. Beauty

163. Pharmaceuticals

164. Biotech

165. Aerospace

166. Defense

167. Energy

168. Utilities

169. Telecommunications

170. Media

171. Entertainment

172. Sports

173. Hospitality

174. Retail

175. Food & Beverage

176. Fashion

177. Beauty

178. Pharmaceuticals

179. Biotech

180. Aerospace

181. Defense

182. Energy

183. Utilities

184. Telecommunications

185. Media

186. Entertainment

187. Sports

188. Hospitality

189. Retail

190. Food & Beverage

191. Fashion

192. Beauty

193. Pharmaceuticals

194. Biotech

195. Aerospace

196. Defense

197. Energy

198. Utilities

199. Telecommunications

200. Media

201. Entertainment

202. Sports

203. Hospitality

204. Retail

205. Food & Beverage

206. Fashion

207. Beauty

208. Pharmaceuticals

209. Biotech

210. Aerospace

211. Defense

212. Energy

213. Utilities

214. Telecommunications

215. Media

216. Entertainment

217. Sports

218. Hospitality

219. Retail

220. Food & Beverage

221. Fashion

222. Beauty

223. Pharmaceuticals

224. Biotech

225. Aerospace

226. Defense

227. Energy

228. Utilities

229. Telecommunications

230. Media

231. Entertainment

232. Sports

233. Hospitality

234. Retail

235. Food & Beverage

236. Fashion

237. Beauty

238. Pharmaceuticals

239. Biotech

240. Aerospace

241. Defense

242. Energy

243. Utilities

244. Telecommunications

245. Media

246. Entertainment

247. Sports

248. Hospitality

249. Retail

250. Food & Beverage

251. Fashion

252. Beauty

253. Pharmaceuticals

254. Biotech

255. Aerospace

256. Defense

257. Energy

258. Utilities

259. Telecommunications

260. Media

261. Entertainment

262. Sports

263. Hospitality

264. Retail

265. Food & Beverage

266. Fashion

267. Beauty

268. Pharmaceuticals

269. Biotech

270. Aerospace

271. Defense

272. Energy

273. Utilities

274. Telecommunications

275. Media

276. Entertainment

277. Sports

278. Hospitality

279. Retail

280. Food & Beverage

281. Fashion

282. Beauty

283. Pharmaceuticals

284. Biotech

285. Aerospace

286. Defense

287. Energy

288. Utilities

289. Telecommunications

290. Media

291. Entertainment

292. Sports

293. Hospitality

294. Retail

295. Food & Beverage

296. Fashion

297. Beauty

298. Pharmaceuticals

299. Biotech

300. Aerospace

301. Defense

302. Energy

303. Utilities

304. Telecommunications

305. Media

306. Entertainment

307. Sports

308. Hospitality

309. Retail

310. Food & Beverage

311. Fashion

312. Beauty

313. Pharmaceuticals

314. Biotech

315. Aerospace

316. Defense

317. Energy

318. Utilities

319. Telecommunications

320. Media

321. Entertainment

322. Sports

323. Hospitality

324. Retail

325. Food & Beverage

326. Fashion

327. Beauty

328. Pharmaceuticals

329. Biotech

330. Aerospace

331. Defense

332. Energy

333. Utilities

334. Telecommunications

335. Media

336. Entertainment

337. Sports

338. Hospitality

339. Retail

340. Food & Beverage

341. Fashion

342. Beauty

343. Pharmaceuticals

344. Biotech

345. Aerospace

346. Defense

347. Energy

348. Utilities

349. Telecommunications

350. Media

351. Entertainment

352. Sports

353. Hospitality

354. Retail

355. Food & Beverage

356. Fashion

357. Beauty

358. Pharmaceuticals

359. Biotech

360. Aerospace

361. Defense

362. Energy

363. Utilities

364. Telecommunications

365. Media

366. Entertainment

367. Sports

368. Hospitality

369. Retail

370. Food & Beverage

371. Fashion

372. Beauty

373. Pharmaceuticals

374. Biotech

375. Aerospace

376. Defense

377. Energy

378. Utilities

379. Telecommunications

380. Media

381. Entertainment

382. Sports

383. Hospitality

384. Retail

385. Food & Beverage

386. Fashion

387. Beauty

388. Pharmaceuticals

389. Biotech

390. Aerospace

391. Defense

392. Energy

393. Utilities

394. Telecommunications

395. Media

396. Entertainment

397. Sports

398. Hospitality

399. Retail

400. Food & Beverage

401. Fashion

402. Beauty

403. Pharmaceuticals

404. Biotech

405. Aerospace

406. Defense

407. Energy

408. Utilities

409. Telecommunications

410. Media

411. Entertainment

412. Sports

413. Hospitality

414. Retail

415. Food & Beverage

416. Fashion

417. Beauty

418. Pharmaceuticals

419. Biotech

420. Aerospace

421. Defense

422. Energy

423. Utilities

424. Telecommunications

425. Media

426. Entertainment

427. Sports

428. Hospitality

429. Retail

430. Food & Beverage

431. Fashion

432. Beauty

433. Pharmaceuticals

434. Biotech

435. Aerospace

436. Defense

437. Energy

438. Utilities

439. Telecommunications

440. Media

441. Entertainment

442. Sports

443. Hospitality

444. Retail

445. Food & Beverage

446. Fashion

447. Beauty

448. Pharmaceuticals

449. Biotech

450. Aerospace

451. Defense

452. Energy

453. Utilities

454. Telecommunications

455. Media

456. Entertainment

457. Sports

458. Hospitality

459. Retail

460. Food & Beverage

461. Fashion

462. Beauty

463. Pharmaceuticals

464. Biotech

465. Aerospace

466. Defense

467. Energy

468. Utilities

469. Telecommunications

470. Media

471. Entertainment

472. Sports

473. Hospitality

474. Retail

475. Food & Beverage

476. Fashion

477. Beauty

478. Pharmaceuticals

479. Biotech

480. Aerospace

481. Defense

482. Energy

483. Utilities

484. Telecommunications

485. Media

486. Entertainment

487. Sports

488. Hospitality

489. Retail

490. Food & Beverage

491. Fashion

492. Beauty

493. Pharmaceuticals

494. Biotech

495. Aerospace

496. Defense

497. Energy

498. Utilities

499. Telecommunications

500. Media

501. Entertainment

502. Sports

503. Hospitality

504. Retail

505. Food & Beverage

506. Fashion

507. Beauty

508. Pharmaceuticals

509. Biotech

510. Aerospace

511. Defense

512. Energy

513. Utilities

514. Telecommunications

515. Media

516. Entertainment

517. Sports

518. Hospitality

519. Retail

520. Food & Beverage

521. Fashion

522. Beauty

523. Pharmaceuticals

524. Biotech

525. Aerospace

526. Defense

527. Energy

528. Utilities

529. Telecommunications

530. Media

531. Entertainment

532. Sports

533. Hospitality

534. Retail

535. Food & Beverage

536. Fashion

537. Beauty

538. Pharmaceuticals

539. Biotech

540. Aerospace

541. Defense

542. Energy

543. Utilities

544. Telecommunications

545. Media

546. Entertainment

547. Sports

548. Hospitality

549. Retail

550. Food & Beverage

551. Fashion

552. Beauty

553. Pharmaceuticals

554. Biotech

555. Aerospace

556. Defense

557. Energy

558. Utilities

559. Telecommunications

560. Media

561. Entertainment

562. Sports

563. Hospitality

564. Retail

565. Food & Beverage

566. Fashion

567. Beauty

568. Pharmaceuticals

569. Biotech

570. Aerospace

571. Defense

572. Energy

573. Utilities

574. Telecommunications

575. Media

576. Entertainment

577. Sports

578. Hospitality

579. Retail

580. Food & Beverage

581. Fashion

582. Beauty

583. Pharmaceuticals

584. Biotech

585. Aerospace

586. Defense

587. Energy

588. Utilities

589. Telecommunications

590. Media

591. Entertainment

592. Sports

593. Hospitality

594. Retail

595. Food & Beverage

596. Fashion

597. Beauty

598. Pharmaceuticals

599. Biotech

600. Aerospace

601. Defense

602. Energy

603. Utilities

604. Telecommunications

605. Media

606. Entertainment

607. Sports

608. Hospitality

609. Retail

610. Food & Beverage

611. Fashion

612. Beauty

613. Pharmaceuticals

614. Biotech

615. Aerospace

616. Defense

617. Energy

618. Utilities

619. Telecommunications

620. Media

621. Entertainment

622. Sports

623. Hospitality

624. Retail

625. Food & Beverage

626. Fashion

627. Beauty

628. Pharmaceuticals

629. Biotech

630. Aerospace

631. Defense

632. Energy

633. Utilities

634. Telecommunications

635. Media

636. Entertainment

637. Sports

638. Hospitality

639. Retail

640. Food & Beverage

641. Fashion

642. Beauty

643. Pharmaceuticals

644. Biotech

645. Aerospace

646. Defense

647. Energy

648. Utilities

649. Telecommunications

650. Media

651. Entertainment

652. Sports

653. Hospitality

654. Retail

655. Food & Beverage

656. Fashion

657. Beauty

658. Pharmaceuticals

659. Biotech

660. Aerospace

661. Defense

662. Energy

663. Utilities

664. Telecommunications

665. Media

666. Entertainment

667. Sports

668. Hospitality

669. Retail

670. Food & Beverage

671. Fashion

672. Beauty

673. Pharmaceuticals

674. Biotech

675. Aerospace

676. Defense

677. Energy

678. Utilities

679. Telecommunications

680. Media

681. Entertainment

682. Sports

683. Hospitality

684. Retail

685. Food & Beverage

686. Fashion

687. Beauty

688. Pharmaceuticals

689. Biotech

690. Aerospace

691. Defense

692. Energy

693. Utilities

694. Telecommunications

695. Media

696. Entertainment

697. Sports

698. Hospitality

699. Retail

700. Food & Beverage

701. Fashion

702. Beauty

703. Pharmaceuticals

704. Biotech

705. Aerospace

706. Defense

707. Energy

708. Utilities

709. Telecommunications

710. Media

711. Entertainment

712. Sports

713. Hospitality

714. Retail

715. Food & Beverage

716. Fashion

717. Beauty

718. Pharmaceuticals

719. Biotech

720. Aerospace

721. Defense

722. Energy

723. Utilities

724. Telecommunications

725. Media

726. Entertainment

727. Sports

728. Hospitality

729. Retail

730. Food & Beverage

731. Fashion

732. Beauty

733. Pharmaceuticals

734. Biotech

735. Aerospace

736. Defense

737. Energy

738. Utilities

739. Telecommunications

740. Media

741. Entertainment

742. Sports

743. Hospitality

744. Retail

745. Food & Beverage

746. Fashion

747. Beauty

748. Pharmaceuticals

749. Biotech

750. Aerospace

751. Defense

752. Energy

753. Utilities

754. Telecommunications

755. Media

756. Entertainment

757. Sports

758. Hospitality

759. Retail

760. Food & Beverage

761. Fashion

762. Beauty

763. Pharmaceuticals

764. Biotech

765. Aerospace

766. Defense

767. Energy

768. Utilities

769. Telecommunications

770. Media

771. Entertainment

772. Sports

773. Hospitality

774. Retail

775. Food & Beverage

776. Fashion

777. Beauty

778. Pharmaceuticals

779. Biotech

780. Aerospace

781. Defense

782. Energy

783. Utilities

784. Telecommunications

785. Media

786. Entertainment

787. Sports

788. Hospitality

789. Retail

790. Food & Beverage

791. Fashion

792. Beauty

793. Pharmaceuticals

794. Biotech

795. Aerospace

796. Defense

797. Energy

798. Utilities

799. Telecommunications

800. Media

801. Entertainment

802. Sports

803. Hospitality

804. Retail

805. Food & Beverage

806. Fashion

807. Beauty

808. Pharmaceuticals

809. Biotech

810. Aerospace

811. Defense

812. Energy

813. Utilities

814. Telecommunications

815. Media

816. Entertainment

817. Sports

818. Hospitality

819. Retail

820. Food & Beverage

821. Fashion

822. Beauty

823. Pharmaceuticals

824. Biotech

825. Aerospace

826. Defense

827. Energy

828. Utilities

829. Telecommunications

830. Media

831. Entertainment

832. Sports

833. Hospitality

834. Retail

835. Food & Beverage

836. Fashion

837. Beauty

838. Pharmaceuticals

839. Biotech

840. Aerospace

841. Defense

842. Energy

843. Utilities

844. Telecommunications

845. Media

846. Entertainment

847. Sports

848. Hospitality

849. Retail

850. Food & Beverage

851. Fashion

852. Beauty

853. Pharmaceuticals

854. Biotech

855. Aerospace

856. Defense

857. Energy

858. Utilities

859. Telecommunications

860. Media

861. Entertainment

862. Sports

863. Hospitality

864. Retail

865. Food & Beverage

866. Fashion

867. Beauty

868. Pharmaceuticals

869. Biotech

870. Aerospace

871. Defense

872. Energy

873. Utilities

874. Telecommunications

875. Media

876. Entertainment

877. Sports

878. Hospitality

879. Retail

880. Food & Beverage

881. Fashion

882. Beauty

883. Pharmaceuticals

884. Biotech

885. Aerospace

886. Defense

887. Energy

888. Utilities

889. Telecommunications

890. Media

891. Entertainment

892. Sports

893. Hospitality

894. Retail

895. Food & Beverage

896. Fashion

897. Beauty

898. Pharmaceuticals

899. Biotech

900. Aerospace

901. Defense

902. Energy

903. Utilities

904. Telecommunications

905. Media

906. Entertainment

907. Sports

908. Hospitality

909. Retail

910. Food & Beverage

911. Fashion

912. Beauty

913. Pharmaceuticals

914. Biotech

915. Aerospace

916. Defense

917. Energy

918. Utilities

919. Telecommunications

920. Media

921. Entertainment

922. Sports

923. Hospitality

924. Retail

925. Food & Beverage

926. Fashion

927. Beauty

928. Pharmaceuticals

929. Biotech

930. Aerospace

931. Defense

932. Energy

933. Utilities

934. Telecommunications

935. Media

936. Entertainment

937. Sports

938. Hospitality

939. Retail

940. Food & Beverage

941. Fashion

942. Beauty

943. Pharmaceuticals

944. Biotech

945. Aerospace

946. Defense

947. Energy

948. Utilities

949. Telecommunications

950. Media

951. Entertainment

952. Sports

953. Hospitality

954. Retail

955. Food & Beverage

956. Fashion

957. Beauty

958. Pharmaceuticals

959. Biotech

960. Aerospace

961. Defense

962. Energy

963. Utilities

964. Telecommunications

965. Media

966. Entertainment

967. Sports

968. Hospitality

969. Retail

970. Food & Beverage

971. Fashion

972. Beauty

973. Pharmaceuticals

974. Biotech

975. Aerospace

976. Defense

977. Energy

978. Utilities

979. Telecommunications

980. Media

981. Entertainment

982. Sports

983. Hospitality

984. Retail

985. Food & Beverage

986. Fashion

987. Beauty

988. Pharmaceuticals

989. Biotech

990. Aerospace

991. Defense

992. Energy

993. Utilities

994. Telecommunications

995. Media

996. Entertainment

997. Sports

998. Hospitality

999. Retail

1000. Food & Beverage





YALE MEDICAL LIBRARY

Manuscript Theses

Unpublished theses submitted for the Master's and Doctor's degrees and deposited in the Yale Medical Library are to be used only with due regard to the rights of the authors. Bibliographical references may be noted, but passages must not be copied without permission of the authors, and without proper credit being given in subsequent written or published work.

This thesis by \_\_\_\_\_ has been used by the following persons, whose signatures attest their acceptance of the above restrictions.

---

---

NAME AND ADDRESS

DATE



