

4. Page LK, Anchor C, Luy E, Kron S, Larson G, Madsen L, et al. Backyard raccoon latrines and risk for *Baylisascaris procyonis* transmission to humans. *Emerg Infect Dis*. 2009;15:1530–1. <http://dx.doi.org/10.3201/eid1509.090128>
5. Kelly TG, Madhavan VL, Peters JM, Kazacos KR, Silvera VM. Spinal cord involvement in a child with raccoon roundworm (*Baylisascaris procyonis*) meningoencephalitis. *Pediatr Radiol*. 2012;42:369–73. <http://dx.doi.org/10.1007/s00247-011-2151-y>
6. Wise ME, Sorvillo FJ, Shafir SC, Ash LR, Berlin OG. Severe and fatal central nervous system disease in humans caused by *Baylisascaris procyonis*, the common roundworm of raccoons: a review of current literature. *Microbes Infect*. 2005;7:317–23. <http://dx.doi.org/10.1016/j.micinf.2004.12.005>
7. Blizzard EL, Yabsley MJ, Beck MF, Harsch S. Geographic expansion of *Baylisascaris procyonis* roundworms, Florida, USA. *Emerg Infect Dis*. 2010;16:1803–4. <http://dx.doi.org/10.3201/eid1611.100549>
8. Miyashita M. Prevalence of *Baylisascaris procyonis* in raccoons in Japan and experimental infections of the worm in laboratory animals. *Journal of Urban Living and Health Association*. 1993;37:137–51.
9. Xie Y, Zhang Z, Niu L, Wang Q, Wang C, Lan J, et al. The mitochondrial genome of *Baylisascaris procyonis*. *PLoS ONE*. 2011;6:e27066. <http://dx.doi.org/10.1371/journal.pone.0027066>
10. Sorvillo F, Ash LR, Berlin OGW, Tatabe J, Degiorgio C, Morse SA. *Baylisascaris procyonis*: an emerging helminthic zoonosis. *Emerg Infect Dis*. 2002;8:355–9. <http://dx.doi.org/10.3201/eid0804.010273>

Address for correspondence: Guangyou Yang, Department of Parasitology, College of Veterinary Medicine, Sichuan Agricultural University, 46 Xinkang Rd, Ya'an, Sichuan, 625014, People's Republic of China; email: guangyouyang@hotmail.com

Novel Divergent Rhabdovirus in Feces of Red Fox, Spain

To the Editor: Rhabdoviruses (family *Rhabdoviridae*) are enveloped single-stranded negative-sense RNA viruses belonging to the Mononegavirales order. The International Committee on Taxonomy of Viruses recognizes 11 genera (*Cytorhabdovirus*, *Ephemerovirus*, *Lyssavirus*, *No-virhabdovirus*, *Nucleorhabdovirus*, *Perhabdovirus*, *Sigmarivirus*, *Sprivivirus*, *Tibrovirus*, *Tupavivirus*, *Vesiculovirus*) (1). In addition, many recently described rhabdoviruses remain unassigned. Rhabdoviruses contain 5 major genes, encoding for nucleoprotein (N), phosphoprotein (P), matrix (M), glycoprotein (G), and RNA-dependent RNA polymerase (L). The *Rhabdoviridae* family includes pathogens of various animal species, humans, and plants. Viruses of the genus *Lys-savirus* are the most relevant to public health because they can cause rabies. Bats are the driving force within this genus; foxes and various other species of wild carnivores also can be infected with lyssaviruses and transmit them to humans and dogs (2).

During a viral metagenomic survey, conducted as described previously (3), of fecal samples collected from 4 red foxes (*Vulpes vulpes*) that were found dead in Álava, Basque Country, Spain, we identified the complete coding sequence and the partial leader and trailer sequence of a novel rhabdovirus, tentatively called red fox fecal rhabdovirus (RFFRV; 15,541 nt, GenBank accession no. KF823814; online Technical Appendix, <http://wwwnc.cdc.gov/EID/article/20/12/14-0236-Techapp1.pdf>) by mapping 8,287 of the 56,519 sequence reads in the sample of a red fox. A proportion of obtained reads contained sequences that were ≥99% identical to mitochondrial DNA of *V. vulpes*, which confirmed

that the sample was collected from a red fox.

The obtained sequence of RFFRV was partially confirmed by specific primers and Sanger sequencing of PCR amplicons. Five major and 3 minor open reading frames (ORFs) were identified that had a genome organization similar to that of other rhabdoviruses (Figure, panel A). No significant hits were obtained by BLAST analysis (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>) of N, P, M, and G nucleotide and amino acid sequences, which was reported previously for novel divergent rhabdoviruses (4).

Predicted N, P, and M genes of RFFRV consist of 1,629, 2,490, and 813 nt, respectively, encoding for 543, 830, and 271 aa (online Technical Appendix Table 1). In addition to the absence of significant hits observed by BLAST analysis, no significant sequence homology was observed with known rhabdovirus proteins in pairwise alignments. Furthermore, no conserved motifs were detected in N, P, and M genes of RFFRV that are commonly observed in rhabdoviruses. However, intergenic regions between all major ORFs contained relatively conserved motifs that could be transcription termination/polyadenylation sequences (A/U) CU₇, similar to other rhabdoviruses (5). Adjacent to this termination signal was a stretch of conserved nucleotides that might function as a transcription initiation signal (online Technical Appendix Table 1).

The amino acid sequence of the G protein consisted of 669 aa and contained an N terminal signal peptide (1-MYHLIVLLVMLGQRA-VA-17), a noncytoplasmic domain (aa 18–646), a transmembrane domain (647-ITALMPLLSLAVVVGI-IMCC-667), and a cytoplasmic tail of 2 aa, similar to other rhabdovirus G proteins as predicted by using Phobius and TMHMM (<http://www.cbs.dtu.dk/services/TMHMM>) (6,7). We predicted 3 potential glycosylation sites in the ectodomain at positions 38–40



(NKT), 554–556 (NAS), and 592–594 (NIS) using NetNGlyc 1.0 (<http://www.cbs.dtu.dk/services/NetNGlyc>).

Between the G and L genes, a complex intergenic region was present that contained 3 ORFs of 246 nt (7,413–7,658 aa), 231 nt (7,716–7,946 aa), and 459 nt (7,893–8,355 aa), of which 2 were overlapping frames (U1–3). Additional ORFs between G and L genes were detected previously in other rhabdoviruses (8,9). We detected transmembrane domains in the amino acid sequences of all 3 additional ORFs, suggesting they might act as viroporin (8,9).

The L gene of RFFRV contained 6,591 nt (2,197 aa). We detected several conserved domains and motifs, including RNA-dependent RNA polymerase, mRNA-capping region, mRNA capping enzyme, and virus-capping methyltransferase. Alignment of the deduced amino acid sequence of the L gene with the L gene of various other viruses belonging to the Mononegavirales order by using MAFFT version 7 (<http://mafft.cbrc.jp/alignment/software/>) and subsequent phylogenetic reconstruction by using a maximum-likelihood tree (WAG+F+I+G model with 100 bootstrap replicates in MEGA5 [<http://www.megasoftware.net>]) suggested that this virus belongs to a novel genus of the *Rhabdoviridae* family. In addition, pairwise identities of the deduced amino acid sequence of the L gene of RFFRV with that of other rhabdoviruses of the *Rhabdoviridae* family were only $\leq 35\%$ (online Technical Appendix Table 2).

Because the fox was found dead and no tissue samples were collected, whether RFFRV played a role in the animal's death is unknown. In addition, multiple attempts to isolate this virus on various cell lines of eukaryotes (Vero E6, MDCK, CRFK, N2a, and BHK cells, primary fox kidney cells) failed because of the absence of cytopathic effects and viral replication by quantitative reverse transcription PCR.

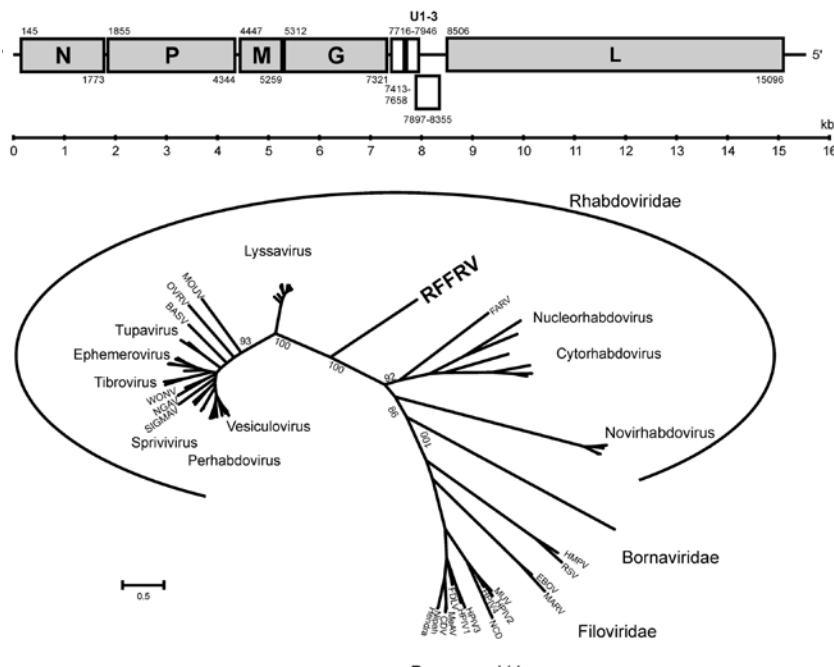


Figure. Genome organization and phylogenetic analysis of RFFRV. A) Genome organization of RFFRV. Indicated are the locations of the major ORFs (including the positions of the first and last nucleotide) and 3 theoretical minor ORFs between the G and L genes. B) Phylogenetic maximum-likelihood tree using the WAG+F+I+G model and 100 bootstrap replicates in MEGA5 (<http://www.megasoftware.net>) of the deduced amino acid sequence of the L genes of various viruses of the order Mononegavirales. G, glycoprotein; L, RNA-dependent RNA polymerase; M, matrix; N, nucleoprotein; ORF, open reading frame; P, phosphoprotein; RFFRV, RFFRV, red fox fecal rhabdovirus. Only bootstrap values in the close proximity of the branch of the RFFRV are indicated. Scale bar indicates nucleotide substitutions per site. Viruses and GenBank accession numbers are shown in the expanded figure legend online (<http://wwwnc.cdc.gov/EID/article/20/12/14-0236-E1.htm>).

despite a high number of reads in the original sample. The fox might have acquired the virus through spillover from a small prey, such as a bat, and additional studies are required to elucidate the prevalence, original host, and pathogenic potential of this novel virus.

Acknowledgments

We thank all researchers and institutions for their invaluable help during sampling and for providing the specimens used in this study, especially Patricia Lizarraga, Laura Elorza, Ricardo Gutierrez, and Luis Javier Chueca.

This work was funded by the European Community's Seventh Framework Program (FP7/2007–2013) under the project "European Management Platform for Emerging and Re-emerging Infectious

Disease Entities” European Community grant agreement no. 223498; the Virgo Consortium; and the Niedersachsen-Research Network on Neuroinfectiology of the Ministry of Science and Culture of Lower Saxony, Germany. In addition, this research was funded partially by the Basque Government through the research group on “Systematics, Biogeography and Population Dynamics” (ref. IT317-10; GIC10/76).

A.R.-G. holds a postdoctoral fellowship awarded by the Department of Education, Universities and Research of the Basque Government (ref. DKR-2012-64) and was awarded a short-visit research grant from the ConGenomics Research networking program of the European Science Foundation to visit the Department of Viroscience, Erasmus Medical Centre and develop the current research project.

Rogier Bodewes,¹
Aritz Ruiz-Gonzalez,¹
Anita C. Schürch,
Albert D.M.E. Osterhaus,
and Saskia L. Smits

Author affiliations: Erasmus Medical Centre, Rotterdam, the Netherlands (R. Bodewes, A.C. Schürch, A.D.M.E. Osterhaus, S.L. Smits); University of the Basque Country, Vitoria-Gasteiz, Spain (A. Ruiz-Gonzalez); National Institute for Environmental Protection and Research, Ozzano dell'Emilia, Italy (A. Ruiz-Gonzalez); and Viroclinics Biosciences, Rotterdam (A.D.M.E. Osterhaus, S.L. Smits)

DOI: <http://dx.doi.org/10.3201/eid2012.140236>

References

- International Committee on Taxonomy of Viruses. Virus taxonomy: 2013 release [cited 2014 Feb 9]. <http://www.ictvonline.org/virusTaxonomy.asp>
- Matha IS, Salunke SR. Immunogenicity of purified Vero cell rabies vaccine used in the treatment of fox-bite victims in India. Clin Infect Dis. 2005;40:611–3. <http://dx.doi.org/10.1086/427700>
- Bodewes R, Ruiz-Gonzalez A, Schapendonk CM, van den Brand JM, Osterhaus AD, Smits SL. Viral metagenomic analysis of feces of wild small carnivores. Virol J. 2014;11:89. <http://dx.doi.org/10.1186/1743-422X-11-89>
- Palacios G, Forrester NL, Savji N, Travassos da Rosa AP, Guzman H, Detoy K, et al. Characterization of Farmington virus, a novel virus from birds that is distantly related to members of the family *Rhabdoviridae*. Virol J. 2013;10:219. <http://dx.doi.org/10.1186/1743-422X-10-219>
- Albertini AA, Ruigrok RW, Blondel D. Rabies virus transcription and replication. Adv Virus Res. 2011;79:1–22. <http://dx.doi.org/10.1016/B978-0-12-387040-7.00001-9>
- Coll JM. The glycoprotein G of rhabdoviruses. Arch Virol. 1995;140:827–51. <http://dx.doi.org/10.1007/BF01314961>
- Käll L, Krogh A, Sonnhammer EL. A combined transmembrane topology and signal peptide prediction method. J Mol Biol. 2004;338:1027–36. <http://dx.doi.org/10.1016/j.jmb.2004.03.016>
- Gubala AJ, Proll DF, Barnard RT, Cowled CJ, Crameri SG, Hyatt AD, et al. Genomic characterisation of Wongabel virus reveals novel genes within the

- Rhabdoviridae*. Virology. 2008;376:13–23. <http://dx.doi.org/10.1016/j.virol.2008.03.004>
- McWilliam SM, Kongswan K, Cowley JA, Byrne KA, Walker PJ. Genome organization and transcription strategy in the complex GNS-L intergenic region of bovine ephemeral fever rhabdovirus. J Gen Virol. 1997;78:1309–17.

Address for correspondence: Rogier Bodewes, Erasmus Medical Centre, Department of Viroscience Dr. Molewaterplein 50, 3015GE Rotterdam, the Netherlands; email: r.bodewes@erasmusmc.nl

Ngari Virus in Goats during Rift Valley Fever Outbreak, Mauritania, 2010

To the Editor: Ngari virus (NRIV) is a single-stranded RNA virus belonging to the family *Bunyaviridae*, genus *Orthobunyavirus*. The genome comprises 3 segments, the small (S), medium (M), and large (L) segments, which encode the nucleocapsid (N) protein, the 2 glycoproteins Gn and Gc, and the RNA-dependent RNA-polymerase, respectively. Sequence analysis showed that NRIV is a reassortant between Bunyamwera virus (BUNV) and Batai virus (BATV), both from the genus *Orthobunyavirus*. S and L segments derived from BUNV, and the M segment derived from BATV (1,2). NRIV is more virulent than BUNV and BATV and is associated with hemorrhagic fever. NRIV was first isolated from *Aedes simpsoni* mosquitoes in 1979 and from humans in 1993, both in Senegal (3). During 1997 and 1998, humans were affected with hemorrhagic fever diseases in Kenya and Somalia that were caused by Rift Valley fever virus (RVFV) and by NRIV (2,4).

In 2010, during an ongoing RVFV outbreak in Mauritania, we collected 163 serum samples (62 from camels, 8 from cattle, and 93 from small ruminants) (5). RVFV RNA was isolated from serum samples as described previously (5). Further molecular testing of the samples was conducted by a SYBRGreen-based real-time reverse transcription PCR (RT-PCR) adapted from a conventional RT-PCR and based on generic primers (bun_group_forw 5'-CTGCTAA-CACCAGCAGTACTTTGAC-3' and bun_group_rev 5'-TGGAGGGTA-AGACCATCGTCAGGAAGTG-3') that target a 250-nt sequence of the S segment of Bunyamwera serogroup members (6). Real-time RT-PCR was performed in a CFX 96 real-time PCR system (Bio-Rad, Hercules, CA, USA) by using 5 µL RNA with a QuantiTect SYBR Green RT-PCR Kit (QIAGEN, Hilden Germany) in a final volume of 25 µL. Cycling conditions included RT at 50°C for 30 min and 95°C for 15 min, followed by amplification with 44 cycles of 95°C for 15 s, 55°C for 25 s, 72°C for 30 s, and 77°C for 5 s. A melting curve analysis was then performed starting with 95°C for 60 s, and a temperature gradient was conducted from 68°C to 94°C in increments of 0.2°C.

Of the 163 serum samples tested, 2 samples from goats resulted in a positive signal with cycle thresholds of 23 (sample 51) and 28 (sample 65), respectively. Both samples showed similar melting peaks at ≈78.2°C and shared the identical partial nucleotide sequence of the S segment. The sequence belongs to the Bunyamwera serogroup, but the short partial sequence was not sufficient for accurate virus determination and identification. For this reason, both serum samples were used to inoculate cell monolayers of Vero E6 cells that were assayed for virus replication. Only sample 51 displayed a cytopathic effect after 72 h and was further analyzed. We isolated the viral RNA from cell culture with TRIzol reagent

¹These authors contributed equally to this article.

Novel Divergent Rhabdovirus in Feces of Red Fox, Spain

Technical Appendix

Technical Appendix Table 1. Characteristics of RFFRV genes and intergenic sequences*

Gene	Length, nt	Length, aa	Conserved intergenic sequence between 2 ORFs†
N	1,629	542	<u>UAG</u> -60nt-ACAAAAAAACUAACCUCAGCUA <u>AUG</u>
P	2,490	829	<u>UAA</u> -75nt-UCAAAAAAAACUAACACAAGCUGCGAA <u>AUG</u>
M	813	270	<u>UAA</u> -23nt-ACAAAAAAACUAACCUAACUAGGUUA <u>AUG</u>
G	2,010	669	<u>UAA</u> -70nt-ACAAAAAAACUAACACAAGCUA <u>AUG</u>
U1	246	81	<u>UAA</u> -29nt-ACAAAAAAACUAACACUACCUUCGUGUU <u>AUG</u>
U2	231	76	NA
U3	459	152	<u>UAA</u> -116nt-ACAAAAAAACUAACACUACUGUGUUAUACAA <u>AUG</u>
L	6,591	2,196	NA

*NA, not applicable; ORF, open reading frame; RFFRV, red fox fecal rhabdovirus.

†Start and stopcodon of the flanking genes are underlined.

Technical Appendix Table 2. Pairwise amino acid identities between the L protein of RFFRV and other rhabdoviruses

RFFRV	LNYV	BEFV	RABV	IHNV	RYSV	PRV	SIGMAV	VSIV	TBIV	BASV	FARV	LBV	SHIBV	OZEV
RFFRV	27	31	34	23	27	33	30	32	31	30	29	35	35	35
LNYV NC_007642	27	25	30	23	35	25	25	25	25	25	33	30	30	30
BEFV NC_002526	31	25	45	24	24	60	57	60	57	52	25	46	46	45
RABV SRV9 AAT48626	34	30	45	25	28	46	44	46	44	40	29	86	87	88
IHNV L40883	23	23	24	25	22	24	24	23	23	20	27	25	26	25
RYSV NC_003746	27	35	24	28	22	25	24	25	23	24	30	28	29	28
PRV HM566195	33	25	60	46	24	25	60	74	54	50	26	47	47	47
SIGMAV Q410979	30	25	57	44	24	24	60	59	52	48	23	46	45	45
VSIV AAA48441	32	25	60	46	23	25	74	59	53	51	27	47	46	46
TBIV GQ294472	31	25	57	44	23	23	54	52	53	52	24	43	43	42
BASV JX297815	30	25	52	40	20	24	50	48	51	52	23	41	41	41
FARV HM627182	29	33	25	29	27	30	26	23	27	24	23	28	28	28
LBV JX901139	35	30	46	86	25	28	47	46	47	43	41	28	92	86
SHIBV ADD84511	35	30	46	87	26	29	47	45	46	43	41	28	92	87
OZEV FJ905105	35	30	45	88	25	28	47	45	46	42	41	28	86	87

Red Fox Fecal Rhabdovirus, Partial Genome (GenBank Accession No. KF823814)*

GGATATCAAGTCCACCAATACCTTATATGTGCATGATCATGCACATCACATTCTACG
 CATACTCCCAGGGAGTACACTAAAAAGACGTAACACAGACTTGAATTACGT
 CAAGTCTAAGTTAAATTGGTTAATTCAATGGATCACGATAACGAAAAGCCAAT
CTCGTACACTCAATAGCGGAAGTTCCTGATAATGTTGCCATTGGGAGCACCATT
CATTCAAGGTGAGCCCATCATCTATTGGAAAATCCGCTGCAACAGGAATTACGCG
GAAAGGGGGGCACAGAAAGACTGGACCAAAGACATGATCCGTGGAGTGAGAGTG
TTCCTGCCCGACTGATGCTAATCTGCTCAATCTCATAGCCGGGAAACCGAAGCC
CCTGAGCTGGAGAAGTACACCATCCAAGATCCGGAAAAGAAGGGCATCTGAAGAA
ATTGAAAGCAAGTGGAGTTGCGAATTGGGCAAACCTGCTGGTCGACTTGCAAA

GCAACACAGGGAACATCCCCAAAGGGAGATTCCCTACTACTCTGCTCTATTCTCGA
TAACTGCCATTAAGGGAGCCCCTGTCCTGGCCCTGCCATGAAAGACCTCGGGGACC
CTGTTATGTGAAAGCTCCCGATGACCTCCACCCACCTACAGGAGACATAGAATGGC
ATGGTGATAAAATTAGTGTGACGAGGCAGCCTACATAGGATATGGAGCATGGCTG
ATCATGCCTAGATTCACTATCAAAGCTGAATCCAAGAAAGATGAAATTGCAGCCAG
CAGCAAGGCATTGACACTCTTAGGCGGTTGCTACCTGAGATCACCAAGCCACAAGT
GTTGGTATCTGTGGTGACGCAGCTCAGATTGGCATATCACGGAACGCTGGTCCCGG
GTCTGCGTACCTGCTGCAGAAGTAGCAATGAGAAGGGCAATGAATATAGAGTATG
ACCTAAAGGCTGACAGAACGGAGTGCAAAGCCGGGAAACACTTCCAGGTTGTCAA
TTAAGAGTCCTGCAGGATATCCCCAATACGACTCAGGCTTGGGCTTGGCAA
GTTGGGCTAGAAATGCCGGATACTCTGCTCTTAATATGCTGCATGCCGGCTGGAC
ATCTACGGAAAACCATTGCTGACCTAAGGATGCTAATCAACTGGAGGTGTTACGA
CAACTACATCGCAGATGAGATTAAGAACGGCCTTGTTAGCAGATGACCCCTGGA
GAGCAGCGTCTTACTTAGCCCCTAATATAAGAACGCCACTAAGCATGGGAAG
CACTCCATTGTAGCGTATTGGGCTATCTATCCAGTCAGCTGCAGCTAACATCAGC
ACAGGGGCTCCATCCCCACCAGAGGGAGTGAAAATGAACGAGCTGATCAGAAAGA
CAGTGTACGATCATGCTGTTGCAATTGTTAGCGAGTGGACAATGACAGGCTTCAAC
CAAGCACCGTCACCACTGTAATGATTGGAGGACAAGTTATCCCCTCAAGGGAGTTG
ACCCTAAGCGAGTGAATGACTTGTCAAGAATGTTCACTCAGAGACAGACCCCTTGT
ATGAAGTACCAACCCACAATCAGAGGAGGGAGCGATCACCCTCTGTTCCAGCGTC
CATACTAGTAGTAGAaGAGATGaTGAAGGGAGCTGGGAAGGAGGCAACGAAGAAGA
GTTGCTGAGAAAACCTCCATGAAAGGCGAGGCCAGTACGAGGAAGATACTAATTAG
GGGGATTATTTCAGCGACATAGACATCTGAAATATTATAAGCTTGTCTTTGA
TTCTTAGCTTAGTATTAGTGCATACAAAAAAACTAACCTCAGCTATGGCTAATCG
TGTAAGGAAAATTCTGTTGTCGTGAATGATCCGGCCGAAGCTTGGCAAGT
TGCTCTCAATATTGGCTAACGGGATCTGTCAACAGATAAGGAGCTGCTCTGCCG
AATTACGAAGTCAATGGTTGCCACATGTCTGTGGGTACGGGAACGTTAAACTCGA
CTGCGACGCCGTTGGAGAGTCTGGGCCCTCCTGCATTAGCATGGATGAAGAGCCC
TGTAGACGGAATTGCCATGTTGGCCTTACTAGAATTGGCGCAAACACGTACAGAG
GCTGCGAGCCGACCTGAGCAAGATGGAGACCCCTAGAATCGATGATTGAGCGGT
TAACAAATACCAGCATTGCCGTGTTCAAAGAAATTCTCAAAGAAAACAGCATC

ACTGACGACACCAAGAAGAAGCTAGAAAAGGCTGCAGCAATCTTGTGGATCTAGG
GAAGAAGTGCAGCTATACTGGATTTGCTGCTATCGGGATGATAGAACGTAGGAG
ATATGATGTCGCGGGTGTAAACGGCTCTGGGCATTGGGATCTGAGGATCCTATTG
GTGTGTGGAAGCCGTTGGAGTTAGATGAGAAAGCACCGTATCTTTGATGAAGTGA
AGCGCATTGGAGGAATCTCTGCAGAAAAACCAAGCCCACCTGAAAAGAAGATGTG
AGGTTATTCGGGAAGAAAGGAAATTGCAGCGCCGAGACAAGCTATAAGAGTGC
GCACCCGTTCTACTCAAGTATCTTCAAAGAACCAACCAAGTTGAAGGTCCGAGG
CGTGGAACTCCCATCTTGATGAAGTGATCGGTCTCTGCCAAGGGAGCAGAGGC
TGCAGAACTTCTGGACGACACCAAGGAGTTGGAAGGCGCATCAGCTGAACCAGAGC
TAGCCGATCATGCCGAGCCGAGCTGACTCAGTCACGCTCTGATTATCAGGGAGATC
AGAATTCACTCTAGCGTGCTCAAGGCTAGTCCCCACATGGACACAGTATAGACT
GCGACTCTAGCCAATCGGGCATCAAACCTTAGCGCTAGGAGCGCAAGCCTCTCC
CCCGAATGGAAGGAACCGGATGGAGGCCACCAAGGGCGATGATCCTGAAGAGGTAG
AGTGCGCCAACGAGATAGTTGGAGGAAATAGAAGGGTTAGCGAAGAACGGCTGCTGC
TTGAATGAAGACACCAGAGAAGAACGGCAGCCTCAATCTGATCAACACCCCTGTGAA
AAATTACAGTAGCTGGCAGAGTCCGAGCGTCCCTGGAAATCAGAAGTCTCTCCAA
GCACGCCGGAGGCTGCTGGATCGGAATAGGCCACTCCATAGCCCACAAGCAGAGG
CAGTTATGGATTCTGTAGGAGTTGCTTCTACCATCAAACCTCTAAAGTCCGGAG
TAACGAGGCCTATGCAGACAGATGAGCCGATGCCATCCTGGAAGCAGGATCGCAG
GGTATTACCAGCCTGTACCCCTCCCTCCAGTTGTAACCCGGAGTATTACTGCCGAGT
TCTCTAAAAGCTCCCTCCCCCGAAGGCGAAAGGGAAATTACCGAAGAGTCGG
ACTGATGTTACGCCGGATACGAAGAGGGCTAGAACAGAGGCCCTCAATTGGGAGG
CCAAATGGTTGACACGCATCTGGAAGGCATTGCAAAGTTGGCGACCCGATACTA
TTGGGTTTCAGCTAGGTAGCTACTTGCTCAACTCATTACAAATCCTTACCTTAAGAG
GGAGGAAACAAACCGTCACCAAAAGAGGATCACGCCCTCCACCTCCATCGGCAC
CTCGAAAAGAACCGTGAATAAATTCAAGTCAAGCTAGGCCCTAGAGCAGAACCGAC
ACCAAAAGAACATCCTGCACCGCACCCCTCTAGAGCAGAACCGACCCAGAAC
CTGGATGACTGGAAACCATCAAGGGTATATCTGCCAGGATGGATGACGTTATGG
AATTTCGCACCAATGAGGCGGAGCTGACAACCTCATTACATCATGTGTGCTGGAGG
ACAAGTTCACTGAGCCATATCCTTAAGCCTAAGGAGTTGTCAAAGAACAGCTAA
AGAATCTCTAGAGGTTGTCACAACATGGACAGAACAGGCTCCAGCTGTTATGTG

AACATTGACATTAAGAAACTACAAGAGCATCTCAGCTTGACAGCTAATTGGCAA
TGAAGATAAAAGCAGAGATGCCAAGGTCCATGGGAATATACTTCACAAGGCTAGGA
GAAGCATATGAAATGCCTCTGGAGGCGCTGCAAGTTGCTACTGCTGGCAGAGA
AGGGATTGACAAGTGGATCAGGGCTGCACTAGTCGTGAAGGCTAACACTGCGAGAG
GGAAAGTTCCCTCCCCCTGGCTGGTTGTGACAACAAGAGCAAAAGACAGCGACAAC
CCCAGCGTCCGTGCAAGAGCAAGTGCCTGTTAAATTCTAAATGAATAATTAG
TCTAACCCCCCTGTAGTTGTTAGTCTGCCTCTATATTAGCTTACAAGTAGTTCTTCATC
TCATCTCATTCAAAAAAAACTAACACAAGCTCGAATATGAGCAATTAGGTATA
CTCTGATGAAATTGCATCAAAGATGTCTTGACATCTAAATCTAAGTATAAGGTAC
TTGGAATTGGTGACGAATTGGGTCATCGAACATTATCCATGAGGGCGAG
GAAGACCATACTCGATTATTAGAGAGGCCATTCTAACATTCTAAAAAGAAAACAAGT
CGCACAAATGAGGCCTGGGTTATCCTACATATCCTGAGCAAAAGAACATAGTTAGA
GGAACACTGCGCGAGCCAGGCCTAACAGAAAGTTGTTCTGTCTATTAACTTAAC
GTCTCTGAGCCTCTGATTTCCGAAGTGGCACAGTCATATCAGACATTCTGAC
GGAATGAAATGTCTGCAATGAGAAAGCTGGTCTGCAAATGATGATCTGTCAACT
AGACCAATGGGAGGTTCTGATGGGTGTTACCGATACTTCAGTAATTGTTT
AGCACCTCATGCTTCCGTCTAGTGCCTAGGAGAGTCATGCATGATTGAGGGAT
TTCTGTCGACCACGTTGATGACAAGACATACTTGGATCATACAGGCTGACATT
GAAGACCGATAAGCTCCAATCATCAATACCCGCTCCCTGTGGTTCACTGTCTT
ACACCAAACGCGCTCAAGCCAGACTTCACCCCCGGACTAGCAGTCTGCAGAGCTAA
GGGAAGTTCTGATCTACCTTGATGACAAATACTTCAGCTCAAAACGTATAGCGAC
AACATCGTGTATTGCTGATGGCCGTGCTCTAACCTAAACTAGGTATAATGTATCA
CCGCTTGCAGTTGGATTTACAAAAAAACTAACCTAAACTAGGTATAATGTATCA
TCTAATTGTTCTCCTTGTATGCTAGGACAAAGAGCAGTAGCAGAGCCAGCTGACAT
TGCACGAGCCGCCAAGATCATACAGTATGCAGAGGAAAGGCTGAGAAATAAGACG
GGTCTGAGGATTGAGCTGTCATCGAGAGATCCGGCTAAAGCGCAGCATCCCCACC
TGATGTGCCTGCGACTAACGAGCCTGCTACCAAGCCCCTGCTACTACAGCTGCCGG
AACTTCAGCCCCGATTAAGGTGGGTTGCCTGAGTCTGAATACCTGCGTCAGGTTGT
GGAGCCTTGGTTCACCGTCTGTGAGAGCAAGGTCCAAGGACACTGACCGATGTAG
TGCATCGAATCCTCAGGAAAGGGATGGCAAAGCTCATAGAATGCTTCGAGGAACA
CCCAATGGAGATGATCACTCCGTTATACAAAATTGGATCCGGATCTCCCTGCA

CCAAGCCGACTGGTAGTCCTAGTGAATCATTGCAAGAAGTGCAAAGGCCGCAACT
ATATCGAGTCTCGAGGCTAAGGTCAAGCTCTACGCCAACCGGGTAGTACTATGCGA
AGTTATGTCTGCATTCTCACCTCTGAATGTCGAAGAGAGATGGAGTGCACGGGAGTACT
GGGTAAGCTGTCTGACACGTCCCGGTTACGCTCAATGTGGTGGACCCGGGTATGT
ACTGACAATGATCCCGCTCTGCTCTGCTCAGCAATGAGGGTCAAACGATGGACCTG
GACTAGCTTCTCAACGTGAAAGAGAGGGCTAGAACGACCGCTGGAGGATAACCCCTG
ACCCTAAAGAATGTGAGTCTCATTGCGAACCAATTTCGAGGAGAAGTGAACCTCGG
CTGTAAGAGGGAGTGTTCCTGATTATTGCTATGCTGGATGTCTACGTGTGAAGTTC
AGGGAGATGTGTACCAAGTCTCTTAGGAACAGCTAAATTCCATAGGTTCTGAACC
AAATAAGAGCCCCTTATTGTGGACGACCCCTGCACCCCATGGGCCATGTAAAG
GATCAGGGGATAGTCTCGTCTTAGCAAAAAATAGAACAGGATAACCCCTCGATTCAAC
ACCATGAATGGGAATTGACGCCAAGGTACAATTGGATACGCACAAAGTTATCA
GGTGCACCTGCCAGGAGTCACAACTAGCATCTGGACGCGTCATCGGTTTCTACA
TGGGGTTATGTGTATTATCAGCTAATGAGCGGCCGCATCGTTAGCGTAGCGTAGG
CACGCTACAATTGGTATAAGGTCAAGCCTCCGATCTGCACAGAACATGGAAGGGC
CGTATATGCCGTTGTCTTACCTGATTCTCAGGTTGCCCTCGACTTCTGAACAGTTGCG
GCAGGACCTGCTCCACTGCCAGACTAGAAAAGAACAGTCGTCCTTAATGCACTGGCAA
CTAAGAGGCTCCCACATCTATAACGCTTTGAAGGCTTGGTTACAAAGGATCGGAGT
CATACGGCTGGTGTCAAGGAAGGGCTCTGTATGCAGCACCTGCCATCAGTAG
AGTACACAGATCTGAGCATGTAGAACATATGGATGGTCGTAAATAATGGG
AGAGAAGTTGGGTGCCTGGATGGAGGGTTAAACTTGCTGTCAAGAGCGGTTGCGT
GGTTAATCCGAACGCATCCGTCTCTATCCTCTAGCGAGTGGAAAGGTAATATCTGA
TAGGGACGGCAAACCTTTAGCAGAACCAATCCAAAGGCAGGTTGGGGAGCATT
CAGCTCTCGAAAACATATCGGCTGCATTGGGGATTACTAGCCAGTTAGAACAGC
CGCCTCTGTGGATGATGGAACGGGCCATTATCATTCTACTAGCAACTAACAG
GAGACCCGGTCATTCCATCGGGCATCATCACTGTGGAGCTATGTCCTAGCCA
GTAAAATCACTGCTATTCTGATGCCCTTTATCCTTAGCAGTAGTGGTAGGGATAAT
AATGTGCTGCCGCAGATAACTGCATATCATGTACAAGAAAGATATTGTTACCTCT
CTCTGATAGCTTTAGTTAATCAAATTAAAAACAAAAAAACTAACACAAAGCTATGA
GTTGTTAACTCAAGATTGAGGAAGGAGATCGAGAGCGGGTGTCTAATAGCATA
AGCCAGAACATCCAAGTGTAAACAACACTTATAATTATCGAAGTTACTATGGTCGGC

ACTTTAGCCTATTCGATAGCGTGGGTGCCATCGTATCTCAGGCTCGCATA
AAGAATTCTGTAGAACGGAGACAACGATTAGGGAGATGGGTCTGCTGCCGA
TACTGGACCTGCGTAATAACTAAATAGGTCGATCCTTAGGTGGTACAAAAAAACTA
ACACTACCTCGTGTATGGAAGAAAAGTGCAGATTCCGATTACTTAAAGAGTT
GACTGACGCAGCTATAGAGGGAGTGTGGCATCTCCTTATACCCAATTACATTGGC
TACGGTCATAATATTCTTACTGCTAATATTGTAGTAGCGTGGAGAGCAGCTGT
GATTGCAAAGATACGACATCGTATAGATGAGTCCACAGCACGGCGCTCCGCCGTGAT
ITCAGGGGTCGACCTGATCACTCTCGTTATCATTGAGTCATAACCGATTCTGTG
GGCAATTATGCAGCCACACTCATCTTAACACTGTACCCTAGTCTGTCTTAGTCTG
CTACAATTATTTCTTTTCATCTGTAAACAACCTAACGGATATCCGAGGCCT
CATCTATCACTTAGGTAAATGAGGTCTCCGAATTGCGTCCATGCTTAGCAGCAGTT
TGTAGCGCACAATCGACATCTGCCAGACTAACAAAGGGAGCTGAACAGTTGCCCTA
AGTGGCCAAGTGACCGAAGTAACACCGAGCTGGGCAACGGCGAGCTGAGCA
TACCACAGCGCTACAGAATGCTTAGTAGAAAAGGACATGTTGGAAAATGAAATG
TTCATTGCAGAGCATTGCTAGTCTAACAAACCTACTTGATTCTCATCC
TCTCTGTCTATGTTGTGAACTGCTTGACTCAGTGATTCTCACATAACAGCCTA
AATATAACCGCAAATTATCTCTAAAAACAAAAAAACTAACACTATCTGTGTTATAC
AAACATGGAAAAATCGTCCCTTTGATCGCATTCCGGACTCTGTGCTGCACAGCCC
CCTTATCCTAGAGAGGAGTTGTGCTTGCCTCCAGACAAGGAAACCAGTGACTA
TCCAGCCATTCCGCAGCCAGACGAATTACAATCTCTGCCGCCGTGAGTTGATAGA
GCTCAGGTCACTGTCACTCACTGGGCAATCTCAGGAGTAGGGTGTATCAAATGGCT
CGCTTATATGAGATTGAATTGGATCGAACGCCAAGGATATTATGAGGCTCATGGCTGA
AGAATACCAATCATCCACAATCCACCCAGAGATAGCATCCGACTCAATAACGTGT
GTGAAGGTCTCTCAAGGCTGCATGACCAGTGGGAGAAAATCAGCATCCAGT
GCGCTTCAGGACATTGGAACGTTTGCGCAGTACCAATCAGACGCCATTATT
GACAACGGACAAAATACCGAACACTCTCAGCAGCTCTTATGGCTAACATCAA
GAGAGAGGCAACCAATCAGAACATTCAAAACAGCATCATTGCTAACAGCAGGATGGGG
AGTCCCGCCTATAATGGAGTGGAAAGTCATAACTCACAAGTTGTTAACTAAAAGAG
AAAGCTTGGAAAGCTAACTGTAGGCGCATACTGGTGTCTGGAGCTGCCAAC
AGCCTCCCAGATTGTCAGAAAATCACTATTGCTGTGAAAAGGACCTGTATTGC
AGCGCTGGATTCACTCTTGTGTAAGTCAAAACGCCCTCGGAGGAATCCCTT

GCCTCAGCCAAAGCACTT GATCAAATGTACCGAATGGGTGACATCATCCTCGCCAC
AGAAGGAAACGAAGGGTACAATGCAATCAAGACAATAGAAAGCATCATGGTCGGTT
CTGCACATAAGCAACTAGAAGACGAGTCAATTCCCCTACAGGTGACAAATTCTACG
AGTTTATCACGAAAGAAGGGCTTCAGCTGCCATCTAGAACATCGTCATCTGCTA
TGAGAAAAATCATCCATATAGGGGAGATAGTAACCCCTGAAGTTGCCATAGAGATG
GCAGGATGCTACAGACACTGGGCCACCCAACTCATCAATCCTGTTGGAGGTCTCACT
GCAGTCCGAGAGAACGCAACTGCCAGCTCCTACGAATGAAAGGCTCATGATATC
ACTAGCTGCTGATCTCAACTATTGTTGAGGAGTTATTGAAGAGCATGGCAG
GTGGCCCCGGAGTGAGATATGAAGCGCGCCAGAGATAGCAAAGACCTGTTCA
ACAAGTGGGTATCAACTAATCAATTCCCTGCCCTACCTCTGCTAGTCAAGTCAGGA
ACAGCTGGTTTTCGTAACATATGACAGCTTATTGACAAAAATCAAGAAATTCCAAT
TCCTCAGCCTAATTCAAGACAAGTCTCACTCAGTTGGACGATCCGCCGTCAACTA
TGTGCCTCAAAAAGAATCTGCTGCTGAGTCCTCACGGAGGGTCTCAATCAACTC
TCAGCTATGCAGAGATCGATGTGAACAAATTGGACTCTATAGACACAGCACAGAA
AATGGTTAACGAAACGACGACTTAGTCATCCTATTGCGAGAAAAaGAGCGAGAGCTC
AAGGTAAAGGGCGGTTCTTCCCTAATGACGTACAAATTGAGGACTTACTTACA
GCAACCGAATACCTAATTGCAAAGCACATTCTGCCGTACTCCCTGAAATTACAATG
ATGCAGGGGCAAATTGACCTGTGGAAGACTTCAAAGGTGCGGTACGCACTGTGAG
TCAAGAGAAGTCTACACATCACATGATTGATGTTGAAAGACAAATTGGAACAAACTT
CCAGAGAGAAGAGTCTACAGCTCCGGTTCCAGATTATGGACAGAGCCTCGGGTG
GTCAAATGTCATATCAAGGACTCATAACTTCTCTAGGTGCTCGTCGGCTATGCT
GGGAGAATAGATATGTCCTATCGGTCTGACAGACAACTGGCCGTGGTGTGGACC
GGACACAAAGGAGGCTTGAAGGGCTAAGACAGAAGGGATGGAGTGTGTTAGGGG
CTTGCTAATACGGCACGTGATGAGACTCACTGGCTGCTGATGGAAAGTGCTAATT
AAGGAGATAACCAGGTAATCATACTTGAGTACCCCTTGCCAGCTCATCGAATACTT
CATCGCTGCTCGAGAGGCATAGGCATTGAAAATGACAACACTCACTTGTCTG
TGTTCTCTGAATTGTCTAACGAGCATCGGATTAAGAACATTAGCCAGAAGAGACATGG
ATATCCTCGAGGATGGTTACTACGGGAAGTCCCTGTGATCGACGGATCGCAAGA
GGAATGGTGCTTAAGAAACTCTGCAGGATCTTGCAATATCAAATGACCTCACTCCT
AGCATCTCAAATAGCATCTCTCTGCACACGTGATGCCGGCTGTATCCAAT
CATCATACGCTGCCAATTACATGTTGCTATCTTATGGCTTCTGCTCCAC

GACTTCTTCGAATACAACCCCTGTCCATGGAGCCTATGATCTTCACATTCA
GAGACGG
GCTTTATTCGGAACAACTGCTGTCGAGACATTACCAAGGTACGAGCGCCCACAACC
AAGGACTCTATCATCATTCTGTAGCAAGAGACTGGGCACCAGCAAGCAATAAGCA
GCAATCGGCTCTCGGACTAGACCTCCTGACCGAGATTCACTCTGGAGGTCTGG
AGGAGCATCCATTAAAGTATATGATCAGGCAGTTCTGATCCTGTACAGAGGG
CCTATCATTGCAAAAGTGGGTTAAGCAAGGCAGAACACCTGAGCTAAAGACCA
TTTCTCTTATGGAGGCCTCCATCAAGTCAGGCTCAGTAGAACTCCTCTAGA
AGCACCCATGTCGATCAATCTGCCATCCTACTCGCCTGTAAATGTACTGCCGCG
GTGGTTGAAGAGAAGATCACACAGAACATGGACACGCTGGTCAACAATCAGCTTG
TGTATCAGGCCTGTCGATGAAGCAAGATATGCGACCGCATTCTGAAGATGTGTG
CATCCCTACAGCCATTCTCCCCGATTGCTTCTCTGTACAGTGCACCCCAT
AGGAGTGGCAGAAACAATTGTTGAAAATTCTCGGGACTAAAAGTGTGCCAGAA
TTGCTCTCAGAGAGGGTGCAGGGCAGCTGGAAAGAAACTAGTAGAAAGCGAAGTT
GAGTTGATAACAACGTATCTAACAGTTCTCAAACGTCTGCTGGAGCATTGGGT
TGTTCTAGCAAATTGGCTGACAGCTATAGGTCGAAAAGTTGGGAGGCCAAATTG
GGGAGTCACAGTGCCTCACCCCTGCAGAGCAGTTACTTAGAGTGGCCAATGACAA
GCAGGGTCTCAAGGAAAGCCTGCTGGATGAATGATTTGCCAAGAACTGTGA
AAGTTAGCTTGAATGAAAGAATGATCTATGGAGACACCTCTCCGCGATGCCGAC
CTATGAATCCTACTTAGGTGGTCGAACTAGTGAGCGGCTCTGTCCAATCATGGC
TAGAGCTCGAAACATCATCTTCTATTGCAAAGAGACTTGTAGAGCTAATAATGCAT
GTGGTTGGGCATATCAAAAGGGTTCGCACTTGCATCACGCAATAAGAAATGCTC
ACACAAATTGACTGGATTAGTGATTCCGAGCGATGCAGTTGTAGAGCTAATAATGCAT
GGGAGTTGGGACATAGGTTCTCAGATCCTAGAGTGTCTCTGGAGCAATGTCAGCC
ACCAACTATAACACTGCCACTCATATTGCGTTCAAGTACAAACGACCTAGTGAGGTTG
AAGCGCGGGGAGGACAACATATGGTTGTTCCAAGGGCTATTGTGTTGACAC
AGCATTATTGGGAGCTAGTAAGGAGAGGAGATAATGGGTCTCCGACTGCCACCT
TCACCCCTAAATGCGAAGAATGCATGATACCATGTCCAGATGTTCAATGGAGCTTC
AAACCCGGTAGTCCCGCTTATTCAATGTTTATCACCTCGCCTCCGAATGATTTC
TTAACGTTGCCGGAGAATCTCTATGTTGACATTGATCCGTTCAAAGTAAAGCCTGAT
CAAATCAAGACTATTGTCAACATCGTCTGCATACAGTACAGTCTGTGACACC
CTCTCGGGTGAGGAGATATCAGATGCTATATCCAATGTTGCAGCAATCATAGCAGTA

ACAAAAACAAATCCTCAAGGAAGATGGACTCTGGGATACGAACTGCCATCCCGT
GTATGCAGGTTACAAGTGCTCGTGGAAAGGTACTTAAGAGCAAGATTGTATTCTGGAC
TTTGCTATGCTGCATTCCGGCACATCAATGAGTCAGAGGAGCATGACTGCTGTGA
GGCTTGTCAGCACATACTTAGAAAGCAAGCAGCAGATCACTGGAACCTGCATAACC
CAACAATGCTGGGAGGGATGCTGAGCAACACGGACATCATTGTCGAGTTGGCTAAA
GAGCAGATTGCGCTCAGCGCAGGATTCCGATATCGCCTATGGAACTTCAAGCGCA
GCAGGACAGTCATTATTGAGCAGTGTCTCCCATGGAAAAGAATTCTCACTACCA
GAATTCCCGAAGGGAGTGATAGCTCTCCGAGAGACATGACAGGAGGTGGAACTC
GGCATATTAAATCGCCTGCATGATGTGCTATGCCGCTTCTCAAAGGAGCCAACAC
GGGACCACTAAAAGCACTTGCAAGGGCTGACCACTGAGATGAAAACATCTGTTATAG
CAAGCCAAGCATTGAAAGAAAGCGAAGGAGTCTGGAGCAGGCTGTAGATATGCTT
CACGAAAACACATCAACCTTGCTGCGTGCTGACATTGACTCAAACAAACTAACA
AAAAGGGGTCAAGACGTAGATCATATTCTCCGGAACATACCTGCTATTCACTT
GATACCGTCAGAGGGCATACTGTATATGGAACACACAGCATCAACAAACAAAGGGCC
TGAATCTTCTTACTTGGAAAGAGGGAGGGAGAGCCTGAGGTGACACAAACCCCTATCA
TGCATCTATTCCGCCCCATCTTAATGTCTGCTAACGGTTCAATCAAGATTCTGTCTGT
CTTGAAAACATCCAGCTCAACCAAAAAATGTTGCAGTGGTCGGAGATGGTAATG
GGGGATTGGGAGGGCACTATTGGGTCTCTACCCTACAGCTGAGTTCTTTCAATTC
CCTATCTGAAATGTCCAACTACGGCGATCAAGGGAGTCACAAATTACCCGCTTG
TGTGCTAGACTCTCCAAGTGATTGGGAAGGGTGATCAATCTGGAGAAGTCTTAAC
ACTGCCTTCAGACTTGCTCTCAGAGAGGTGGCCTCCGAGGTAACGAAATACTGGAT
GAAGAAATGCTTGACCCAGACCTGCTGGTGTGCGATGCAGAGAGCTATGATTCTAT
CAAGAGGTCAGAAATCTGAGCAACCTCCTCAAGGTATGCGGCCAGCGGTCTCACG
TCATAATCAAAGCCCACTTGTCAAGGATGCAGGACGTCCAGAGCTGGTCAATGCGC
ACTTGCATTCACACAATGTGATGTACTTGAGAGCCCCTACAGCAACTGGGGAAAGT
TTGAGGTGTACTTGTTGACACTCGAATTGTAGAAAGAAACGATATGAGGTATGCAC
CAAAAGATACATATCTCATGTGGACTAGAGCTGAATTCAACGCTCACCCCTAGCTCC
AATTAAACAGCGTCCACAGCTGGTTTCCCTCATCTTGAAAAAGACGATAGACTCC
GAGCTGTCCGGCGGATGCTAACGCTACGACGGCACATTAGCAGGTACACTAAGATA
AGACCGCAAGAGCTGTCACGAGTAGAGAGGAAAGCAAAAGAGCACGGAGACATGG
GGTTCTGTGAATTGCTGTCACTTAGGATCCAAGTTCATATCATCTATGAACA

ATTCTTTATCAGGGCAACTCACATGCTGGGAGATTGAGAGGCCAGCTAAAGCGAT
CGGAGCTGATCGGGCTAATAGGGACTGCTATCGGTGCCTCATGCTTGGTGTCAAGCCA
TTTCTCATCCGCATGATTACTGCTTGTCTTGAACTTACAAGAATGGGGTGGATGT
TGTTTGGAATGATAAAGGAATATCAAGCCAATTGTGGTCAGAGCCTGGTCAG
GTGAGTCCACTCCTGTAACTATAACCCCAAAGGAACTCAAAAAGCCTGTTCAATTCTA
TGGTAGAGATACTTGGGGGCTTATTGTGGAGCTCTGGTATTGATAGAATCACTCAAA
GGTTAGAGTGGCTGGTAAGAGAAAAATATGAACCTCGGCTCTCGGAAGGGTACAAT
CACTTGGCAGTAGCCACTTGGGATAACGATACCTAGACGTAGATCCATTGGAGTGT
ATAAGATGGGAGATGGAACAAGCTGCCGTCGAAGAAATCGATTCGGGGGATAAT
TGATGAATAGCACTGATTCTAAAATAGTATCAGAGAAAAACAAAAACTAAAAA
AACAAAAAAACATATACATATAAAATTAGGAGCAATAACTGAAGATCAAAGAG
CATTAAAAGAGATCTAAGAGAAAAAGAGAGAAGGACCTGGCAAAAGCTAAAAGA
TAGAAATAAGGAAACAAAGAATTTCCTGAGAGGGCAGCACATAAGGATGATAGG
GATCACATGAAGGCTTCGGAACCTTGAAGCTTGTAAAAAAAGCTCAAAGACAAAA
ATTGCAGAGAAGAGGGAACCCAAGTTGCTCTGATTGCTGACTGGCGACATCTGAC
CATTTGCTCTTACTGAAAGACCAATCCTGATATAGATGACTTCTAGTTGTCTGGCT
ATGAGTGTCACTCGTACAAGGCATGGGTATGGGGGTATGGAGGGCAGCACCATT
GTGGGT

*Coding sequences of identified open reading frames are underlined; deduced amino acid sequences below, overlapping sequence of open reading frames U2 and U3 gene are in italics.

Deduced Amino Acid Sequences RFFRV Genes

N Gene RFFRV

MDHDNEKPISYTSIAEVPDNVAIGSTIYIQGEPIIYFGKSAATGITRKGGAQKDWTKDMIR
GVRVFLPQTDANLLNLIAGETEAPELEKYTIQDPEKKGILKKFESKWEFANWANLLVDL
QSNTGNIPKGRFPYYSSALFSITAIKGAPVLAPAMKDLGDPVYVKAPDDLHPPTGDIEWH
GDKISVDEAA~~YIGYGAWLIMPRFTIKAESKKDEIAASSKAFDTLRLLLPEITKPQVLVSVV~~
TQLRLAYHGTLVPGSAYLAAEVAMRRAMNIEYDLKADRTECKAGEHFGCQLRVLQDI
PQYDSGFWGFGQVGLEMAGYSALNMLHAGLDIYGKTIADLRMLINWRCYDNYIADEIK
EGPLLADDPWRAASYLLAPNIRPLSMGKHSIVAYLGLSIQSAAANISTGAPSPEGVKM
NELIRKTVDHA~~V~~AVIVSEWDNDRLQPSTVTTVMIGGQVIPFKGVDPKRVNDLSRMFTQR

QTPLYEVPPHNQRRERSPSVSSVHTSSRRDDEGSWEGGNEEELLRKLHERRGQYEEDTN
LGGFYSAT.

P Gene RFFRV

MANVRKIHSVVSVNDPAEAQVALQYWAKRILSTDKELLCRITKSMVAHMSVVYGN
VKLCDAVWRVWALLALAWMKSPVDGIAMLALLEFGAKHVQRLRADLSKIGDPRIDD
FERLTNTSIARVPKKFFKENSITDDTKKKLEKAAAILSDLGKKCSYTGFCCYADDSNVGD
MMSRVLTASGALGSEDPIGVWKPLEKAPYLFDDEVKRIGGISAEKPSPEKKMCRLF
GKKKGKFAAPETSYKSAHPFSTQVSSKEPPKLKVRGVELPSLDEVIGLLPKGAEEAELLDD
TKELEGASAEPADHAEPQLTQSRSDYQGDQNFTSSVLKASPPHHSIDCDSSQIGASN
LSARSASLSPEWKEPDGATKGDDPEEVECANEIVGGNRRVSEEGCLNEDTREEGSLQS
DQHPVKNYSSWAERFRASLEEVSPKHAEAAGSAIGHSHSPQAEAVMDSVGVASSTIKPL
KSGVTRPMQTDEPHAILEAGSQGITSLYPPLPVPGVLLPSSLKSSLPPKAKGKFTEEFGL
MLRGIRRGLEERGLNWEAKWFDTLEGIAKFGDPDTIGFQLGSYFAQLITNPYLKREET
KPSPKEDHAPPPSAPRKESVNKFQARPHSQVPTKEDPAPHPPRAEEPDRSWMTGKPS
RVYLPGSDDVMFRTNEAELDNFITSCVLEDKFTEPYILKPKELSKEQLKNLLEVVSQHG
QKASQLLCEHLTLRNYKSISALTANWAMKIKAEEMPRSMGIYFTRLGEAYGNAFWEALQ
VATAGREGIDKWIRAALVVKANTARGKVPPPGWFVTTRAKDSNDNSVRARASALFKFL
NE.

M Gene RFFRV

MSNFRTLMKFASKMSLTSKSKYKVLGIGDELGQSNVNIIHEGEEDHTSIYSESPSSSKK
KTSRTMRPWVYPTYPEQKNIVRGLRGARPKKVVL SINLTGVSEPLDFPEVATVISSDILD
GMEMSAMRKLVLQMMILSTRPMGRFSDGCYRYIFSNCFSTSCPSSCLGEFMHDSDGDFC
STTFDDKTYFGSYRLTFEDAIGSNHQYPLPLWFHLSYTNALKPDFTPGLAVCRAKGKFLI
YLDDKYFSFKTYSNDNIVLLLGRAPKAIKQK.

G Gene RFFRV

MYHLIVLLVMLGQRAVAEPADIARAQKIIQYAEERLRNKTGLRIELSSRDPAQSAASPPD
VPATKQPATKPAATTAAAGTSAPIKVGLPESEYLRQVVEPWFTVCESKVPRTLTDVVHRIL
RKGDGKAHRMLRGTPNGDDHSVYTKFGSGISPAPSRLVVLSESLQEVRPQLYRVSRRL
SSSTPHRVVLCEVMSAFSPLNVEEMECTGVLGKLSDTSSVTLNVVDPGYVLTMRVCSC
SAMRVKRWTWTSFFNVKERLEDRWRIPPDKECESHCEPIFAGEVTSAVRGSPDYCYA

WMSTCEVQGDVYQVSLGTAKFHRFLNQIRAPFIVDDPCTPSAPCKGSGDSLVLAKIEED
NPRFTTMNGELTPRYNWDTHKVYQVHLPGVTSILDASCGFLHGGYVYYQLMSGRIVS
VSVGTLQFGDKVKPPICTEWKGPYMPFVLPDSQVASTSEQLRQDLLHCQTRKEVVLNA
LATKRLPSITLFEGLGYKGSESYGLVSRKGLLYAAPCPSVEYTDLEHVEGANIWMVVNN
REVGCLDGGLNFAVKSGCVVNPNASVSILLGEWKVISDRDGKLLAEPIPKAGWGSIPAL
ENISAAFGDYLASLEQPPLWDDGNGPIIPTSNSTGDPVIHSGASSLWSSMSLASKITAILM
PLLSLAVVVGIIIMCCRR.

U1 Gene RFFRV

MSCLTQDLRKEIRERVSNSISQNPSVTTLIIIIEVILWVGTLAYISIALGCHRYLQARIKNSVE
RRQRFRGDGVLLPDTGPA.

U2 Gene RFFRV

MEEKCSDSDYFKELTDAAIEGVWASPLYPITLATVIIFLILLIFVVAWRAAVIAKIRHIDE
STARRLRPDFGGRP.

U3 Gene RFFRV

MSPQHGASALISGVDPDHFSLIIESYPILVGNYAATLILTWTLSFLATIIFSSVNNLT
DIRGVLIYHLGNEVSEFASHALAAVCSAQSTSARLTRELEQFRLSGQVTEVTPSSGATGE
LSIPQRYRMLLVEKDMLENEMFIAEHSV.

L Gene RFFRV

MEKSFLFD RIPDSVLHSPLILREELCFCRPDKETSDYPAIPQPDEFTISRRPELIELRSLSLTG
AISGVGCIKWLAYMRLNWIEAKDIMRLMAEEYQSSTIHPEIASDFNNVCEGLFSRLHDQ
WGENQHPSALQDIWNVFVAQYQSDAIIDNWTKYRILSAALLWSNHKERGNQSEFQNSII
AKAGWGVRAINGVEVITHKLTKRERLGKLTVGAYWCLELANQPPRLFRKSLLLCEK
DLYLQRWISLFLCNCQNAFGGIPLPQPKHLIKMYRMDIILATEGNEGYNIAKTIESIMVG
SAHKQLEDESISPTGDKFYEFITKXGLSACHLESSSAMRKIIHIGEIVTPEVAIEMAGCYR
HWGHPIINPVGGLTAVRENATAQLPTNERLMISLAADLNYLLRSYFEEHGRWPPGVRY
EAAPEIAKDLFNKWVSTNQFPGPTSASQVRNSWFFVTYDSLFDKNQEIPILSISDKSHSV
GRSALSTMCLKNLLSPSRRVLQSTLSYAEIDVNKFQIDSIDSTENGLSNDDLVILLREKE
RELKVKGRRFLMTYKLRTYFTATEYLIAKHILPVLPEITMMQGQIDLWKTFKGAVRTV
SQEKSTHHMIHVDFEKWNNFQREESTAPVFQIMDRAFTGWSNVISRTHNFFSRCFVGYAG
RIDMFPIGLTDNWPWCWTGHKGFFEGLRQKGWSVVGALLIRHVMRLTGLHGKVLIQG

DNQVIILEYPLASSNTSSLALERHRHSKMTTHFLSVFSELSKSIGLRIKPEETWISSRMVY
YGKFPVIDGSARGMVLKKLCRIFAISNDLTPSISNSISSLHTSCIAGCIQSSYAVPIYMFAIF
YGFLLHDFFEYNPLSMEPMIFTFRRALFGTTAVRDITKVRAPTTKDSIIISVARDWAPAS
NKQQSALGLDLLTRDSSLGGASILKYMIRQFPDPVTEGLSFAKVGLSKAETPELKTI
FSSYGGPLIKSGSVELLLEAPMSINLPSSTRPVNLRRFVEEKITQNMDTLVNNQLVYQA
LSMKQDMRPPFLKMCASLQPFFPRFASSLYSATPIGVAETIVGKFLGTKTVARIALREGA
GQLGKKLVESEVELITTYLTVLKLSALEHWGCSSKLADSYRSKSWGGQIWGVTVPHPAE
QFYLEWPMTSGCFKESLAWMNDFAEETVKVSLNERMIYGDTSPRSAGPMNPYLGGRTS
ERSSVQSWELETSSSIAKRLVELINACGWAYQKGSHLHAIKEMLTQLTGLVIPSDAVC
RITESGSLGHRFSDPRVSSGAMSATNYNTATHIAFSTNDLVRALKRGEDNYMVLFQGLFV
YFTSIIGELVRRGDNGSPTAHLHPKCEECMIPCPDVSMEASNPPVPLYSMFLSPRLPNDFL
TL PENLYVDIDPFKVKPDKIQTIVNIVPAYSTSVDTLSGEEISDAISNVAAIIAVTKTNPQ
GRWTLGIRTAIPVYAGYKCSWKVLKSKIVFWTFAMLAFRHINESEEHDCCEACQHILRK
QAADHWNLNHNPMLGGMLSNTDIIVELAKEQIALSAGFPISPMELOSSAAGQSLFEQCVS
HGKEFSLPEFPKGVIALPRDMTGGGTPAYLIACMMCYAGFFKGANTGPLKALAGLTTE
MKT SVIASQAFEESEGVWSRLVDM LHENTSTLCCVLDIDFKTLTKRGQDVHDHILPGSIPA
ISLDTVRGAYCIWNTASTTKGPESSYLEEEGEPEVTQTPIMHLFRPILMSANGSIKILSVLK
TIQLQPKNVA VVGDGNGGFLALLGLYPTAEFFFNSLSEMSNYGDQGEFTNYPACVLD S
PSDWGRVINLGEVLTLPSDLSSERWPSEVTKYWMKKCFDPDLLVCDAESYDSIKRSEILS
NLLKVCQQRSHVIKAHLSRMQDVQSLVNAHLHSHNVMYLRAPYSNWGKFEVYLWYT
RIVERNDMRYAPKDTYLMWTRAEFNAHPQLQLTAFHSWVFPHLEKDDRLRAVRRLMS
LRRHISRYTKIRPQELSVERKAKEHGDGMGFCELLHYLGSKFISSMNNSVIRATSHAGR
LRASLKRSELIGLIGTAIGASCLVSAISHPHDYCLFLEYKNGVDVVLGNDKGNIKPIVVR
AWSGESTPVTITPKELKKPVHSMVRYLGGLLWSSGIDRITQRLEWLVREKYELGFSEGY
NHLAVATWDNDTLDVDPLECIRWEMEQAAVEEIDFGGIIDE.