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## Supplementary Data for "Sequence and annotation of the 369-kb NY-2A and the 345-kb AR158 viruses that infect *Chlorella* NC64A": Appendix C: Gene Names c001R - c814L

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### SUPPLEMENTARY DATA FOR

# Sequence and annotation of the 369-kb NY-2A and the 345-kb AR158 viruses that infect *Chlorella* NC64A

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

**Abstract:** Viruses NY-2A and AR158, members of the family *Phycodnaviridae*, genus *Chlorovirus*, infect the fresh water, unicellular, eukaryotic, chlorella-like green alga, *Chlorella* NC64A. The 368,683-bp genome of NY-2A and the 344,690-bp genome of AR158 are the two largest chlorella virus genomes sequenced to date; NY-2A contains 404 putative protein-encoding and 7 tRNA-encoding genes and AR158 contains 360 putative protein-encoding and 6 tRNA-encoding genes. The protein-encoding genes are almost evenly distributed on both strands, and intergenic space is minimal. Two of the NY-2A genes encode inteins, the large subunit of ribonucleotide reductase and a superfamily II helicase. These are the first inteins to be detected in the chlorella viruses. Approximately 40% of the viral gene products resemble entries in the public databases, including some that are unexpected for a virus. These include GDP-d-mannose dehydratase, fucose synthase, aspartate transcarbamylase, Ca<sup>++</sup> transporting ATPase and ubiquitin. Comparison of NY-2A and AR158 protein-encoding genes with the prototype chlorella virus PBCV-1 indicates that 85% of the genes are present in all three viruses.

Keywords: Chlorella viruses, Phycodnaviridae, Virus NY-2A, Virus AR158, Genome sequence

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, Peptide 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 b002R – b797R Appendix B: Gene Names B001L – B886R Appendix C: Gene Names c001R – c814L Appendix D: Gene Names C006R – C815L

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Gene	Genome	A.A.	Peptide	pl	CDD Hit	COGs COG Definition	Bit Score E-value	% % Identity Positive	Query	Hit from-	BLASTp Hit	Hit	BLASTp Definition E	it Score E-value	% entity Pos	% C	luery Hit
Name c001R	Position 8581208	lenath 117	Mw 13,216	10.00	Number	Hit Found		Identity Positive	from-to	to	Number	Accession No Hit Found	·	Id	entity Pos	sitive tr	om-to trom-to
c002L	1117-914	68	7,355			Hit Found						No Hit Found					
c003L	14071162	82	9,208			Hit Found						No Hit Found					
c004R	13451566	74	8,633	7.51	No H	Hit Found						No Hit Found					
c005L	27682553	72	7,560	6.39	No F	Hit Found						No Hit Found					
c007L	28532656	66	8,324	8.67	No I	Hit Found						No Hit Found					
c009R	41074307	67	8,180	7.70	No H	Hit Found						No Hit Found					
c010R	41954416	74	8,305	10.98	No I	Hit Found						No Hit Found					
c012R	47395137	133	14,851			Hit Found						No Hit Found					
c013L	52494968	94	10,864			Hit Found						No Hit Found					
c014L c016R	60595835	75	9,020			Hit Found						No Hit Found					
c016R c017R	59236135 66536874	71 74	8,679 8,908			Hit Found Hit Found						No Hit Found					
c019L	85068288	73	8,018			Hit Found						No Hit Found					
c020L	86368364	91	10,668			Hit Found						No Hit Found					
c022R	96299838	70	7,637			Hit Found						No Hit Found					
c023R	1001310267	85	9,780			Hit Found						No Hit Found					
c025L	10493-10260	78	8,776	12.21	No I	Hit Found						No Hit Found					
c026R	10400-10885	162	15,129	6.51	No H	Hit Found						No Hit Found					
c027R	10482-10724	81	7,004	7.01	No H	Hit Found						No Hit Found					
c028L	11138-10896	81	9,300	9.03	No H	Hit Found						No Hit Found					
c030L	12030-11833	66	7,378	12.02	No I	Hit Found						No Hit Found					
c032L	12699-12505	65	7,548	11.07	No H	Hit Found						No Hit Found					
c033R	14085-14363	93	10,260	4.54	No H	Hit Found						No Hit Found					
c034R	14457-14663	69	7,533			Hit Found						No Hit Found					
c035R	1540515620	72	8,118			Hit Found						No Hit Found					
c038L	2011419836	93	10,815			Hit Found						No Hit Found					
c039R	20154-20471	106	13,496			Hit Found						No Hit Found					
c040R c041R	20225-20644 21263-21613	140 117	15,363 12,757			Hit Found Hit Found						No Hit Found No Hit Found					
c042R	21734-21976	81	8,902			Hit Found						No Hit Found					
c043R	21986-22390	135	14.717			Hit Found						No Hit Found					
c045R	23050-23385	112	12,025	3.86	No I	Hit Found						No Hit Found					
c046R	23722-23919	66	7,153	3.83	No I	Hit Found						No Hit Found					
c047R	24526-24927	134	15,178	5.19	No I	Hit Found						No Hit Found					
c049L	26322-26053	90	10,915	10.53	No H	Hit Found						No Hit Found					
c050L	27023-26817	69	8,732	10.36	No H	Hit Found						No Hit Found					
c053R	29160-29459	100	11,626	8.45	No H	Hit Found						No Hit Found					
c054R	30036-30260	75	8,463	6.50	No I	Hit Found						No Hit Found					
c058L	32400-32086	105	11,811	11.09	No H	Hit Found						No Hit Found					
c059R	32134-32358	75	9,266	8.65	No I	Hit Found					1	NP_048391	a43R	160.23 1.61E-38	92%	94%	1-75 1-75
c060L	32599-32345	85	9,926			Hit Found						NP_048520	a172L	90.12 2.06E-17	49%	71%	485 486
c067R	36452-36772	107	12,931			Hit Found						NP_048400	a52R	72.79 3.37E-12	88%	91%	3974 136
c070L	37569-37273	99	11,285			Hit Found						No Hit Found					
c071R c076R	37404-37646 40834-41088	81 85	9,390 9,619			Hit Found						No Hit Found					
c077R	41317-41559	81	8,642			Hit Found						NP 048874	a518R	82.42 4.19E-15	61%	66%	1-74 175
c079R	41856-42158	101	11,879			Hit Found						No Hit Found	40.00	02.42 4.102.10	0170	00%	
c080R	42043-42243	67		11.66		Hit Found						No Hit Found					
c082L	42700-42482	73	8,199			Hit Found						No Hit Found					
c083L	42949-42734	72	7,961			Hit Found						No Hit Found					
c084L	43118-42810	103	12,209			Hit Found						No Hit Found					
c085R	42826-43095	90	10,283	12.37	No H	Hit Found						No Hit Found					
c087L	43830-43534	99	11,655	10.86	No I	Hit Found						No Hit Found					
c088L	44198-43725	158	18,278	6.91	No H	Hit Found					1	NP_048417	contains Gly-rich Gx motif LGGGLG (5X); contains type I hydrophobic trai	106.30 2.89E-22	60%	78%	77158 182
c091R	4541445683	90	10,722	12.10	No H	Hit Found						No Hit Found					
c092L	4597345743	77	8,822	4.37	No I	Hit Found						NP_048503	a155R	88.20 7.74E-17	67%	81%	1175 2286
c094L	4649846280	73	9,099	11.44	No H	Hit Found						No Hit Found					

Name	Position	lenath	Peptide Mw	pl	CDD Hit COGs Number	COG Definition	Bit Score	E-value	% % Identity Positive	Query from-to	to	BLASTp Hit Number	Hit Accession	BLASTp Definition	Bit Score	E-value lo	% entitv P	% ositive	Query from-to
c095R	4638446647	88	10,427	6.51	No Hit Found								No Hit Found						
c096R	4655946783	75	8,714	11.72	No Hit Found								No Hit Found						
c099R	4734847569	74	8,973	11.96	No Hit Found								No Hit Found						
c101L	47831-47637	65	7,735	11.80	No Hit Found							1	NP_048424	a76L	81.26	9.47E-15	59%	72%	7-6
c103R	4812748393	89	10,619	9.04	No Hit Found								No Hit Found						
c105L	48553-48332	74	8,787	6.23	No Hit Found								No Hit Found						
c106R	4847748920	148	16,988	12.25	No Hit Found								No Hit Found						
c107R	4866749050	128	15,872	12.73	No Hit Found								No Hit Found						
c108L	4903048701	110	12,147	4.53	No Hit Found								No Hit Found						
c109R	4895149169	73	8,248	4.44	No Hit Found								No Hit Found						
c111L	50241-50035	69	7,818	7.69	No Hit Found								No Hit Found						
c113L c113L	50991-50605	129	14,954	8.53	No Hit Found								NP_048428 NP_048808	a80L similar to PBCV-1 ORF a80L, encoded by GenBank Accession Number U	83.19	2.45E-15 4.64E-06	56% 55%	73% 77%	4
	51552-51854	404	11.969	5.03	No US Formal								=	Similar to PBCV-1 ORP about, encoded by Genbank Accession Number of	52.57	4.04E-00	3376	1176	1
	52181-52435	101	10.012		No Hit Found								No Hit Found						
	52181-52435	81	9.832	10.33	No Hit Found								No Hit Found						
					No Hit Found								No Hit Found	-001	445.40	5.005.05	000/	040/	
	53301-53107	65	6,979		No Hit Found								NP_048434	a86L	115.16	5.92E-25	88%	91%	1-4
	53809-53456	118	14,006		No Hit Found								No Hit Found						
	54576-54854	93	10,295		No Hit Found								No Hit Found						
	5600856250	81	9,887	12.34	No Hit Found								No Hit Found						
	5721857024	65	6,945		No Hit Found								No Hit Found						
	57627-57304	108	12,676		No Hit Found								No Hit Found						
	58671-58321	117	13,915	7.33	No Hit Found								No Hit Found						
	58873-58634	80	9,314	11.01	No Hit Found								NP_048449	a101L	80.49	1.60E-14	65%	87%	1
	6077860999	74	8,490	5.70	No Hit Found								No Hit Found						
c136L	6200561682	108	12,746	11.14	No Hit Found								No Hit Found						
c138L	62271-62071	67	7,631	11.05	No Hit Found							1	NP_048450	a102L	74.71	8.81E-13	67%	70%	1
c139R	6221862478	87	10,553	8.38	No Hit Found								No Hit Found						
c141R	63522-63800	93	10,443	9.04	No Hit Found								No Hit Found						
c144R	64162-64443	94	11,576	10.77	No Hit Found								No Hit Found						
c145R	6456364808	82	10,093	8.66	No Hit Found								No Hit Found						
c147R	6496465188	75	8,899	9.58	No Hit Found								No Hit Found						
c148L	6558465348	79	9,420	9.90	No Hit Found								No Hit Found						
	65751-65419	111	13,009		No Hit Found								NP_048458	a110L		4.30E-44	85%	89%	101
c151L	6640266112	97	10,606	9.77	No Hit Found							1	NP_048460	a112L	191.05	8.51E-48	96%	96%	1
c152L	66829-66602	76	8,774	9.66	No Hit Found							1	NP_048463	a115L	97.44	1.28E-19	61%	68%	1
c153R	66732-66965	78	9,066	7.94	No Hit Found							1	NP_048464	a116R	105.15	6.10E-22	70%	75%	1
c154L	6761867190	143	16,723	6.22	No Hit Found							1	NP_048465	a117L	177.56	9.63E-44	63%	80%	71
c156L	6890368421	161	19,134	8.25	No Hit Found								No Hit Found						
c158L	6954869270	93	10,616	7.09	No Hit Found								No Hit Found						
c159L	70317-69982	112	13,401	9.58	No Hit Found								No Hit Found						
c161L	7067170441	77	8,834	6.50	No Hit Found								No Hit Found						
c163L	72952-71135	606	66,489	4.07	No Hit Found							1	XP_499823	hypothetical protein	53.53	2.42E-05	23%	36%	2315
c164L	7344473241	68	7,426	4.32	No Hit Found								No Hit Found						
c165L	74207-73557	217	25,658	10.46	No Hit Found								No Hit Found						
c166L	7387673613	88	9,636	6.62	No Hit Found								No Hit Found						
c168L	7517974985	65	7,619	11.13	No Hit Found							1	NP_048473	a124L	59.31	3.85E-08	58%	66%	18
c170L	7605575735	107	12,611	10.91	No Hit Found								No Hit Found						
c172R	76271-76723	151	17,718	9.90	No Hit Found							1	NP_048480	a132R	59.31	3.81E-08	47%	67%	611
c173L c173L	77323-76730	198	23,198	4.95	No Hit Found								AAK23092 7P 00811430	hypothetical protein Methyltransferase FkbM		2.78E-10 1.80E-09	29% 30%	52% 51%	401 431
c173L														methyltransferase FkbM		4.01E-09	34%	52%	591
c176R	7808178329	83	9,177	10.51	No Hit Found								No Hit Found						
c178R	7912079335	72	8,465	7.77	No Hit Found								No Hit Found						
c181L	7987579660	72	7,514	5.54	No Hit Found								No Hit Found						
c183L	81681-80872	270	28,002	5.53	No Hit Found							1	NP_048490	a142L	172.17	1.47E-41	78%	78%	103-2
c183L	00004 00000	404	F0.0C-		Ma 199 From 4								NP_048491	a143L		1.54E-06	89%	89%	20
c184L	8226480882	461	53,389	8.44	No Hit Found							1	NP_048492 117632	a144L hypothetical protein a141L - Chlorella virus PBCV-1	526.17	8.95E-148 1.91E-73	57% 60%	67% 70%	54 2354
c184L c184L												-	NP_048495	Phe-, Gly-rich protein: RCGF 3X, GCGF 11X, RSGF 5X, GSGF 2X	210.20	1.82E-07	37%	47%	401

Gene Name c185L	Genome Position	A.A. lenath	Peptide Mw	pl	CDD Hit COGs Number	COG Definition Bi	it Score E	E-value Ide	% % entity Positive	Query from-to	Hit from- to		Hit ession 8492	BLASTp Definition a144L	Bit Score E-val	lder	% ntitv Po	% sitive 64%	Query from-to f	Hit rom-to 92155
c186L	83049-82717	111	10,598	9.60	No Hit Found							No Hit I	Found							
c188R	83010-83234	75	8,439	8.43	No Hit Found							No Hit F								
c190L	84315-83878	146	16,674	10.89	No Hit Found							1 NP_048		contains prokaryotic membrane lipoprotein lipid attachment site	75.87 3.90E	-13	53%	61%	67143	985
c192L c196R	84820-84611 86464-86658	70 65	8,351 7,842	10.54 9.24	No Hit Found No Hit Found							No Hit F								
c199R	87125-87364	80	8,733	4.43	No Hit Found							No Hit F								
c201L	8797687776	67	7,962	6.91	No Hit Found							1 NP_048		a167L	61.23 1.01E	-08	48%	62%	1-66	166
c205L	9099390712	94	11,356	6.86	No Hit Found							No Hit F	Found							
c207R	9166291946	95	11,440	11.54	No Hit Found							1 NP_048	8391	a43R	80.88 1.25E	-14	54%	65%	175	175
c209L	92782-92531	84	10,068	10.74	No Hit Found							No Hit F	Found							
c210R	92834-93049	72	7,509	5.54	No Hit Found							No Hit I								
c211L	93130-92852	93	11,611		No Hit Found							No Hit I								
c213R c214R	93448-93792 94069-94278	115 70	13,719 8,428	8.19 8.40	No Hit Found No Hit Found							No Hit F								
c215R	94322-94600	93	10.894	7.81	No Hit Found							No Hit I								
c218R	95123-95371	83	10,219		No Hit Found							No Hit I								
c219R	95277-95480	68	7,747	6.80	No Hit Found							1 NP_048		a251aL	73.94 1.50E	-12	67%	76%	1-56	83141
c221L	96277-96062	72	8,732	12.02	No Hit Found							No Hit I								
c222L	9669096298	131	13,927	10.62	No Hit Found							No Hit F	Found							
c223L	9656396369	65	7,794	11.02	No Hit Found							No Hit F	Found							
c224R	97366-97599	78	9,691	11.50	No Hit Found							No Hit F	Found							
c225L	9784697577	90	10,690	5.48	No Hit Found							No Hit F	Found							
c226L	9807397873	67	7,104	5.33	No Hit Found							No Hit I	Found							
c229L	9958599382	68	7,374	11.50	No Hit Found							No Hit I								
c231L	99829-99557	91	11,323 9.087		No Hit Found							No Hit I		.107	51.60 8.02E	- 00	59%	87%	4 00 0	40.047
	100234100004	77 68	7,244	8.20 12.23	No Hit Found No Hit Found							1 NP_048 1 NP_048		a187L a188L	105.15 6.06E		83%	86%	1-32 2 1-68	
	102087101800	96	11,556	8.52	No Hit Found							No Hit I		41002	100.10 0.002		0070	0070		0 /2
c237L	102706102467	80	9,028	4.91	No Hit Found							No Hit I	Found							
c238R	102616103026	137	16,220	11.53	No Hit Found							No Hit F	Found							
c240L	107496107272	75	8,593	10.37	No Hit Found							No Hit F	Found							
c243R	108582108794	71	8,346	8.35	No Hit Found							1 NP_048	8545	a198R	88.97 4.45E	-17	54%	75%	1-70	574
c247L	110412110158	85	10,356	9.47	No Hit Found							No Hit I								
	110609110232		13,951	10.58	No Hit Found							1 NP_048		A201L	135.96 3.23E	-31	66%	79%	24116	193
	110539110739	67	7,624	10.92	No Hit Found							No Hit I								
	110992110705 111325111116	96 70	11,788 7.097	10.16 9.46	No Hit Found  No Hit Found							No Hit I 1 NP_048		a204L	125.56 4.31E	- 20	92%	94%	170	1.70
	112026111583	148	16.166	5.80	No Hit Found							No Hit I		d2U%L	125.50 4.515	20	9276	3470	1-70	170
	112858112664	65	7,957	9.66	No Hit Found							No Hit I								
	113487113155	111	13,135	8.09	No Hit Found							No Hit I								
c262R	115076115321	82	9,540	10.65	No Hit Found							No Hit I	Found							
c266L	118141117878	88	10,558	9.46	No Hit Found							No Hit F	Found							
c267L	118131117883	83	9,410	11.05	No Hit Found							1 T17711	1	hypothetical protein a221L - Chlorella virus PBCV-1	70.09 2.14E	-11	75%	88%	1-44	144
	118076118285	70	7,704	9.74	No Hit Found							1 NP_048		a223R	102.06 5.10E		71%	78%	170	170
	118757118551	69	8,032	9.41	No Hit Found							1 NP_048		a224L	78.57 6.06E	-14	90%	92%	140	49-88
	120301120582		10,998	7.83	No Hit Found							No Hit I								
	120507120313		7,395	9.43	No Hit Found							No Hit I								
	120772120999 121048121383		8,512 12,845		No Hit Found No Hit Found							No Hit F								
	121413121619		8,266		No Hit Found							1 NP_048		a232R	72.79 3.32E	-12	56%	73%	665	665
	122093121860		9,730		No Hit Found							No Hit I								
	123270123076		7,071		No Hit Found							No Hit I								
c283L	123376123122	85	10,277	8.92	No Hit Found							No Hit I	Found							
c284R	123303123560	86	10,147	11.32	No Hit Found							No Hit F	Found							
c287L	124418124002	139	16,327	7.51	No Hit Found							No Hit I	Found							
	124187124480	98		11.21	No Hit Found							No Hit F								
c289L	124901124662	80	9,384	9.63	No Hit Found							No Hit F	Found							

Gene Genor Name Positi	ion I	A.A. I enath	Peptide Mw 11.173	<b>pl</b> 8.70	CDD Hit Number COGs	COG Definition	Bit Score E-valu	e % Identitv F	% Qu Positive fro	uery F om-to	lit from- BLASTp Hit Hit to Number Accessio	BLASTp Definition	Bit Score E-value Id	% entitv Po	% Positive	Query Hit from-to
c293L 1268611:		117	13,667	5.15	No Hit Found						1 NP_048590	a242L	174.48 8.20E-43	74%	79%	1-117 1-117
c294L 1271511	126864	96	10,473	10.81	No Hit Found						No Hit Found					
c295L 1275741:	127317	86	9,989	9.40	No Hit Found						No Hit Found					
c296L 1276111:	127354	86	9,966	9.45	No Hit Found						No Hit Found					
c297L 1279731	127674	100	11,308	10.05	No Hit Found						No Hit Found					
c301L 1313971	131020	126	15,612	10.02	No Hit Found						No Hit Found					
c302L 1312851	131067	73	8,030	11.07	No Hit Found						No Hit Found					
c304R 1320441	132301	86	10,130	10.16	No Hit Found						No Hit Found					
c305R 1329801	133429	150	17,984	10.04	No Hit Found						No Hit Found					
c308L 1347601		84	10,506	9.57	No Hit Found						1 NP_048603	a252aL	90.51 1.58E-17	66%	80%	2584 2584
c310R 1352381		209	23,434	7.79	No Hit Found						No Hit Found					
c312R 1359521		67	7,723	9.56	No Hit Found						1 NP_048618	a264R	100.91 1.15E-20	67%	85%	161 161
c314R 1363321		107	11,951	10.71	No Hit Found						No Hit Found					
c316R 1369261		84	9,454	10.79	No Hit Found						1 NP_048624	a270R	82.03 5.62E-15	63%	67%	1-68 1-68
c317R 1371081		201	23,972	6.45	No Hit Found						1 NP_048626	a272R	71.63 1.50E-11	52%	74%	2890 567
c320R 1391751		123	13,615	8.82	No Hit Found						1 NP_048635	a281R	174.48 8.29E-43	83%	85%	8113 70176
c322R 1395611		72	8,843	9.57	No Hit Found						1 NP_048633	a279R	158.69 4.56E-38	95%	95%	1-72 1-72
c323R 1406571		93	10,245	3.62	No Hit Found						1 NP_048634	a280R	174.10 1.05E-42	93%	93%	1-93 1-93
c325L 1416641- c326R 1426061-		67 67	7,712		No Hit Found						1 NP_048637	a283L	138.27 6.49E-32	94%	97%	167 167
c328L 1430601		88	8,415 10,735	10.20	No Hit Found No Hit Found						No Hit Found No Hit Found					
c330R 1442221		80	9.135	10.73	No Hit Found						No Hit Found					
c331L 1446561		89	11.112		No Hit Found						No Hit Found					
c332R 1444091		70	8.349	11.47	No Hit Found						No Hit Found					
c333R 1446181		79	9,404	11.24	No Hit Found						No Hit Found					
c335l 1459621		93	10,511	8.11	No Hit Found						No Hit Found					
c336l 1460981	145880	73	8,842	11.23	No Hit Found						No Hit Found					
c338R 1465271	146901	125	15,299	8.38	No Hit Found						No Hit Found					
c340L 1475271	147114	138	16,602	10.27	No Hit Found						1 BAA20344	ORF245	137.50 1.09E-31	60%	74%	1115 125243
c341L 1473421		73	8,987	10.31	No Hit Found						No Hit Found					
c343R 1480341	148369	112	12,380	10.29	No Hit Found						1 NP_048647	a293R	149.83 2.13E-35	73%	79%	1102 67168
c345R 1487271	148927	67	7,924	6.08	No Hit Found						No Hit Found					
c347L 1502201	149966	85	9,964	7.48	No Hit Found						No Hit Found					
c350R 1518471	152107	87	10,253	8.53	No Hit Found						No Hit Found					
c352L 1524821	152276	69	8,208	9.95	No Hit Found						No Hit Found					
c353L 1526641	152470	65	7,404	10.57	No Hit Found						No Hit Found					
c355R 1527251	152958	78	8,970	11.50	No Hit Found						No Hit Found					
c356L 1535121	152814	233	27,426	7.38	No Hit Found						1 NP_048503	a155R	92.43 1.13E-17	66%	86%	1683 2794
c357L 1533431	153101	81	9,010	9.52	No Hit Found						No Hit Found					
c359R 1538691	154096	76	9,221	10.65	No Hit Found						No Hit Found					
c360R 1541161	154376	87	9,662	8.83	No Hit Found						No Hit Found					
c361R 1546291	154838	70	7,340	10.81	No Hit Found						No Hit Found					
c362L 1551161	154841	92	10,661	11.19	No Hit Found						No Hit Found					
c364L 1557501	155502	83	9,737	9.91	No Hit Found						No Hit Found					
c366R 1570111: c366R	157211	67	7,781	8.41	No Hit Found						1 CAB92310 2 AAF74028	DNA methyltrasferase M.Hpy188I	64.70 9.11E-10 64.70 9.11E-10	51% 51%	67% 67%	1-66 348-415 1-66 348-415
c368L 1575521	157358	65	7,724	8.65	No Hit Found						No Hit Found					
c370L 1582881		132	16,358		No Hit Found						No Hit Found					
c371R 1580931		76	8,992		No Hit Found						No Hit Found					
c373R 1585411:	158852	104	12,534	7.89	No Hit Found						No Hit Found					
c375L 1602251	159938	96	10,814		No Hit Found						No Hit Found					
c376L 1604941	160288	69	7,990		No Hit Found						No Hit Found					
c383R 1636041	163978	125	14,390	8.56	No Hit Found						No Hit Found					
c385R 1640621	164265	68	8,318		No Hit Found						No Hit Found					
		147	13,781	4.87	1 pfam05277	DUF726, Protein of unknown function (DUF726). This family consists of s	38.34 1.41E-	03 45%	57%	32120						
c388L 1652041																
c388L 1652041	165239	122	14,249	11.34	No Hit Found						No Hit Found					
		122 82	14,249 8,693	11.34 9.57	No Hit Found No Hit Found						No Hit Found 1 NP_048675	a319L	148.29 6.19E-35	87%	90%	1-82 1-82

Gene Name	Genome Position	A.A. lenath	Peptide Mw	pl	CDD Hit COGs	COG Definition E	it Score E	E-value	% % entity Positive	Query from-to	Hit from- BLAS	STp Hit Hit mber Accessi	DD BLASTp Definition	Bit Score E-value lo	%	%	Query	Hit rom-to
	166963167274	104	12,249	12.09	No Hit Found			140				No Hit Foun						0111 10
c395L	167296167054	81	9,032	9.53	No Hit Found							No Hit Foun						
c399R	168424168642	73	7,772	10.81	No Hit Found							No Hit Foun	i					
o401R	169138169344	69	7,754	9.38	No Hit Found							No Hit Foun						
	169430169825		15,519	10.69	No Hit Found							No Hit Foun						
c403L	170154169894	87	10,415	9.01	No Hit Found							No Hit Foun						
c405R	170653170910	86	10,333	10.62	No Hit Found							1 NP_048683		79.72 2.77E-14	64%	75%	2586	24-85
c407L	171282171010	91	11,145	10.73	No Hit Found							No Hit Foun	i					
c408L	171736171488	83	9,597	8.64	No Hit Found							No Hit Foun	i					
c409R	172546172800	85	9,366	10.41	No Hit Found							No Hit Foun	1					
c410R	172862173068	69	8,172	6.46	No Hit Found							No Hit Foun						
c411L	173150172947	68	7,941	8.81	No Hit Found							No Hit Foun						
c412L	173465173232	78	9,004	4.36	No Hit Found							No Hit Foun						
c416R	175246175455	70	7,673	10.25	No Hit Found							No Hit Foun	1					
	175512175745	78	9.550	11.08	No Hit Found							No Hit Foun						
	176799176599	67	7,351	9.43	No Hit Found							No Hit Foun						
	177097176882		8,469	4.76	No Hit Found							No Hit Foun						
	177102176896	69	7,865	9.63	No Hit Found							No Hit Foun						
	177516177734	73	8,106	8.19	No Hit Found							No Hit Foun						
	178650178339		12,773	12.59	No Hit Found							No Hit Foun						
	178438178758			7.07	No Hit Found							No Hit Foun						
	179432179037		12,519															
		132	16,002	11.55	No Hit Found							No Hit Foun						
	179380179592	71	8,587	10.24	No Hit Found							No Hit Foun						
	180919181362	148	16,592	8.66	No Hit Found							1 NP_048701		143.28 2.03E-33	60%	77%	34148	1115
	181894182229	112	13,285	11.17	No Hit Found							No Hit Foun						
	182240182512	91	10,959	11.10	No Hit Found							No Hit Foun						
	183172182717	152	18,139	6.83	No Hit Found							1 NP_048704		92.05 5.29E-18	57%	72%	83152	170
	185762186091	110	12,244	10.57	No Hit Found							No Hit Foun						
	185875186207	111	13,069	11.06	No Hit Found							No Hit Foun						
	186389186598	70	7,646	4.83	No Hit Found							No Hit Foun						
c445L	186897186688	70	7,690	12.35	No Hit Found							No Hit Foun	i					
c446R	186739186942	68	8,981	12.71	No Hit Found							No Hit Foun	1					
	187744188016	91	9,560	4.77	No Hit Found							No Hit Foun	1					
c450L	188399188181	73	8,160	8.21	No Hit Found							No Hit Foun	1					
c451L	188608188411	66	8,354	8.68	No Hit Found							No Hit Foun	1					
c452L	188952188668	95	10,699	7.95	No Hit Found							No Hit Foun	i					
c453L	189466189254	71	8,352	10.01	No Hit Found							No Hit Foun	1					
c454L	189905189540	122	13,554	8.20	No Hit Found							No Hit Foun	i					
c455L	190553190311	81	8,974	4.64	No Hit Found							No Hit Foun	1					
c456L	190890190396	165	19,722	9.86	No Hit Found							No Hit Foun	1					
c457L	191383191150	78	9,128	7.98	No Hit Found							No Hit Foun	1					
c458L	191654191370	95	10,029	3.14	No Hit Found							No Hit Foun	i					
c460R	191923192192	90	10,868	10.95	No Hit Found							No Hit Foun	i					
c461R	192227192478	84	9,664	8.22	No Hit Found							No Hit Foun	i					
c463R	192330192695	122	15,030	10.78	No Hit Found							1 NP_048724	a367R	69.32 3.64E-11	41%	49%	1-91	191
c464R	192682192903	74	8,075	3.54	No Hit Found							No Hit Foun	1					
c465R	193494193793	100	11,295	12.65	No Hit Found							1 NP_048734	a377R	86.27 2.92E-16	55%	63%	14100	691
c468R	194418194948	177	20,563	11.10	No Hit Found							1 NP_048737	a380R	144.05 1.77E-33	71%	82%	49149	1101
c469R	194668194871	68	7,734	11.11	No Hit Found							1 NP_048738	a381R	127.49 1.14E-28	89%	91%	1-68	976
c471L	195563195276	96	10,916	8.81	No Hit Found							No Hit Foun	1					
c472L	195962195699	88	9,947	12.15	No Hit Found							No Hit Foun	i					
c473L	196443196084	120	14,174	8.04	No Hit Found							No Hit Foun	1					
c474L	197139196792	116	12,847	10.36	No Hit Found							1 NP_048742	a385L	59.31 3.86E-08	85%	85%	3569	135
c476R	196916197356	147	17,664	12.63	No Hit Found							1 NP_048743	Gly-rich, AGLG (9x); similar to herpesvirus hypothetical protein 5, corresp	61.62 7.82E-09	80%	80%	6196	57-92
c477R	197986198198	71	8,501	11.94	No Hit Found							1 NP_048745	a388R	50.83 1.34E-05	78%	78%	2-33	
	198384198680		11,302		No Hit Found							1 NP_048748		146.36 2.39E-34	76%	77%		195
	199076198861	72		6.73	No Hit Found							No Hit Foun						
	199524199727	68		6.49	No Hit Found							No Hit Foun						

Gene Name	Genome Position	A.A. lenath	Peptide Mw	pl	CDD Hit 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	%	% Positive	Query from-to	Hit from to
	199906199709	66	8,079	6.08	No Hit Found				identity rositive	110111-10	10		No Hit Found			identity	rositive	110111-10	ii oiii-to
c488R	202540202737	66	7,352	5.69	No Hit Found								No Hit Found						
c489R	202826203029	68	8,180	8.38	No Hit Found								No Hit Found						
c495L	205972205670	101	10,913	4.64	No Hit Found								No Hit Found						
c499L	207957207397	187	19,795	13.59	No Hit Found								No Hit Found						
c500L	208594208388	69	7,216	4.54	No Hit Found								No Hit Found						
	208895209098	68	9,523		No Hit Found								No Hit Found						
	209135208896	80	7,695	10.89	No Hit Found								No Hit Found						
	210028210228		7,716		No Hit Found								NP_048712	a355L	69.71 2.83E-1	1 59%	62%	2-65	1470
	210244210450	69 123	8,191 13,737	7.34	No Hit Found No Hit Found								No Hit Found NP_048766	a409R	73.94 1.52E-1	2 82%	86%	1 46	37-82
	212597212268	110	12,766		No Hit Found								No Hit Found	840811	73.84 1.322-1.	0270	0070	1-40	37-02
c514R	213721213972	84	9,652	8.82	No Hit Found								No Hit Found						
c516R	214835215071	79	8,394	10.76	No Hit Found								No Hit Found						
c517R	215109215402	98	9,869	6.50	No Hit Found								No Hit Found						
c520R	216571216768	66	8,038	7.12	No Hit Found								No Hit Found						
c524L	218761218330	144	17,685	6.94	No Hit Found								No Hit Found						
c525L	218547218350	66	7,822	7.78	No Hit Found								No Hit Found						
c526R	218409218711	101	11,774	10.92	No Hit Found								No Hit Found						
	219598219834	79		12.42	No Hit Found								No Hit Found						
	222712222963		10,680	11.63	No Hit Found								No Hit Found						
	222970222776			4.64	No Hit Found								No Hit Found						
	223213223422 224827225084	70		11.50	No Hit Found No Hit Found								No Hit Found						
	225889226092	86 68	10,233 8.318	10.15	No Hit Found No Hit Found								NP_048799	a442R	74.33 1.15E-1	2 47%	70%	1.66	266
	228137227904	78		8.19	No Hit Found								No Hit Found	044217	74.33 1.132-1.	4770	7070	1-00	2-00
	228274228053	74	9,185	12.38	No Hit Found								No Hit Found						
	228515228820	102	11,578	10.84	No Hit Found								No Hit Found						
c552R	228980229195	72	7,853	9.49	No Hit Found							1	NP_048803	a446R	102.45 3.88E-2	1 89%	91%	1772	156
c553R	229460229654	65	7,222	11.63	No Hit Found							1	NP_048804	a447R	126.72 1.97E-2	3 95%	98%	165	3599
c555L	230640230437	68	8,309	9.20	No Hit Found								No Hit Found						
c557L	231355231155	67	7,965	8.72	No Hit Found								No Hit Found						
c561R	233698233922	75	8,478	5.50	No Hit Found							1	NP_048810	a453R	88.58 5.96E-1	7 64%	66%	175	1589
	234730235155				No Hit Found								NP_048814	a457R	115.16 5.90E-2		80%	32107	176
	235249235001	83			No Hit Found								NP_048815	a458L	107.07 1.58E-2	2 60%	74%	1-83	183
	235008235334 235497235772	109	13,234	6.23	No Hit Found								No Hit Found						
	235840236046	92 69	10,922 7,712		No Hit Found  No Hit Found								No Hit Found						
	239382239621	80	9,499	5.58	No Hit Found								No Hit Found						
	241380241135	82	8,847	9.44	No Hit Found								NP_048825	a469L	54.30 1.22E-0	3 42%	52%	8-82	376
c576R	242004242207	68	8,156	12.37	No Hit Found								No Hit Found						
c578L	243006242806	67	7,615	7.28	No Hit Found								No Hit Found						
c581R	244422244925	168	20,013	9.69	No Hit Found							1	NP_048835	Phe-rich	63.16 3.42E-0	34%	36%	1107	1-107
c582L	244885244649	79	9,482	12.03	No Hit Found								No Hit Found						
c585R	245549245776	76	8,634	8.17	No Hit Found								No Hit Found						
	247798247472		12,914	6.61	No Hit Found								No Hit Found						
	248180247986	65		8.15	No Hit Found								No Hit Found						
	248677248378		11,404		No Hit Found								No Hit Found						
	248861248628			8.82	No Hit Found								No Hit Found						
	250572250826 251481251077	85 135		9.04	No Hit Found								No Hit Found NP_048854	Pro-rich, IPPPNMSLPLS (3x)	108 61 5 50F-2	3 52%	52%	1 120	1120
	251366251136	135		8.11	No Hit Found								NP_048854 NP_048855	a499L	56.23 3.26E-0		92%		1120
	251958252380	141	12,711		No Hit Found								No Hit Found		0.EUL-0	0070	UL 70	. 2/	. =-
c606R	253462253725				No Hit Found								NP_048860	a504R	176.79 1.65E-4	3 90%	95%	1-87	187
c607L	253954253745			10.53	No Hit Found								No Hit Found						
c609R	254024254254	77	9,572	8.82	No Hit Found								No Hit Found						
c610R	254295254501	69	8,436	11.12	No Hit Found							1	NP_048863	a507R	87.43 1.30E-1	60%	78%	165	367
c611R	254634254846	71	8,705	11.52	No Hit Found							1	NP_048863	a507R	77.80 1.03E-1	3 57%	71%	1-71	116186

Gene Name	Genome Position	A.A. lenath	Peptide Mw	pl	CDD Hit COGs	COG Definition	Bit Score E-v	value %	% Positive	Query from-to	Hit from- to	BLASTp Hit Number	Hit Accession	BLASTp Definition	Bit Score E-va	lue   %	tity Po	% sitive f	Query rom-to f	Hit rom-to
	255250255477	76	8,867	10.98	No Hit Found			identity	1 Oslave	110111-10	10		NP_048866	a510R	93.59 1.85		89%	92%		256
c614L	256042255743	100	11,629	11.36	No Hit Found								No Hit Found							
c617R	256457256780	108	13,055	3.84	No Hit Found							1	No Hit Found							
c623R	260350260727	126	14,999	10.45	No Hit Found							1	No Hit Found							
c625R	260645260917	91	10,217	4.48	No Hit Found							1 1	NP_048503	a155R	83.57 1.88	E-15	61%	79%	1682	27-93
c626R	261053261349	99	12,307	11.85	No Hit Found							١	No Hit Found							
	261949262200	84	9,930	6.00	No Hit Found								No Hit Found							
c631L c631L c631L	262842262534	103	11,102	6.48	No Hit Found							2 1	NP_048874 NP_048885 NP_049041	a518R a529L a685R	123.64 1.63 112.85 2.88 52.37 4.62	E-24	64% 86% 53%	72% 88% 59%	3197	3100 167 1566
c633L	263400263116	95	10,738	7.34	No Hit Found							1	No Hit Found							
c637R	265541265768	76	9,000	8.21	No Hit Found							1	No Hit Found							
	267680267480	67	7,432		No Hit Found								No Hit Found							
	267684267893	70	7,788	8.08	No Hit Found								No Hit Found							
	268730268954 269556269795	75 80	9,167 8,610	10.45	No Hit Found								No Hit Found							
	270148270354	69	8,068	10.67	No Hit Found No Hit Found								No Hit Found							
	270408270824	139	15,586	10.08	No Hit Found								No Hit Found							
	270560270859	100	12,308	8.19	No Hit Found								No Hit Found							
	272131272412	94	10,313		No Hit Found								No Hit Found							
c654R	272479272742	88	9,942	7.69	No Hit Found							1	No Hit Found							
c655R	273079273276	66	7,138	4.14	No Hit Found							1 1	NP_048897	a541R	59.69 2.94	E-08	51%	62%	366	34-96
c656R	274303274548	82	8,845	4.71	No Hit Found							1	No Hit Found							
c657L	275164274937	76	9,244	11.55	No Hit Found							1	No Hit Found							
c659L	276357275908	150	17,931	8.50	No Hit Found							1	No Hit Found							
c660L	276176275958	73	8,154	10.82	No Hit Found							1 1	NP_048901	a545L	132.11 4.74	E-30	87%	91%	173	173
	277646277912	89	10,287	5.06	No Hit Found								No Hit Found							
	278090278368	93	10,919	6.62	No Hit Found								NP_048905	a549R	74.33 1.14		41%	62%		180
	278744278977	78	9,229	6.35	No Hit Found								NP_048906	a550R	77.80 1.04	E-13	44%	74%	178	55132
	279068279433 280152279628	122 175	14,344 21,526	6.84 9.79	No Hit Found No Hit Found								No Hit Found							
	280659281009	117	13,112		No Hit Found								No Hit Found							
	280985281263	93	10.824	8.37	No Hit Found								NP 048911	a555R	168.32 5.78	E-41	89%	91%	1-93	193
c673R	281654281881	76	8,378	8.06	No Hit Found								No Hit Found							
c674L	282272282066	69	7,971	10.96	No Hit Found							1	No Hit Found							
c677R	283694284077	128	14,105	6.98	No Hit Found							1 1	NP_048916	a560R	72.79 3.33	E-12	65%	75%	362	45104
c679R	284289284639	117	13,356	4.85	No Hit Found							1	No Hit Found							
c680R	284566284850	95	10,558	7.72	No Hit Found							1	No Hit Found							
c682R	285472285705	78	9,588	11.24	No Hit Found							1	No Hit Found							
c683R	285888286115	76	8,753	7.57	No Hit Found							1 1	NP_048922	a566L	95.13 6.35	E-19	95%	95%	3576	4990
	286278286030	83	9,586	12.35	No Hit Found								No Hit Found							
	288667288263	135	15,524	7.64	No Hit Found								NP_048922	a566L	168.70 4.48	E-41	88%	88%	1-88	190
	290040290249	70	8,717	8.45	No Hit Found								No Hit Found							
	291082290816 291831292052	89 74	10,577 8,008	8.53 6.65	No Hit Found No Hit Found								No Hit Found							
	292684292448	79	8.679	10.91	No Hit Found								NP 048874	a518R	64.31 1.19	= 00	53%	66%	1778	1.62
	294008294205	66	7.959	8.96	No Hit Found								No Hit Found	85101	04.51 1.10	L-09	3376	0070	1770	1-02
c702R	294495294737	81	10,045	12.26	No Hit Found							1	No Hit Found							
c703L	295008294775	78	9,649	11.87	No Hit Found							1 1	NP_048938	a582L	122.87 2.82	E-27	78%	85%	1-78	178
c704R	295166295399	78	9,098	11.65	No Hit Found							1	No Hit Found							
c706R	295636295863	76	8,209	9.61	No Hit Found							1	No Hit Found							
c708R	296776297015	80	9,519	4.84	No Hit Found							1	No Hit Found							
c709R	297878298177	100	11,570	11.19	No Hit Found							1	No Hit Found							
	298147298341	65	7,526	7.53	No Hit Found							1 1	NP_048943	a587R	112.08 5.01	E-24	80%	89%	165	165
	298310298549	80	8,692		No Hit Found								No Hit Found							
	298822299517	232	26,620		No Hit Found								NP_048944	a588R	109.38 8.85	E-23	50%	68%	1116	1-116
	299039299362	108	12,116		No Hit Found								No Hit Found			- 40	200/	000:		40.0-
	301023300682 300930301148	114 73	11,519		No Hit Found No Hit Found								NP_048951 No Hit Found	a595L	65.86 4.16	E-10	62%	68%	1-59	19-82
6/1/R	JUU93U3U1148	/3	8,461	11.74	NO HIT FOUND							,	NO TIL FOUND							

Gene Name	Genome Position	A.A. lenath	Peptide Mw	pl	CDD Hit CO	OGs COG Definition	Bit Score	E-value	% Identity	% Positive	Query from-to	Hit from- I	BLASTp Hit Number	Hit Accession	BLASTp Definition	Bit Score I	E-value Id	% entity Po		Query from-to f	Hit rom-to
c720R	301999302544	182	21,575	6.23	No Hit	Found								No Hit Found							
c721R	302351302581	77	8,832	10.50	No Hit	Found							1	NP_048956	a600R	95.52	4.85E-19	67%	75%	3-76	11-81
c724L	303571303308	88	10,598	5.09	No Hit	Found								No Hit Found							
c730R	306186306458	91	10,984	9.52	No Hit	Found							1	NP_048966	a610R	173.71	1.38E-42	89%	89%	1-91	191
c732R	307630307887	86	9,951	6.95	No Hit	Found								No Hit Found							
c734R	308977309249	91	9,847	9.69	No Hit	Found							1	NP_048972	a616R	116.32	2.62E-25	62%	67%	2-91	493
c738R	310936311184	83	9,070	5.59	No Hit	Found								No Hit Found							
c743R	313791314033	81	10,084	11.26	No Hit	Found								No Hit Found							
c746L	315067314846	74	8,396	10.19	No Hit	Found							1	NP_048982	a626L	164.85	6.57E-40	100%	100%	1-74	174
c749R	317565317786	74	8,495	11.36	No Hit	Found							1	NP_048986	a630R	95.13	6.39E-19	67%	67%	1-74	174
c750R	317883318134	84	9,368	10.23	No Hit	Found								No Hit Found							
c751L	318217317990	76	8,645	9.54	No Hit	Found								No Hit Found							
c752R	319348319650	101	11,949	9.40	No Hit	Found								No Hit Found							
c758L	321419321105	105	12,528	4.79	No Hit	Found								No Hit Found							
c760L	321983321588	132	14,744	9.00	No Hit	Found							1	NP_048995	a639L	250.37	1.19E-65	95%	96%	1132	1132
c761R	321812322048	79	8,848	11.89	No Hit	Found							1	NP_048996	a640R	152.14	4.33E-36	94%	97%	179	179
c762L	322373322119	85	9,710	11.11	No Hit	Found							1	NP_048997	a641L	168.32	5.94E-41	96%	97%	1-85	185
c764L	323418323110	103	11,914	11.57	No Hit	Found								No Hit Found							
c768R	325179325496	106	12,145	7.92	No Hit	Found								No Hit Found							
c769L	325587325351	79	9,349	11.86	No Hit	Found							1	NP_049011	a655L	73.17	2.56E-12	57%	70%	978	170
c770R	325499325759	87	11,038	11.44	No Hit	Found								No Hit Found							
c775R	327489327767	93	43,880	8.00	No Hit	Found								No Hit Found							
c776R	327685327906	74	8,238	4.18	No Hit	Found							1	NP_048503	a155R	84.34	1.13E-15	66%	76%	573	1884
	328022327756	89	9,671	10.81	No Hit									No Hit Found							
c778R	327854328186	111	12,186	8.25	No Hit	Found								No Hit Found							
c779L	328307328026	94	11,341	12.45	No Hit	Found							1	NP_048504	a156L	56.61	2.44E-07	39%	59%	6-93	19107
	328556328332	75	8,562	12.23	No Hit									No Hit Found							
	329124328930	65		7.92	No Hit									No Hit Found							
	329322329543	74		8.07	No Hit									No Hit Found							
	330130330351	74	8,035	10.78	No Hit									No Hit Found							
	330745331128			7.85	No Hit									No Hit Found							
	332984333271	96		7.85	No Hit									NP_049023	similar to Chlorella virus CVK2 unknown ORF, corresponds to GenBank A		3.24E-47	91%	97%	196	196
	333374333568	65		7.72	No Hit									NP_049024	similar to Chlorella virus CVK2 unknown ORF, corresponds to GenBank A		5.72E-28	87%	93%	165	165
	333483333698	72	8,040	11.46	No Hit									NP_049025	similar to Chlorella virus CVK2 unknown ORF, corresponds to GenBank A		2.12E-35	100%	100%		12-83
	334913335278	122			No Hit								1	NP_049026	similar to Chlorella virus CVK2 unknown ORF, corresponds to GenBank A	223.79	1.16E-57	87%	92%	1122	1122
	336130335867	88		4.66	No Hit									No Hit Found							
	336125336397	91		11.12	No Hit									No Hit Found		50.04	0.005.00	400/	000/	40 440	
	336198336527 337668337270			10.96	No Hit									NP_048630 NP_049033	similar to PBCV-1 ORF a80L, corresponds to GenBank Accession Numbe a677L		3.82E-08 5.93E-25	48% 76%	66% 82%	49110	
	337668337270	133	14,418	7.92									'	NP_049033 No Hit Found	867/L	115.16	5.93E-25	76%	82%	1-69	175
	339024339272	206	9,941 23.724	7.92	No Hit									NP 048354	a6L	67.40	3.02E-10	47%	67%	74143	4 74
	339521339748			9.74	No Hit									No Hit Found	acc	67.40	3.02E-10	4/70	0776	/4143	1/1
	340292340516	75		4.17	No Hit									No Hit Found							
	340292340516	75	8,658	10.19	No Hit									No Hit Found							
	340919341197	93	10.890	10.19	No Hit									No Hit Found							
	341935342282	116		5.54	No Hit								1	NP_048354	a6L	57.38	1.47E-07	51%	72%	4086	147
	342680342471	70	8.233	5.13	No Hit									No Hit Found		07.00	01	/0			
	342546342746	67	7.515	7.18	No Hit								1	NP 049046	identical to PBCV-1 terminal repeat ORF, corresponds to GenBank Acces	58 54	6.53E-08	50%	68%	160	12-66
	343070342846	75	.,	6.66	No Hit									No Hit Found	2 2 7 Termina report or a consequent to Gendank Acces	55.54		00,0	00,0		.2 00
	343967343539	143	16,443		No Hit									No Hit Found							
			,																		