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## Supplementary Data for "Sequence and annotation of the 314-kb MT325 and the 321-kb FR483 viruses that infect *Chlorella* Pbi": Appendix A: Gene Names m002R through m843L

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Fitzgerald, Lisa A.; Graves, Michael V.; Li, Xiao; Feldblyum, Tamara; Hartigan, James; and Van Etten, James L., "Supplementary Data for "Sequence and annotation of the 314-kb MT325 and the 321-kb FR483 viruses that infect Chlorella Pbi": Appendix A: Gene Names m002R through m843L" (2007). Virology Papers. 2.

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Published in Virology 358:2 (February 20, 2007), pp. 459–471. doi:10.1016/j.virol.2006.08.034 Copyright © 2006 Elsevier Inc. Used by permission. Submitted July 27, 2006; revised August 18, 2006; accepted August 23, 2006; published online October 4, 2006. Supplementary data originally published online at doi:10.1016/j.virol.2006.08.034.

#### SUPPLEMENTARY DATA FOR

# Sequence and annotation of the 314-kb MT325 and the 321-kb FR483 viruses that infect *Chlorella* Pbi

Lisa A. Fitzgerald<sup>a</sup>, Michael V. Graves<sup>b</sup>, Xiao Li<sup>b</sup>, Tamara Feldblyum<sup>c</sup>, James Hartigan<sup>d</sup>, and James L. Van Etten<sup>e, f, \*</sup>

**Abstract:** Viruses MT325 and FR483, members of the family Phycodnaviridae, genus *Chlorovirus*, infect the fresh water, unicellular, eukaryotic, chlorella-like green alga, *Chlorella* Pbi. The 314,335-bp genome of MT325 and the 321,240-bp genome of FR483 are the first viruses that infect *Chlorella* Pbi to have their genomes sequenced and annotated. Furthermore, these genomes are the two smallest chlorella virus genomes sequenced to date, MT325 has 331 putative protein-encoding and 10 tRNA-encoding genes and FR483 has 335 putative protein-encoding and 9 tRNA-encoding genes. The protein-encoding genes are almost evenly distributed on both strands, and intergenic space is minimal. Approximately 40% of the viral gene products resemble entries in public databases, including some that are the first of their kind to be detected in a virus. For example, these unique gene products include an aquaglyceroporin in MT325, a potassium ion transporter protein and an alkyl sulfatase in FR483, and a dTDP–glucose pyrophosphorylase in both viruses. Comparison of MT325 and FR483 protein-encoding genes with the prototype chlorella virus PBCV-1 indicates that approximately 82% of the genes are present in all three viruses.

Supplementary data associated with this article is archived in this repository as 4 separate files: Appendices A–D. Each document, in spreadsheet format, shows Gene Name, Genome Position, A.A. length, Peptid e Mw, pI, CDD Hit Number, COGs, COG Definition, Bit Score, E-value, % Identity, % Positive, Query from-to, Hit from-to, BLASTp Hit Number, Hit Accession, BLASTp Definition, Bit Score, E-value, % Identity, % Positive, Query from-to, and Hit from-to.

Appendix A: Gene Names m002R through m843L Appendix B: Gene Names M001L through M807R Appendix C: Gene Names n001L through n849R Appendix D: Gene Names N003L through N847R

Appendix A: Gene Names m002R through m843L

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Gene Name	Genome Position	A.A. length	Peptide Mw		CDD Hit Number COGs	COG Definition	Bit Score	E-value	% Identity	% Positiv	from to	LASTp Hit lumber	Hit Accession	BLASTp Definition	Bit Score E-value Ide	% %	Query F e from-to	Hit from- to
m002R	4061008	201	21,987	11.12	No Hit Found					·			No Hit Found					
m004R	13451566	74	8,032	7.02	No Hit Found								No Hit Found					
m006L	19841775	70	8,219	11.40	No Hit Found								No Hit Found					
m008R	27513119	123	14,086	9.94	No Hit Found								No Hit Found					
m013L	53235072	84	9,564	6.53	No Hit Found							1	NP 048541 a194R		74.33 1.17E-12	49% 589	% 179	1-79
m016R	80028271	90	9,632	10.81	No Hit Found								No Hit Found					
m017L	94459146	100	11,875	12.25	1 pfam05743	Tsg101, Tumour susceptibility gene 101 protein (TSG101). This family consists of the eukaryotic tumour susceptibility gene 101 protein (TSG101). Altered transcripts of this gene have been detected in sooradic breast cancers and many other human mallonanc	37.01	1.80E-03	22%	43%	978 244313		No Hit Found					
m018R	94739682	70	7,726	7.15	No Hit Found								No Hit Found					
m020R	1055010780	77	8,549	10.27	No Hit Found								No Hit Found					
m021L	1098510761	75	8,826	12.85	No Hit Found								No Hit Found					
m022R	1227812502	75	8,225	10.51	No Hit Found								No Hit Found					
m027L	1391913683	79		12.51	No Hit Found								No Hit Found					
m028R	1388314077	65		11.07	No Hit Found							1	NP_048404 contains type 1 hydro	phobic transmembrane region	59.31 3.85E-08	59% 669		43104
m029R	1398414631		23,949	7.66	No Hit Found							1	NP_048403 a55L		72.79 7.93E-12	24% 589	6 1122	10131
m031L	1537814911	156		12.18	No Hit Found								No Hit Found					
m033R m035R	1623416551 1670416919	106 72		6.24 9.71	No Hit Found No Hit Found								No Hit Found  No Hit Found					
m036R	1721217415	68		11.66	No Hit Found								No Hit Found					
m038L	1767017449	74		4.99	No Hit Found								No Hit Found					
m039L	1811417800	105		8.68	No Hit Found								No Hit Found					
m040L	1842118068	118		9.90	No Hit Found								No Hit Found					
m041L	1869918385	105		6.49	No Hit Found								No Hit Found					
m043L	1952219256	89	9,827	4.45	No Hit Found								No Hit Found					
m044L	1997519673	101	11,052	4.36	No Hit Found								No Hit Found					
m045L	2157721281	99	10,863	4.53	1 pfam01698	FLO_LFY, Floricaula / Leafy protein. This family consists of various plant development proteins which are homologues of floricaula (FLO) and Leafy (LFY) proteins which are floral meristem identity proteins. Mutations in the secuences of these croteins af	35.77	4.63E-03	24%	33%	1872 660		No Hit Found					
m046R	2177621973	66	7,552	10.84	No Hit Found								No Hit Found					
m048L	2262222185	146	16,111	4.33	No Hit Found								No Hit Found					
m049L	2292822632	99	10,983	8.40	No Hit Found								No Hit Found					
m050L	2334523046	100		4.71	No Hit Found								No Hit Found					
m051L	2436624076	97		4.31	No Hit Found								No Hit Found					
m052L	2465324366 2492324687	96		3.85 8.43	No Hit Found								No Hit Found  No Hit Found					
m053L m054L	2558025374	79 69		9.24	No Hit Found								No Hit Found					
m056L	2737126628	248		4.65	No Hit Found								No Hit Found					
m057R	2689427187		11,613	6.89	No Hit Found								No Hit Found					
m058L	2770727489	73		4.16	No Hit Found								No Hit Found					
m059L	2839728056	114	12,391	4.58	No Hit Found								No Hit Found					
m060L	3098430757	76	8,621	7.81	No Hit Found								No Hit Found					
m062L	3158431147	146	15,831	3.99	No Hit Found								No Hit Found					
m063L	3230731633	225	23,962	4.66	No Hit Found								No Hit Found					
m064L	3291632575	114	12,445	4.37	No Hit Found								No Hit Found					
m065L	3417733806	124	14,508	12.15	No Hit Found								No Hit Found					
m066L	3497734780	66		9.51	No Hit Found								No Hit Found					
m067L	3551135083	143		10.10	No Hit Found								No Hit Found					
m068L	3558735117		18,213	8.39	No Hit Found								No Hit Found					
m069R	3540336698		49,571	9.77	No Hit Found							1	NP_048711 A354R		198.36 3.93E-49	42% 589	6 187432	2235
m072L m073L	3735636865 3828938026		17,831 9,774	10.78	No Hit Found								No Hit Found  No Hit Found					
m073L m074L	3828938026 3893138665	88		4.40	No Hit Found No Hit Found								No Hit Found No Hit Found					
m075L	3917438968	69		4.88	No Hit Found								No Hit Found					
m076R	3931639564		9,924	8.93	No Hit Found								No Hit Found					
m077L	4015539955	67		9.42	No Hit Found								No Hit Found					
m079R	4136941662	98	11,476	12.04	No Hit Found								No Hit Found					
m080L	4290942484	142	16,161	9.58	No Hit Found								No Hit Found					

Gene Name m082L	Genome Position 4310342711	A.A. length	Peptide Mw 14,511		DD Hit umber COGs	COG Definition	Bit Score	E-value	% Identity	% Positiv e	Query from-to	HIL HOIH	BLASTp Hit Number	Hit Accession  No Hit Found	BLASTp Definition	Bit Score E-value Id	% entity Po	% ositive f	Query I rom-to	Hit from- to
m084L	4364643356		10,346	11.01	No Hit Found								1	NP 048548 A201L		119.40 3.15E-26	78%	91%	2593	2492
m087L	4583845353	162	18,201	10.11	No Hit Found									No Hit Found						
m089L	4660145978	208	22,820	11.93	1 COG3119	AsIA, Arylsulfatase A and related enzymes [Inorganic ion transport and metabolism].	35.87	3.82E-03	3 27%	43%	68143	3 871		No Hit Found						
m092L	4815547943	71	8,214	10.98	No Hit Found								1	NP_048647 a293R		57.77 1.10E-07	56%	65%	1771	559
m094L	4908948862	76	8,968	9.95	No Hit Found									No Hit Found						
m095R	4904249245	68	7,667	7.51	No Hit Found									No Hit Found						
m096R m098R	4913949405 4963650043	89 136	9,821	8.27 9.47	No Hit Found									No Hit Found  No Hit Found						
m102L	5246352128		11,772	8.35	No Hit Found									No Hit Found						
m104L	5359853269	110	12,338	12.81	No Hit Found									No Hit Found						
m105R	5350153740	80	8,771	10.60	No Hit Found									No Hit Found						
m106R	5359253804	71	8,282	11.67	No Hit Found									No Hit Found						
m108L	5421053923	96	10,934	8.46	No Hit Found									No Hit Found						
m110L	5537855175	68	7,587	11.46	No Hit Found									No Hit Found						
m112L	5611055808	101	11,820	8.08	No Hit Found									No Hit Found						
m113L m115R	5642156131 5685057080	97 77	11,314 9,170	9.26 10.32	No Hit Found No Hit Found									No Hit Found No Hit Found						
m117R	5707057549	160	18,272	8.74	No Hit Found									No Hit Found						
m118R	5719757397	67	7,617	11.31	No Hit Found									No Hit Found						
m119R	5777258005	78	8,655	9.54	No Hit Found									No Hit Found						
m123L	5914358925	73	8,218	8.43	No Hit Found									No Hit Found						
m125R	5918359386	68	7,610	9.23	No Hit Found									No Hit Found						
m126L	5954659349	66	7,634	10.39	No Hit Found									No Hit Found						
m127R	5996960166	66	6,979	11.55	No Hit Found									No Hit Found						
m129L m130R	6143660564 6098261284	291 101	32,015 10,868	7.60 11.97	No Hit Found No Hit Found									No Hit Found No Hit Found						
m131L	6145061181	90		10.73	No Hit Found									No Hit Found						
m132R	6134861668	107	11,636	10.71	No Hit Found									No Hit Found						
m134L	6308062343	246	27,343	10.28	No Hit Found									No Hit Found						
m135L	6346663209	86	9,385	11.43	No Hit Found									No Hit Found						
m138R	6399364250	86	9,861	8.21	No Hit Found									No Hit Found						
m141R	6555465787	78	8,665	11.88	No Hit Found									No Hit Found						
m142R m144R	6556265795 6609566400	78 102	8,803 12.245	7.15 7.97	No Hit Found									No Hit Found No Hit Found						
m145R	6645666686	77	8,012	4.24	No Hit Found									No Hit Found						
m146L	6706266835	76	8,599	10.65	No Hit Found									No Hit Found						
m147R	6758467784	67	7,447	11.07	No Hit Found									No Hit Found						
m151L	6899068790	67	7,942	10.38	No Hit Found									No Hit Found						
m153L	7006669116	317	35,140	6.06	No Hit Found								1 2 3 4 5 6 7 8	AAC39771 hepatitis A virus CAA66906 HAVcr-1 protein CAE74002 Hypothetical prot BAB03062 unnamed protein AAC39774 hepatitis A virus	icellular receptor 1 long form cellular receptor 1 short form the cellular receptor 1 short form cellular receptor 1 long form cellular receptor 1 short form	77.80 4.92E-13 58.92 2.37E-07 58.92 2.37E-07 58.15 4.03E-07 57.77 5.27E-07 56.61 1.17E-06 56.23 1.53E-06 56.23 1.53E-06 53.53 9.94E-06	29% 28% 28% 33% 22% 27% 27% 27% 26%	43% 36% 36% 41% 38% 38% 35% 35% 40%	9234 9234 108242 3253	149-404 152-356 147-351 147-305 202-452 868-1133 152-360 147-355 80-335
													10	XP_502320 hypothetical pro	tein	52.76 1.70E-05	26%	39%		311-532
m154R	6988970083	65	7,314	6.51	No Hit Found									No Hit Found						
m155L	7094570082	288	31,696	4.48	No Hit Found								1 2 3 4 5 6 7 8 9	AAC39774 hepatitis A virus	icellular receptor 1 long form cellular receptor 1 short form cellular receptor 1 long form cellular receptor 1 short form receptor receptor 1 short form receptor 1 short form receptor 1 short form product	82.80 1.31E-14 65.08 2.82E-09 65.08 2.82E-09 63.54 8.21E-09 59.69 1.19E-07 58.15 3.45E-07 55.54 1.71E-06 55.07 2.92E-06 51.60 3.23E-05	28% 28% 28% 29% 29% 29% 29% 22% 24% 26%	46% 37% 37% 36% 36% 36% 38% 39% 34% 39%	2288 31231 31231 31231 31231 27231 21285 4285 24269 2	154-332 149-327 149-314 137-305 286-535 198-499
m156L	7165970991	223	24,748	4.54	No Hit Found								1	XP_499823 hypothetical pro	tein	65.08 1.78E-09	25%	44%	2211	153409
m157L	7247872140		12,142	5.81	No Hit Found									No Hit Found						
m158L	7389773517		13,996	7.91	No Hit Found									No Hit Found						
m159L	7473673984		28,065	9.84	No Hit Found								1			117.86 2.89E-25	50%	54%	99231	14164
m161L m162l	7505874843 7528775090		8,381 7,833	9.45 8.34	No Hit Found No Hit Found									No Hit Found No Hit Found						
102L	. 020, -10000	00	,,,,,,,	0.04	140 FIRE FOUND															

Gene Name	Genome Position		Peptide Mw	pl CDD Numb		COG Definition	Bit Score E-value	% Posit	Query Hit from BLASTp W from-to to Number	Hit Accession	BLASTp Definition	Bit Score E-value	% % ntity Positi	Query	Hit from-
m164L	7555575361	length 65		11.52	No Hit Found			identity e	from-to to Number	No Hit Found		ide	nuty Fositi	ve mom-to	, 10
m165R	7544775647	67		11.06	No Hit Found					No Hit Found					
m168R	7700277304	101		8.08	No Hit Found					No Hit Found					
m169R	7705777332 7794778162	92 72	10,280 8,084	5.66 7.32	No Hit Found					No Hit Found  No Hit Found					
m172R m175R	7867978897	72		10.14	No Hit Found					No Hit Found					
m176R	7921379434	74		5.86	No Hit Found					No Hit Found					
m181R	8308983283	65		9.38	No Hit Found					No Hit Found					
m182R	8312083344	75	8,985	9.41	No Hit Found					No Hit Found					
m185R	8429884501	68	7,612	9.01	No Hit Found					No Hit Found					
m187L	8463784401	79	9,091	10.42	No Hit Found					No Hit Found					
m188R	8544285699	86		7.59	No Hit Found					No Hit Found					
m189L	8605685853	68		10.47	No Hit Found					No Hit Found					
m191R	8606286256	65		12.50	No Hit Found					No Hit Found					
m192L m193L	8651886288 8697886649	77		11.89 8.47	No Hit Found					No Hit Found  No Hit Found					
m193L m194R	8694487165	110 74		10.23	No Hit Found					No Hit Found					
m195L	8761187258	118		6.88	No Hit Found					No Hit Found					
m196L	8790887678	77		7.62	No Hit Found					No Hit Found					
m197L	8867588475	67		4.11	No Hit Found					No Hit Found					
m198L	8903588763	91	7,908	3.58	No Hit Found					No Hit Found					
m199R	8877789040	88	9,997	12.36	No Hit Found					No Hit Found					
m202L	9030590036	90	10,585	10.88	No Hit Found					No Hit Found					
m203L	9036190113	83	9,179	9.43	No Hit Found					No Hit Found					
m205R	9124391446	68		5.66	No Hit Found					No Hit Found					
m207L	9254991956		21,901	9.47	No Hit Found					No Hit Found					
m209R	9207692375	100		11.09	No Hit Found					No Hit Found					
m211R m212R	9272392977 9279993008	85 70		5.93 8.17	No Hit Found No Hit Found					No Hit Found  No Hit Found					
m214L	9321493002	71		11.22	No Hit Found					No Hit Found					
m216L	9419293989	68		11.32	No Hit Found					No Hit Found					
m217L	9472594324	134	14,250	11.13	No Hit Found					No Hit Found					
m220R	9528295596	105	11,594	10.62	No Hit Found					No Hit Found					
m222R	9579096113	108	11,755	11.02	No Hit Found				1	NP_048598 a249L		53.91 1.62E-06	44% 50	0% 1810	17102
m223R	9585296346	165	19,121	10.66	No Hit Found					No Hit Found					
m224L	9665196373	93		10.36	No Hit Found					No Hit Found					
m226R	9666997106	146		11.60	No Hit Found					No Hit Found					
m227R	9724297442	67		10.52	No Hit Found					No Hit Found					
m228L m229R	9801097807 9800498408	68 135		10.79	No Hit Found					No Hit Found  No Hit Found					
m230R	9854198750	70		5.63	No Hit Found					No Hit Found					
m231R	9864898857	70		9.09	No Hit Found					No Hit Found					
m234R	9950499701	66	7,387	11.07	No Hit Found				1	NP 048586 a238L		83.19 2.48E-15	66% 72	2% 26	6 48112
m235R	99846100055	70	7,784	9.64	No Hit Found					No Hit Found					
m236R	100367100780	138	15,833	8.18	No Hit Found					No Hit Found					
m239L	101646101323	108	12,058	8.36	No Hit Found					No Hit Found					
m244R	103086103475	130	14,802	9.99	No Hit Found				1	NP_048575 A227L		162.93 2.43E-39	60% 78	3% 312	9 10136
	103420103190	77		10.69	No Hit Found				1	NP_048576 a228R		72.79 3.36E-12	51% 62	2% 17	7 176
	103875-104153		10,394	11.82	No Hit Found					No Hit Found					
	104786-104517 104705-105112		9,825 15,210	11.81 8.24	No Hit Found No Hit Found					No Hit Found  No Hit Found					
	104705-105112		7,566	11.93	No Hit Found					No Hit Found					
	105328-105609		10,949	8.36	No Hit Found					No Hit Found					
	105987-105775		7,609	4.99	No Hit Found					No Hit Found					
	106227106427		7,671	9.01	No Hit Found					No Hit Found					
m257R	106575106811	79	8,744	10.82	No Hit Found					No Hit Found					
m259R	107614107814	67	7,648	12.22	No Hit Found					No Hit Found					
m260R	107861-108202	114	13,241	10.49	No Hit Found					No Hit Found					

Gene Name	Genome Position	A.A. length	Peptide Mw	pl	CDD Hit Number	COGs COG Definition	Bit Score E-val	lue   %	% Positiv	Query H	lit from BLASTp	Hit Accession	BLASTp Definition	Bit Score E-value	% entity Pos	% Quer	/ Hit from-
	108103-108387	95		8.52		Hit Found			е		Number	No Hit Found					
	108508108278			10.64		Hit Found						No Hit Found					
m265L	110071-109826	82	9,249	10.32	No I	Hit Found						No Hit Found					
m267L	110331-110119	71	7,953	4.56	No I	Hit Found						No Hit Found					
m268R	110517-110729	71	8,512	11.89	No I	Hit Found						No Hit Found					
m270R	111648111842	65	7,318	7.29	No I	Hit Found						No Hit Found					
m271L	111986-111708	93	11,395	10.66	No I	Hit Found						No Hit Found					
m273R	111941112186	82	9,539	9.91	No I	Hit Found						No Hit Found					
	112244-112468	75		6.40		Hit Found						No Hit Found					
	112616112897	94		11.93		Hit Found						No Hit Found					
	113591-113304		10,835	11.12		Hit Found						No Hit Found					
	113988-113377 113597-113959	204	22,758 13,681	8.55 12.56		Hit Found Hit Found						No Hit Found  No Hit Found					
	114363-114136	76		7.32		Hit Found						No Hit Found					
	114563-114369	65		12.23		Hit Found						No Hit Found					
	115387-115025			9.48		Hit Found						No Hit Found					
m285L	115368-115105	88	9,895	11.95	No I	Hit Found						No Hit Found					
m286L	115960-115682	93	10,741	10.24	No I	Hit Found						No Hit Found					
m287L	116160115906	85	9,533	10.91	No I	Hit Found						No Hit Found					
m290R	117315117527	71	8,140	8.46	No I	Hit Found						No Hit Found					
m291L	117744-117439	102	11,319	4.95	No I	Hit Found						No Hit Found					
m292L	117991-117611	127	14,385	8.73	No I	Hit Found						No Hit Found					
m293L	118314118069	82	9,088	10.16	No I	Hit Found						No Hit Found					
m295L	119073118456	206	23,413	10.50	No I	Hit Found						No Hit Found					
	119251-118901	117		8.81		Hit Found						No Hit Found					
	119390119163			4.83		Hit Found						No Hit Found					
	120203119898		11,861	9.48		Hit Found						No Hit Found					
	120447-120220 120462-121241	76	9,522 26,935	13.28 12.50		Hit Found Hit Found						No Hit Found					
	120771-120532	80		13.03		Hit Found						No Hit Found					
	120878-121216	113		6.46		Hit Found						No Hit Found					
	121304-121783		17,675	5.42		Hit Found						No Hit Found					
m309R	122771-123256	162		11.83		Hit Found						No Hit Found					
m311R	123590-123814	75	7,705	10.62	No I	Hit Found					1	NP_048551 a204	4L	84.34 1.12E-15	65%	67% 2-	74 678
m316R	125397-125696	100	11,765	9.99	No I	Hit Found						No Hit Found					
m318R	126300-126710	137	14,710	11.07	No I	Hit Found						No Hit Found					
m320R	126788-127180	131	13,962	4.67	No I	Hit Found						No Hit Found					
m321L	127242-127021	74	8,170	11.30	No I	Hit Found						No Hit Found					
	129162-128968	65		10.55		Hit Found						No Hit Found					
	129461-129204		10,035	10.29		Hit Found						No Hit Found					
	129967-130338	124		6.23		Hit Found						No Hit Found					
	130163-130543 131493-131143	127	14,296 13,038	10.96		Hit Found Hit Found						No Hit Found  No Hit Found					
	132326-131694		24,704	9.01		Hit Found					1		igi	241.89 9.33E-63	58%	77% 14-2	08 1197
	132317-132763	149		11.09		Hit Found						No Hit Found	.oc	241.00 0.002.00	0070		
	132869-133270			9.07		Hit Found						No Hit Found					
	133035133403			8.82		Hit Found						No Hit Found					
m340L	134014-133568	149	15,783	4.36	No I	Hit Found						No Hit Found					
m342L	134328-134092	79	8,598	9.54	No I	Hit Found						No Hit Found					
m343L	134957-134694	88	10,005	11.68	No I	Hit Found						No Hit Found					
m345L	136343-136149	65	7,482	9.44	No I	Hit Found						No Hit Found					
m347R	136321-136632	104	11,830	8.39	No I	Hit Found						No Hit Found					
	136790-137071		9,874	11.44		Hit Found						No Hit Found					
	137303-136971		12,566	12.11		Hit Found						No Hit Found					
	137080-137358		10,302	7.18		Hit Found						No Hit Found					
	137918-137451	156		10.73		Hit Found						No Hit Found					
m353L	138776138432	115	13,428	9.08	No I	Hit Found						No Hit Found					

Gene Name	Genome Position	length	Peptide Mw	pi N	DD Hit COGs	COG Definition	Bit Score	E-value	% Identity	% Positiv e	Query Hit from BLASTp from-to to Hit Number	Hit	Accession	BLASTp Definition	Bit Score E-value	% Identity F	% (	Query F rom-to	lit from- to
m355R		95		11.28	No Hit Found								No Hit Found						
	138988-139191	68		12.04	No Hit Found								No Hit Found						
	141751-140726 142845-143192		13,254	12.13	No Hit Found No Hit Found								No Hit Found						
	143505-143173		14,534	13.51	No Hit Found								No Hit Found						
						COG3889, Predicted solute binding protein [General function prediction	45.40	5 00F 00	0001	500	40 400 750 050								
	143202-143642		14,331	6.43	1 COG3889	only).	45.42	5.98E-06	30%	59%	12103 758850		No Hit Found						
	144406144146			9.57	No Hit Found								No Hit Found						
	144817-144425		14,771	7.46	No Hit Found								No Hit Found						
	145139-144936			8.09	No Hit Found								No Hit Found						
	147588-147247		12,771	11.00	No Hit Found														
	147919-147722 150262-149987	66 92		4.16 11.55	No Hit Found No Hit Found								No Hit Found  No Hit Found						
	150262-149987	69		8.67	No Hit Found								No Hit Found						
	151760-151497	88		11.31	No Hit Found								No Hit Found						
	152087-151890	66		8.17	No Hit Found								No Hit Found						
	152546152349	66		8.65	No Hit Found								No Hit Found						
	152797153084	96		8.04	No Hit Found								No Hit Found						
	153385154275		33,023	6.53	No Hit Found						1	1	NP_048635 a281R		114.01 5.60E-24	50%	55%	124277	14164
	155619-155380			5.13	No Hit Found								No Hit Found						
m391L	156030155731	100	11,267	6.48	No Hit Found								No Hit Found						
m392L	156173155943	77	8,749	9.47	No Hit Found								No Hit Found						
m393R	155957-156175	73	8,628	10.76	No Hit Found								No Hit Found						
m394L	156620-156312	103	11,566	7.34	No Hit Found								No Hit Found						
m396R	156675156941	89	10,042	11.44	No Hit Found								No Hit Found						
m398R	157206157541	112	12,842	11.36	No Hit Found								No Hit Found						
m400L	158861-158496	122	13,544	10.38	No Hit Found								No Hit Found						
m402L	159108158845	88	10,297	10.48	No Hit Found								No Hit Found						
m405L	160671-160357	105	11,781	11.79	No Hit Found						1	1	NP_048803 a446R		56.61 2.52E-07	60%	66%	55105	2-52
m406L	161079-160885	65	7,660	11.13	No Hit Found								No Hit Found						
m408L	162340-161600	247	23,928	4.24	1 pfam05642	Sporozoite_P67, Sporozoite P67 surface antigen. This family consists of several Theileria P67 surface antigens. A stage specific surface antigen of Theileria parva, p67, is the basis for the development of an antisorozoite vaccine for the control of Ea	35.54	4.72E-03	39%	44%	106194 185275		No Hit Found						
m410L	163253-163038	72	8,079	9.69	No Hit Found								No Hit Found						
m412R	163120-163332	71	8,024	6.34	No Hit Found								No Hit Found						
m414R	163833164213	127	13,143	9.29	No Hit Found						1 2 3	2	ZP_00372892 hypothetical protein BAE63414 unnamed protein p ZP 00372930 hypothetical protein	roduct	64.31 1.19E-09 59.31 3.82E-08 59.31 3.82E-08	42% 40% 39%	49%	19127 3127 11127	15122 38166 7122
m415R	163984164427	148	16,959	11.95	No Hit Found								No Hit Found						
	165741-166007	89		4.43	No Hit Found								No Hit Found						
	166383166703	107		8.68	No Hit Found								No Hit Found						
	166897167292	132		12.08	No Hit Found								No Hit Found						
	168066167671	132		10.56	No Hit Found								No Hit Found						
	168014167781	78		11.94	No Hit Found								No Hit Found						
	168393-168199			4.91	No Hit Found								No Hit Found						
	168933-168712			7.65	No Hit Found								No Hit Found						
	169269-169072 169803-170045	66 81		7.97 11.40	No Hit Found								No Hit Found						
	170094-170339			10.25	No Hit Found								No Hit Found						
	170566-170781			12.61	No Hit Found								No Hit Found						
	170706-171065		13,910	5.86	No Hit Found								No Hit Found						
	170959-171360		15,335	10.45	No Hit Found						1	1	NP_048778 A421R		86.66 2.25F-16	53%	68%	60128	2391
	171801-171595			10.60	No Hit Found						'		No Hit Found		22.23	30,0	/-		
	172878-172141		28,123	8.64	No Hit Found								No Hit Found						
	172616-172221		15,721	10.86	No Hit Found								No Hit Found						
	172973-173236			10.35	No Hit Found								No Hit Found						
	173255173524		10,727	9.39	No Hit Found								No Hit Found						
	173707-173970			10.11	No Hit Found								No Hit Found						
	174059-174325		10,279	10.83	No Hit Found								No Hit Found						
m453R	175006175218	71	8,470	8.55	No Hit Found								No Hit Found						

Gene Name	Genome Position	A.A. length	Peptide Mw	pl	CDD Hit Number	COGs	COG Definition	Bit Score E-valu	e %	% ity Positiv	Query from-to	Hit from	BLASTP Hit Number	Hit Accession	BLASTp Definition	Bit Score E-value	% Identity I	% ( Positive fi	Query H	lit from- to
m456L	176463176071	131	15,409	11.10		No Hit Found				-				No Hit Found						
m458L	176799-176533	89	10,130	10.99		No Hit Found								No Hit Found						
m459L	176915176643	91	10,412	7.81		No Hit Found								No Hit Found						
m461L	177809-177327	161	18,843	10.84		No Hit Found								No Hit Found						
m462L	177892-177671	74	8,117	4.22		No Hit Found								No Hit Found						
m464R	179187179435	83	9,781	9.42		No Hit Found								No Hit Found						
m466R	180176180388	71	8,158	9.73		No Hit Found								No Hit Found						
m468L	181119180901	73	8,736	10.25		No Hit Found								No Hit Found						
m469L	181264-180977	96	10,878	11.76		No Hit Found								No Hit Found						
m470L	181909181709	67	7,565	10.71		No Hit Found								No Hit Found						
m471L	183057-182830	76	8,083	11.66		No Hit Found								No Hit Found						
m473L	183499183272	76	8,279	7.86		No Hit Found								No Hit Found						
m474L	183918-183592	109	12,268	10.19		No Hit Found								No Hit Found						
m477R	184604184816	71	7,675	10.57		No Hit Found								No Hit Found						
m480L	185780-185550	77	8,473	11.31		No Hit Found								No Hit Found						
m481L	185980185774	69	8,077	11.91		No Hit Found								No Hit Found						
m483R	186328186606	93	10,660	11.26		No Hit Found								No Hit Found						
m486L	187726187436	97	10,790	5.04		No Hit Found								No Hit Found						
m487R	187873—188391	173	20,718	4.10		No Hit Found							1 2 3 4 5 6 7 8 9	ZP_00054046 XP 344436 AAB27498 XP_75005 CAF92931 AAA91036 XP_584941 XP_875111	Insonhetical protein Breal 10 (10.04924 COGGS210 Large exproteins involved in heme utilization or adhesion PREDICTED: similar to Hypothetical protein CBG20540 PREDICTED: similar to transmembrane protease, serine 2 unamed protein product Wi-1 adhesion PREDICTED: similar to Epsin 1 isoform 1 PREDICTED: similar to Epsin 1 isoform 1 PREDICTED: similar to Epsin 1 isoform 7 PREDICTED: similar to Epsin 1 isoform 7	59.31 5.32E-08 57.77 1.55E-07 56.23 4.50E-07 53.91 2.24E-06 51.99 8.49E-06 50.45 2.47E-05 48.52 9.39E-05 48.52 9.39E-05 48.52 9.39E-05	30% 31% 24% 28% 29% 41% 21% 34% 34% 33%	38% 42% 49% 43% 35% 45% 35% 37% 37% 37% 38%	16-122 28-134 22-133 43-119 24-140 29-131 29-131	1115 843966 260336 556667 278388
m489R	187941-188480	180	19,136	8.04		No Hit Found							1 2 3 4 5 6 7 8 9	NP_703149 AAB16964 XP_421396 ZP_00283849 XP_780848 ZP_0067800 BAE24926 AAF34245	Extraembryonic, spermatogenesis, homeobox 1-like extraembryonic, spermatogenesis, homeobox 1-like collagen-like protein generate protein spermatogenesis, homeobox 1-like collagen-like protein generate protein with a bacterial SH3 domain homeoboue precipion initial to Anyl hydrocarbon receptor precursor (Al hypothetical protein TeryDRAFT 0036 unnamed protein product juntament protein teryDRAFT 0036 unnamed protein product juntament juntame	68.94 7.44E-11	45% 45% 29% 40% 37% 45% 35% 36% 36% 36%	52% 53% 54% 53% 53% 53% 55% 42% 53% 53% 53%	7102 4102 796 2100 594 2102 7102	140-238 504-594 155-253 381-481 329-429 277-372 277-372
m490L	188607188398	70	8,412	11.87		No Hit Found								No Hit Found						
m492L	189190-188975	72	8,399	12.17		No Hit Found								No Hit Found						
m493L	189269189006	88	9,350	3.65		No Hit Found								No Hit Found						
m494L	189635189432	68	7,425	8.43		No Hit Found								No Hit Found						
m498L	191788191483	102	12,357	9.89		No Hit Found								No Hit Found						
m499R	191600-191800	67	7,016	7.26		No Hit Found								No Hit Found						
m500L	191928-191734	65	6,824	9.68		No Hit Found								No Hit Found						
m503R	192663192923	87	9,505	7.85		No Hit Found								No Hit Found						
m505R	193511-193777	89	10,512	6.76		No Hit Found								No Hit Found						
m509L	195125-194832	98	10,361	9.29		No Hit Found								No Hit Found						
m510L	195823-195557	89	10,289	8.82		No Hit Found								No Hit Found						
	196352-196678			11.25		No Hit Found								No Hit Found						
m514L	197128196385	248	28,609	5.32		No Hit Found							1	NP_048893	A537L	107.84 2.92E-22	30%	46%	4248	10263
	197407-197183	75	8,403	6.78		No Hit Found								No Hit Found						
m517R	197246197749	168	19,026	10.39		No Hit Found							1 2	NP_048895 NP_048482	A539R similar to bacteriophage T4 intron-associated endonuclease, correspond to Swiss-Prot Accession Number P13299	177.95 9.53E-44 s 50.45 2.29E-05	50% 53%	70% 68%	1168 2874	1173 1155
m519L	198025-197813	71	8,027	12.02		No Hit Found								No Hit Found						
m520R	199333199668	112	12,587	10.19		No Hit Found								No Hit Found						
m521L	201067-200873	65	7,479	11.17		No Hit Found								No Hit Found						
m523L	202412-202161	84	9,653	8.46		No Hit Found								No Hit Found						
m524R	203196-203426	77	9,149	8.12		No Hit Found								No Hit Found						
m527R	205326205580	85	9,179	5.59		No Hit Found							1	NP_048897	a541R	49.68 3.08E-05	40%	48%	179	1086
m528R	206769-207014	82	9,367	4.03		No Hit Found								No Hit Found						
m529R	207508-207714	69	8,312	11.00		No Hit Found								No Hit Found						
m532R	208354208560	69	7,721	7.26		No Hit Found								No Hit Found						
m533R	208901-209155	85	9,403	7.89		No Hit Found								No Hit Found						
m534R	209205209474	90	10,036	4.17		No Hit Found								No Hit Found						

Gene Name	Genome Position	A.A. length	Peptide Mw	pl	CDD Hit Number	COGs	COG Definition B	Bit Score E-value	% Identity	% Positiv	Query Hit from from-to to	BLASTp Hit Number	Hit Ac	ccession BLASTp Definition	Bit Score E-value	% Identity	% Positive	Query from-to	Hit from- to
m536R	209245209604	120	13,509	11.01	N	lo Hit Found				·		· · · · · · · · · · · · · · · · · · ·	N	No Hit Found					
m537R	209650210063	138	14,807	10.13	N	No Hit Found						1	)	XP_719031 hypothetical protein CaO19_6069	51.22 1.03E-05	47%	57%	73138	171
m538R	210184210414	77	8,142	9.88	N	No Hit Found							No	No Hit Found					
m539R	210439-210873	145	16,355	7.46	N	No Hit Found							No	No Hit Found					
m540L	211104210829	92	10,460	8.25	N	No Hit Found							Ne	No Hit Found					
m541R	210931-211455	175	19,176	10.91	N	No Hit Found						1	ZP	P_00886604 hypothetical protein CsacDRAFT 2497	51.22 1.50E-05	36%	45%	73169	5112
m542R	211603-211881	93	10,213	10.82	N	No Hit Found							No	No Hit Found					
m544L	212294212097	66	7,409	10.36	N	No Hit Found							No	No Hit Found					
m545L	212862-212653	70	8,227	7.74	N	No Hit Found							No	lo Hit Found					
m547R	213534213743	70	8,062	8.84	N	No Hit Found							Ne	lo Hit Found					
m548L	214314213919	132	14,428	12.46	N	No Hit Found							Ne	lo Hit Found					
m549L	214851-214564	96	10,868	10.35	N	No Hit Found						1	1	NP_048942 a586R	73.94 1.52E-12	60%	68%	164	1275
m550L	215851-214862	330	36,801	7.75	N	o Hit Found							No	lo Hit Found					
m551L	215751-215542	70	7,776	12.41	N	o Hit Found						1	1	NP 048940 contains prenyl group binding site (CAAX) box	78.95 4.62E-14	58%	70%	170	2998
m552L	216399-215962	146	17,293	9.37	N	o Hit Found							Ne	lo Hit Found					
m554L	216622-216275	116	13,759	9.68	N	o Hit Found							No	No Hit Found					
m555L	216794216498	99	11,134	9.37	N	o Hit Found							No	No Hit Found					
m558R	218073218720	216	28,188	11.65	N	No Hit Found						1		CAE56520 Hypothetical protein CBG24243 XP_509745 PREDICTED: similar to Coagulation factor VII (Serum prothrombin	71.25 2.31E-11	41%	53%		37166
												2		XP_509745 PREDICTES animal to coagulation factor vii (Serum production)  AAX27666 unknown	63.93 3.68E-09 62.39 1.07E-08	25% 35%	50% 35%	112177 3 87184	39804045 14111
												4 5		AAB01097 glycine-rich protein XP 364198 hypothetical protein MG09043.4	60.85 3.12E-08 56.61 5.88E-07	43% 28%	48% 40%	91172 96200	2099 370472
												6 7		CAE57148 Hypothetical protein CBG25081 EAL26857 GA19788-PA	56.23 7.68E-07 53.91 3.81E-06	36% 46%	44% 53%		206342 421503
												8 9		ABA47855 hypothetical protein BURPS1710b 1103 BAC86188 unnamed protein product	53.14 6.50E-06 52.76 8.49E-06	40% 30%	41% 32%	105181 139213	58139 438533
												10		CAE73786 Hypothetical protein CBG21336	51.60 1.89E-05	33%	40%	87210	90164
m559R	218182-218796	205	16,941	3.64	N	lo Hit Found							No	lo Hit Found					
	218995219360		14,412	12.56		lo Hit Found								lo Hit Found					
	219308219763		17,062	7.97		No Hit Found								No Hit Found					
	220197-220544	116		7.38		lo Hit Found								lo Hit Found					
	220465220707	81		6.41		lo Hit Found								lo Hit Found					
	220749-221102	118		5.13		to Hit Found								No Hit Found					
	222557-222784	76		11.01		lo Hit Found								No Hit Found					
	222990-223187	66		7.82		lo Hit Found								No Hit Found					
	223021-223305		10,472	8.56		No Hit Found								lo Hit Found					
	224482-224252	77		8.84		No Hit Found								lo Hit Found					
	224324-224605		11,130	9.55		No Hit Found								lo Hit Found					
	224481-224771		11,724	10.66		No Hit Found								No Hit Found					
	225246-224911		12,616	4.54		No Hit Found						1		NP 048751 A394R	59.69 2.90E-08	50%	67%	1771	63116
	225310-225513 225640-225969	68		7.29		No Hit Found								No Hit Found					
	225711-225953	81	12,466 9,833	7.60		to Hit Found								No Hit Found					
	225961-225755	69		11.74		to Hit Found								No Hit Found					
	226414-226163	84		8.50		to Hit Found								No Hit Found					
							Lipoprotein_14, Actinobacillus constitutively-expressed outer membrane												
m592L	226406-226182	75	7,909	11.73	1 pf	fam03346	lipoprotein A.	39.74 2.82E-04	33%	48%	256 64118	1 2		XP_536513 PREDICTED: similar to AHNAK nucleoprotein isoform 1  AAN60242 unknown	64.70 9.22E-10 60.85 1.33E-08	72% 60%	74% 82%	2167 2267	237283 71116
												3	1	NP_849949 lipid binding NP_046976 tail protein	60.85 1.33E-08 55.84 4.28E-07	60% 40%	82% 65%	2267 2171	91136 639690
												5	,	XP 365380 predicted protein  XP_343560 PREDICTED: similar to LAF-4 protein (Lymphoid nuclear protein related	53.91 1.63E-06 50.06 2.35E-05	38%	76%	2167	3177 17351778
												7		XP_926450 PREDICTED: hypothetical protein XP 921357	49.68 3.07E-05	52% 50%	77% 70%		13317/8
												8		EAL47534 hypothetical protein 100.t00015	48.14 8.93E-05	50%	61%	2366	67110
	226879-226571					lo Hit Found								lo Hit Found					
	227162-226689		16,876	7.39		No Hit Found						1		NP_048746 a389R	57.77 1.18E-07	52%	53%	55118	264
	226710226958		9,668	10.72		No Hit Found								lo Hit Found					
	227504-227190		10,901	11.09		No Hit Found								No Hit Found			_		
m597R	227549-227782	78	9,371	12.11	N	lo Hit Found						1 2	1	CAE73829 Hypothetical protein CBG21394 NP_033126 repetin	59.69 2.94E-08 56.61 2.49E-07	28% 54%	77% 58%	1672	337417
												3 4		CAE62907 Hypothetical protein CBG07098  XP_584748 PREDICTED: similar to concentrative Na+-nucleoside cotransporter	52.37 4.69E-06 51.60 8.00E-06	27%	71% 53%	369 371	2292 568640
												5 6		XP_136142 PREDICTED: similar to periphilin 1 isoform 1  EAL47849 fimbriae-associated protein, putative	51.60 8.00E-06 48.91 5.18E-05	35% 28%	67% 57%	270	248306 743815
												7 8		AAW31832 TraC XP_918493 PREDICTED: similar to periphilin 1 isoform 1	48.52 6.77E-05 48.14 8.84E-05		54% 64%	271 1169	551621 256314
m598L	228408-228196	71	8,177	6.37	N	No Hit Found							Ne	lo Hit Found					
m600L	229103-228675	143	13,338	4.46	N	lo Hit Found							No	lo Hit Found					

Gene Genome Name Position	A.A. Peptide length Mw	pl CDD Hit Number		COG Definition	Bit Score E-value	% Positiv	Query Hit from BLAST from-to to Numbe	Hit	Accession	BLASTp Definition	Bit Score E-value	% Identity	% Positive	Query from-to	Hit from- to
m602R 229665-229874	70 7,777	4.71	No Hit Found			. 6	Numbe	ər	No Hit Found						
m603R 229877-230146	90 10,142	9.02	No Hit Found						No Hit Found						
m604R 230175-230435 m608R 231258-231482	87 10,144	10.03	No Hit Found						No Hit Found						
m608R 231258-231482 m609L 232244-231798	75 8,553 149 16,808	11.18	No Hit Found						No Hit Found  No Hit Found						
m611L 233193-232747	149 10,006	9.49	No Hit Found  No Hit Found						No Hit Found						
m613L 233551-233306	82 9,898	10.64	No Hit Found						No Hit Found						
m614L 234221-233913	103 10,728	9.55	No Hit Found						No Hit Found						
m617R 234588234998	137 12,172	10.48	No Hit Found						No Hit Found						
m619L 235658235443	72 8,060	9.59	No Hit Found						No Hit Found						
m621R 236035-236253	73 7,561	12.50	No Hit Found						No Hit Found						
m623R 236877-237314	146 16,741	10.23	No Hit Found						No Hit Found						
m624L 237337-237107	77 9,004	11.30	No Hit Found						No Hit Found						
m626L 238229-237798	144 16,890	11.00	No Hit Found					1	NP_048841 A485R		176.41 2.14E-43	57%	78%	4144	7147
m630R 239412-239957 m632R 240321-240938	182 20,851 206 24,644	10.12	No Hit Found No Hit Found						No Hit Found  No Hit Found						
m633R 240856-241290	145 16.724	9.34	No Hit Found					1	NP_048635_a281R		54.30 1.22E-06	68%	73%	73112	1050
m636L 242219-241983	79 8,978	8.34	No Hit Found						No Hit Found		04.00	0070	7070	70 112	10 00
m639R 243064-243258	65 7,680	6.87	No Hit Found						No Hit Found						
m640L 243558-243079	160 16,920	4.28	No Hit Found						No Hit Found						
m641L 243676243473	68 8,028	8.83	No Hit Found						No Hit Found						
m642L 243786243568	73 7,563	8.37	No Hit Found						No Hit Found						
m643L 243970-243713	86 9,840	6.71	No Hit Found						No Hit Found						
m644L 244353-244102	84 8,333	4.17	No Hit Found						No Hit Found						
m645R 244473-244868	132 15,774	11.24	No Hit Found						No Hit Found						
m646L 244759-244535	75 9,037	11.43	No Hit Found						No Hit Found						
m648L 245910245242	223 24,608	4.73	No Hit Found					1	NP 048635 a281R		58.54 1.66E-07	44%	47%	140217	70164
m650L 246234-246025	70 7,650	9.58	No Hit Found						No Hit Found						
m652L 247565-247158	136 15,259	10.07	No Hit Found						No Hit Found		07.00 0.045.40	45%	53%	4136	1133
m654R 247737-248300	188 21,003	10.58	No Hit Found					2 3 4 5	CAG98466 unnamed protein pr AAL79317 unknown NP_851581 CPXV051A protein NP_063682 ribonucleoside-diph AAW23723 ribonucleoside-diph	osphate reductase	97.06 2.84E-19 54.68 1.62E-06 50.83 2.34E-05 48.91 8.87E-05 48.91 8.87E-05	45% 41% 53% 53% 53%	51% 63% 65% 65%	4136 67136 85131 85127 85127	2-71 31-77 31-73 31-73
m655L 248307248044	88 9,770	12.20	No Hit Found						No Hit Found						
m656R 248276-248581	102 11,361	5.51	No Hit Found						No Hit Found						
m658R 248739-249017	93 10,389	10.78	No Hit Found						No Hit Found						
m659L 249484249281	68 7,714	12.51	No Hit Found						No Hit Found						
m661R 249468-249791	108 11,503	6.73	No Hit Found						No Hit Found						
m662R 249604-249804	67 7,526	12.17	No Hit Found						No Hit Found						
m663L 250168-249914	85 9,431	10.81	No Hit Found						No Hit Found						
m665R 250297-250515 m666R 250307-250546	73 8,493	8.56 4.61	No Hit Found No Hit Found						No Hit Found  No Hit Found						
m668L 252326-251328	80 8,886 333 37,720	8.88	No Hit Found						No Hit Found						
m669L 252400-252110	97 10.877	10.55	No Hit Found						No Hit Found						
m671L 252679252404	92 10,622	12.29	No Hit Found					1	NP_048822 a466L		67.40 1.39E-10	52%	58%	2491	1986
m673R 252619252858	80 8,851	8.52	No Hit Found						No Hit Found						
m675L 254829254323	169 19,043	10.45	No Hit Found					1	NP 048817 a460R		55.07 9.28E-07	31%	60%	67142	2-77
m676L 254668-254471	66 7,208	6.65	No Hit Found						No Hit Found						
m677L 255539254910	210 23,333	10.44	No Hit Found					1 2	NP_048816 a459R NP_048814 a457R		105.92 8.03E-22 75.87 8.89E-13	72% 60%	78% 64%	131205 1488	175 276
m678R 255023-255283	87 9,356	9.89	No Hit Found					-	No Hit Found		70.07 0.082-13	00.70	J*# /0	00	2 70
m679R 255756-256115	120 13,417	10.35	No Hit Found						No Hit Found						
m681L 256491-256267	75 8,246	7.95	No Hit Found						No Hit Found						
m682L 256816-256610	69 7,908	7.25	No Hit Found						No Hit Found						
m684R 256735-257034	100 11,393	7.09	No Hit Found						No Hit Found						
m685R 257437-257700	88 9,272	7.06	No Hit Found						No Hit Found						
m687L 257723-257484	80 8,779	12.52	No Hit Found						No Hit Found						
m688R 257588-257965	126 12,965	4.66	No Hit Found						No Hit Found						
m689L 259013-258789	75 8,358	8.85	No Hit Found						No Hit Found						

Gen Nam	e Positio	n lei	ngth	Peptide Mw	pl CDD H Numbe	er COGS	COG Definition	Bit Score E-val	lue   %	ity Positiv	from to	LASTp Hit lumber	Hit Accession	BLASTp Definition	Bit Score E-value	% Identity I	% Positive	Query from-to	Hit from- to
	91L 259232-25 92L 260188-25		72	7,860 26.392	9.69 7.64	No Hit Found						1	No Hit Found NP 048924 A	1032 / 1	74.71 2.29E-12	27%	53%	41194	29173
	96L 261340-26		104	11,865	12.43	No Hit Found							No Hit Found	SOUL	74.71 2.250-12	2170	3370	41-104	25-173
m69	8R 261490-26	1708	73	8,055	9.38	No Hit Found							No Hit Found						
m70	2R 263252-26	3500	83	10,082	11.72	No Hit Found							No Hit Found						
m70	3L 263952-26	3533	140	15,929	7.34	No Hit Found							No Hit Found						
	05L 265052-26		114		11.86	No Hit Found							No Hit Found						
	7R 264862-26		446		5.21	No Hit Found						1	NP 048861 A	A505L	456.45 8.38E-127	56%	72%	48443	73478
	08L 26540326 09L 26569426		92 78		10.78 9.47	No Hit Found No Hit Found							No Hit Found						
	OR 265685-26		69	7,834	12.50	No Hit Found							No Hit Found						
	I1L 266105-26		127		12.13	No Hit Found						1	NP_048862 a	1506R	85.11 6.51E-16	62%	70%	38107	1-70
	2R 265925-26		133		10.72	No Hit Found							No Hit Found						
m71	3R 26628626	017	244	27,642	5.49	No Hit Found						1	NP_048859 c	contains prokaryotic membrane lipoprotein lipid attachment site	183.34 5.29E-45	37%	62%	6243	38297
m71	IGL 267588-26	304	95	10,824	10.44	No Hit Found							No Hit Found						
m71	8R 268194-26	3400	69	7,789	10.21	No Hit Found							No Hit Found						
m72	OR 269695-27	0036	114	13,297	8.26	No Hit Found						1	NP_048573 a	1225L	48.91 5.26E-05	39%	69%	62114	2-54
	22L 27031427		79		8.21	No Hit Found							No Hit Found						
	3R 270224-27		84	9,947	11.87	No Hit Found						1	NP_048572 a		82.80 3.29E-15	55%	61%	182	6-88
	4R 270551-27		75 169	8,091 19,093	11.61	No Hit Found						1	NP_048571 a	nypothetical protein a221L - Chlorella virus PBCV-1	59.31 3.87E-08 50.83 1.75E-05	41% 37%	48% 64%	275 66124	1992 157
	6R 271199-27		87		6.78	No Hit Found						'	No Hit Found	rypotnetical protein azz IL - Cilioretta virus PBCV-1	50.63 1.75E-05	3/76	0470	00124	1-57
	8R 271756-27		67	7,785	11.50	No Hit Found							No Hit Found						
m73	OR 272215-27	2433	73		3.99	No Hit Found							No Hit Found						
m73	1R 272950-27	3153	68	7,677	12.13	No Hit Found							No Hit Found						
m73	32L 273939-27	3745	65	7,597	6.35	No Hit Found							No Hit Found						
m73	3R 273860-27	1807	316	37,047	9.72	No Hit Found						1		imilar to Variola virus orf E10L, corresponds to Swiss-Prot Accession Number P33801		48%	69%	4315	6318
												2	YP_142754 S	S/T protein kinase, similar to Paramecium bursaria chlorella virus 1	60.85 6.19E-08	24%	42%	7245	105358
				7.000	40.04	N. 105 F						3		hypothetical protein Y73B6A.1	51.22 4.91E-05	23%	42%	4219	167375
	36L 275201-27 7R 275006-27		69 68	7,983 7,204	12.91	No Hit Found No Hit Found							No Hit Found No Hit Found						
	39L 276077-27		95		10.79	No Hit Found							No Hit Found						
	I3L 277075-27		84	9,212	11.75	No Hit Found							No Hit Found						
m74	I4L 277130-27	852	93	10,647	5.56	No Hit Found							No Hit Found						
m74	I5L 277792-27	7586	69	7,813	12.42	No Hit Found							No Hit Found						
m74	16L 278098-27	805	98	10,800	9.60	No Hit Found							No Hit Found						
	17L 279611-27		70	7,840	11.37	No Hit Found							No Hit Found						
	9R 279997-28		159		10.18	No Hit Found							No Hit Found						
	OR 281024-28		84		11.31	No Hit Found							No Hit Found						
	1R 281164-28 4R 281720-28		102		6.99	No Hit Found No Hit Found							No Hit Found No Hit Found						
	55L 282855-28		86	9.347	9.33	No Hit Found							No Hit Found						
	59L 284429-28			13,717	8.20	No Hit Found							No Hit Found						
	60L 284676-28			13,329	7.92	No Hit Found							No Hit Found						
m76	1R 284444-28	5031	196	21,931	11.13	No Hit Found							No Hit Found						
m76	4R 285340-28	687	116	12,055	11.37	No Hit Found							No Hit Found						
m76	5R 286458-28	760	101	11,498	8.81	No Hit Found							No Hit Found						
	7R 286794-28		121	13,630	10.82	No Hit Found							No Hit Found						
	88L 287392-28			8,796	9.11	No Hit Found							No Hit Found						
	9R 287562-28		70		10.27	No Hit Found							No Hit Found						
	1R 287942-28 72L 288195-28		134	15,317 7.467	10.40 7.04	No Hit Found							No Hit Found						
	2L 288195-28 4R 288510-28			7,648	9.97	No Hit Found							No Hit Found						
	75L 288923-28			13,206	9.29	No Hit Found						1	NP_048989 A	A633R	157.53 1.03E-37	61%	80%	1114	2115
m77	76L 288978-28	3706		10,642	9.47	No Hit Found							No Hit Found						
m77	8R 28911328	316	68		9.64	No Hit Found							No Hit Found						
m77	9R 289217-29	179	321	35,384	7.69	No Hit Found							No Hit Found						
m78	OR 289629-28	9868	80	8,856	11.97	No Hit Found							No Hit Found						

MINIT   2004-2-1010   70	Gene Genome Name Position	A.A. length	Peptide Mw		D Hit COGs	COG Definition	Bit Score E-value	% Positiv e	Query Hit from BLASTp from-to to BLASTp Hit Number	Hit Accession	BLASTp Definition	Bit Score E-va	lue Id	% entity Po	% ositive f		lit from- to
18-11   18-12   18-1	m781R 2903152906	39 125	13,025	10.01	No Hit Found				1	NP_048987 a631L		51.22 1.05	-05	41%	62%	64125	465
Part	m782R 2908432910	52 70	7,387	4.20	No Hit Found					No Hit Found							
Marie Mari	m783R 2910392912	90 84	9,703	8.84	No Hit Found					No Hit Found							
Marie   Mari	m784R 291512-2919	31 150	16,785	9.69	No Hit Found					No Hit Found							
Profile   2008-1-2007   1916	m786L 2918342915	39 82	9,155	12.29	No Hit Found					No Hit Found							
Month   Mont	m787R 2925392927	72 78	8,637	11.76	No Hit Found					No Hit Found							
Model   Mode	m788L 2929282927	16 71	8,080	11.04	No Hit Found					No Hit Found							
Marie   Mari	m790R 2930752933	05 77	9,084	10.37	No Hit Found					No Hit Found							
Mail	m792L 2938042933	32 141	15,179	11.22	No Hit Found					No Hit Found							
Model   Paralle   Model   Pa	m793L 2944702940	30 137	15,030	8.52	No Hit Found					No Hit Found							
M78E   286148-296814   11   12.491   8.64   Mol Nel Found			23,601		No Hit Found												
Model   1868																	
MRINEL 2009008-20094089   142   16,811   0.76   No HR Found   No HR Fo	m798L 2961462958	14 111	12,491	8.64	No Hit Found												
MRIOUR   207748   2																	
MeBURA   287846-288189   108   12,010   7.43   No Hil Found   No																	
M805R   208684 - 208743 - 2089715   91   7,840																	
M808   28874-299015   91   7,840   4.11   No Hit Found   No Hit																	
M808																	
M80El   30075-29881   65   7,514   12 15   No Hit Found   No Hit																	
m812R         300887-301033         149         17,652         9.95         No Hit Found           m815R         302491-302888         66         7,546         10.94         No Hit Found           m816R         302543-3022770         76         8,360         11.88         No Hit Found           m821.         304786-304820         89         10,147         9.99         No Hit Found           m828.         305535-305107         143         13,300         8.84         No Hit Found           m824.         305839-305423         139         16,292         8.84         No Hit Found           m825.         306579-308381         73         8,114         8.84         No Hit Found           m826.         307347-308856         164         19,181         9.83         No Hit Found           m827.         307665-307446         70         7,766         6.76         No Hit Found           m831R         308002-308882         87         10,140         9.67         No Hit Found																	
m815R         302491-302888         66         7,546         10,94         No Hit Found           m816R         302543-302770         76         8,880         11,88         No Hit Found           m821L         304786-304820         89         10,147         9,99         No Hit Found           m823L         30533-305107         143         13,300         8,84         No Hit Found           m824L         305839-305423         139         16,292         8,64         No Hit Found           m825L         306579-303810         73         8,114         8,84         No Hit Found           m826L         307347-308856         164         19,181         9,63         No Hit Found           m827L         307655-307446         70         7,766         6,76         No Hit Found           m831R         30802-308809         95         10,948         11,00         No Hit Found           m831R         30802-308802         87         10,140         9,67         No Hit Found																	
m816R         302543-302770         76         8.880         11.88         No Hit Found           m821L         304786-304520         89         10,147         9.99         No Hit Found           m823L         305538-305107         143         13,300         8.84         No Hit Found           m824L         305839-305423         139         16,292         8.64         No Hit Found           m825L         306579-306361         73         8,114         8.84         No Hit Found           m826L         307347-308856         164         19,181         9.83         No Hit Found           m827L         307865-307446         70         7,766         6.76         No Hit Found           m831R         30802-308802         87         10,140         9.67         No Hit Found																	
m821L         304786-304520         89         10,147         9.99         No Hit Found           m823L         30538-305107         143         13,300         8.84         No Hit Found           m824L         305839-305423         139         16,292         8.64         No Hit Found           m825L         306579-306361         73         8,114         8.84         No Hit Found         No Hit Found           m826L         307347-308856         164         19,181         9.83         No Hit Found         No Hit Found           m831R         30802-308809         95         10,948         11,00         No Hit Found         No Hit Found           m831R         30802-308862         87         10,140         9.67         No Hit Found         No Hit Found																	
m823L         305535-305107         143         13,300         8.84         No Hit Found           m824L         305839-305423         139         16,292         8.64         No Hit Found           m825L         306579-305301         73         8,114         8.84         No Hit Found         No Hit Found           m826L         307347-308856         164         19,181         9.83         No Hit Found         No Hit Found           m827L         307655-307446         70         7,766         6,76         No Hit Found         No Hit Found           m831R         30802-308862         87         10,140         9,67         No Hit Found																	
m824L 305839-305423       139 16,292 8,64 No Hit Found       No Hit Found         m825L 306579-305861       73 8,114 8,84 No Hit Found       1 NP_048675 a319L       64.31 1,21E-09 48% 54% 1-         m826L 307347-306856       164 19,181 9,83 No Hit Found       No Hit Found         m827L 307855-307446       70 7,766 8,76 No Hit Found       No Hit Found         m831R 30802-308880       95 10,948 11,00 No Hit Found       No Hit Found         m831R 30802-308882       87 10,140 9,67 No Hit Found       No Hit Found																	
m825L 306579-306361 73 8,114 8.84 No Hit Found 1 NP_048675 a319L 64.31 1.21E-09 48% 54% 1- m826L 307347-308856 164 19,181 9.83 No Hit Found  m827L 307655-307446 70 7,766 6.76 No Hit Found  m830R 308525-308809 95 10,948 11.00 No Hit Found  m831R 308002-308862 87 10,140 9.67 No Hit Found  No Hit Found																	
m828L 307347_308856 164 19,181 9.63 No Hit Found No Hit Found m827L 307655_307446 70 7.766 6.76 No Hit Found m830R 30825_308809 95 10,948 11.00 No Hit Found m831R 308002_308862 87 10,140 9.67 No Hit Found No Hit Found									1			64.31 1.21	-09	48%	54%	173	182
m830R 308525-308809 95 10,948 11.00 No Hit Found No Hit Found m831R 308602-308862 87 10,140 9,67 No Hit Found No Hit Found																	
m830R 308525-308809 95 10,948 11.00 No Hit Found No Hit Found m831R 308602-308862 87 10,140 9,67 No Hit Found No Hit Found	m827L 307655-3074	16 70	7,766	6.76	No Hit Found					No Hit Found							
m832R 309017–309241 75 8,779 8.93 No Hit Found No Hit Found	m831R 308602-3088	32 87	10,140	9.67	No Hit Found					No Hit Found							
	m832R 309017-3092	11 75	8,779	8.93	No Hit Found					No Hit Found							
m834L 309526-309287 80 9,021 9.23 No Hit Found No Hit Found	m834L 309526-3092	37 80	9,021	9.23	No Hit Found					No Hit Found							
m835R 309618-309971 118 13,226 5.06 No Hit Found No Hit Found	m835R 309618-3099	71 118	13,226	5.06	No Hit Found					No Hit Found							
m836R 308842-310261 140 16,55 12,54 No Hit Found No Hit Found	m836R 309842-3102	31 140	16,555	12.54	No Hit Found					No Hit Found							
m837L 310068-309865 68 7,845 12.56 No Hit Found No Hit Found	m837L 310068-3098	35 68	7,845	12.56	No Hit Found					No Hit Found							
m839R 310328-310765 146 16.481 5.10 No Hit Found No Hit Found	m839R 310328-3107	35 146	16,481	5.10	No Hit Found					No Hit Found							
m840L 311207-319866 74 8,032 7,02 No Hit Found No Hit Found	m840L 311207-3109	36 74	8,032	7.02	No Hit Found					No Hit Found							
m841R 311031-311405 125 13,988 9.87 No Hit Found No Hit Found			40.000	0.07	No US Found												
	m841R 311031-3114	05 125	13,988	9.07	NO HIL FOULIG					No Hit Found							