

1 **Supplemental Information**

2 *Paramecium bursaria* Chlorella Virus 1 Proteome Reveals Novel Architectural and
3 Regulatory Features of a Giant Virus

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9 ***Genome re-sequencing with Roche 454 Life Sciences GS FLX Titanium***

10 Preliminary proteomic analyzes using the existing PBCV-1 gene annotations
11 (NCBI Refseq: NC_000852) revealed possible errors in the genome sequence, which
12 prompted re-sequencing of the PBCV-1 genome. PBCV-1 DNA was purified from
13 virions that had been treated with 10 units DNase I per ml in 50 mM Tris-HCl pH 7.8/1
14 mM CaCl₂/10 mM MnCl₂ at 37°C for 1 hr, then prepared as described above, using the
15 UltraClean®Blood DNA Isolation Kit (MO BIO Laboratories, Carlsbad, CA). The DNA
16 was evaluated for quantity and quality by measuring absorbance at 260 and 280 nm with
17 a Thermo Scientific NanoDrop 2000 spectrophotometer, and by measuring fluorescence
18 of dye-augmented DNA using the PicoGreen and a Qubit fluorometer (Invitrogen,
19 Carlsbad, California). Using the Roche Rapid Library Preparation method for GS FLX
20 Titanium chemistry (Roche 454 Life Sciences, Branford, Connecticut), sample DNA was
21 fragmented by nebulization. DNA fragments were end repaired with polynucleotide
22 kinase and T4 DNA polymerase, then purified by size exclusion chromatography.

23 Selected DNA fragments were ligated to a Rapid Library Multiplex Identifier (MID)
24 adaptor designed for GS FLX Titanium chemistry. The MID adaptors were designed
25 with a unique decamer sequence to facilitate multiplex sequencing with the 454
26 technology, such that the resulting library reads can be reliably sorted after sequencing
27 using SFF software tools. MID adaptor ligated DNA fragments were again size selected
28 by chromatography, quantified with a TBS-380 mini-fluorometer (Promega, Madison,
29 Wisconsin). The Rapid Library quality was assessed with an Agilent Bioanalyzer High
30 Sensitivity DNA chip (Agilent Technologies, Santa Clara, California). The average
31 fragment length was between 600 bp and 900 bp, with the lower size cut-off at less than
32 10% below 350 bp. Pooled DNAs were titrated to obtain the optimal copies per bead
33 (cpb). After titration, 3 cpb was chosen as the best DNA and bead ratio and
34 corresponding amounts of DNA were added to the subsequent emPCR reactions.
35 EmPCR was performed with the 454/Roche Lib-L (LV) kits following manufacturer's
36 protocol for the Roche 454 GS FLX Titanium.

37 *Deep pyrosequencing and data processing*

38 The emPCR products were sequenced using Roche-454 GS FLX Titanium
39 chemistry. PBCV-1 DNA was sequenced twice giving a total of 98,258 quality-filtered
40 reads. Raw reads were filtered according to length and quality criteria. Filter-pass reads
41 were parsed into sample-barcoded bins and ported to a local database

42 The raw data from the 454 pyrosequencing machine were first processed through
43 a quality filter that removed sequence reads that did not meet the following criteria: i) A
44 complete forward primer and barcode. ii) ≤ 2 "N" in a sequence read, where N is

45 equivalent to an interrupted and resumed signals from sequential flows. iii) 200 nt \leq
46 sequence length \leq 500 nt. iv) Average quality score \geq 20. After filtering, each read was
47 trimmed to remove 3' adapter and primer sequences and was parsed by a MID adaptor
48 barcode. The corresponding QUAL file also was updated to remove quality scores from
49 reads not passing quality filters. The files are associated with sample information in a
50 hierarchical manner in MySQL tables. The processed data and the MySQL database
51 tables were stored on a database server and shared within the assembly/annotation team.

52 *Genome assembly*

53 Genome assembly was achieved using both automated pipeline and hand-curation
54 methods. Initially, the sff files were processed with the 454 Life Sciences GS De novo
55 Assembler software (program version 2.3). The sample was passed through the program
56 using default settings. Adjustments to those default settings gave no significant
57 improvement in the assembled sequence. The sample assembled into 12 contigs.
58 Additionally, 454 Life Science GS Mapper software was used with the original PBCV-1
59 genome used as the reference sequence; this too resulted in multiple contigs (>12)
60 assigned to PBCV-1.

61 The average sequence depth of coverage was >40X over the entire PBCV-1
62 genome from duplicate sequencing runs; however, some regions had approximately 100X
63 coverage while there was no coverage in other regions (i. e., 11 contig gaps). As part of
64 this sequencing project, 10 PBCV-1 antigenic mutants were also sequenced (samples
65 prepared and processed as described above). The mutants were also passed through the
66 GS De Novo Assembler and Mapper software programs. Both methods yielded multiple

67 contigs for each viral genome. One persistent major gap region observed in all of the
68 PBCV-1 isolates involved the *a122/123r* gene region, which contains highly repetitive
69 sequences, perhaps explaining why the gsMapper and gsAssembler programs were
70 unable to assign sequencing reads to this region. This region was re-sequenced with
71 Sanger-based methods and the revised *a122/123r* gene sequence was used to fill in the
72 gap between these two adjacent contigs. Sequencing the PBCV-1 antigenic mutants also
73 proved helpful, because we obtained at least one read from at least one antigenic mutant
74 for each of the remaining gaps. The final sequencing problem occurred at the 5' and 3'
75 ends of the sequence, which are reverse complementary repeats of one another (5). One
76 of the PBCV-1 antigenic mutants had reads that extended to near one end of the genome.
77 This region was reverse complemented and used to fill in the gap at the other end of the
78 genome.

79 ***Identification of the PBCV-1 contigs***

80 The PBCV-1 contigs were identified by BLAST searches (1) against the NCBI
81 non-redundant/nucleotide database limited to Chlorovirus (tax id: 181083). The contigs
82 were assumed to match viruses that infect *C. variabilis* (PBCV-1, NY-2A or AR158).
83 Each contig was assigned to a virus based on the best BLAST hit. Most of the reads
84 (99.28%) fit PBCV-1, seven relatively small contigs were ambiguous as to their BLAST
85 search and discarded.

86 ***Genome annotation***

87 The genome annotation was conducted using a semi-automatic in-house
88 annotation pipeline. The majority of protein-coding genes were annotated with the

89 Genome Annotation Transfer Utility (GATU) tool using the PBCV-1 genome and gene
90 calls (Refseq: NC 000852.4) as the reference genome (7). The GATU tool was used
91 iteratively, initially to accept ORFs that were substantially unchanged (100% identity,
92 Unassigned ORFs >95% identity), creating a list of 613 ORFs. The second pass
93 identified ORFs that had substantially changed, e.g., a few cases two adjacent ORFs were
94 found to be a single ORF due to a single nucleotide polymorphism (SNP) resulting in a
95 stop-codon. The third pass selected all New Annotations and all Unassigned-ORFs with
96 percent identity values above zero. Where two ORFs had identical starts or stops, the
97 larger ORF was selected and the smaller one was not considered further. In the fourth
98 iteration, GATU was used where the "third iteration" with all ORFs (both assigned and
99 unassigned) was the reference sequence and the newly 454 sequenced and assembled
100 PBCV-1 was annotated. The result was a list of New Annotations of the ORFs, with no
101 Unassigned-ORFs. These were evaluated by eye to determine if there were overlapping
102 and in-frame ORFs; there were none. The list contained 802 ORFs. These gene calls
103 were then examined for predicted function.

104 Functional annotations (gene and protein properties) of the called genes were
105 assigned to give both qualitative and quantitative data for the determined ORFs. Transfer
106 RNAs were predicted using the tRNAscan-SE software (3). Gene calls were checked
107 using the following 5-step process: i) putative homologs were identified by searching the
108 protein sequences against the Swiss-Prot (8) and RefSeq (9) protein databases using the
109 BLASTp (1) program (e-value < 1e-5). Because of very strong similarities between *C.*
110 *variabilis* NC64A infecting viruses, hits from PBCV-1, NY2A and AR158 viruses were
111 discarded in RefSeq while only previously existing entries from PBCV-1 were ignored in

112 the Swiss-Prot alignments. Only the best hit from the two databases was kept and
113 assigned as possible homologs of the predicted ORFs. ii) KEGG terms were assigned by
114 submitting the entire predicted proteome to the KEGG Automatic Annotation Server
115 [KAAS] (4), using the Bidirectionnal Best Hit method with an extended "Representative
116 for GENE" reference dataset, combining 30 complete genomes ranging from eukaryotes
117 to prokaryotes. iii) Genes were also assigned to a functional cluster using the BLASTp
118 program against the Clusters of Orthologous Groups [COG] (6) dataset and assigning the
119 best-hit's cluster to the query protein (e-value < 1e-5). iv) Protein domain/family
120 predictions were made using PfamScan software (e-value < 1e-5), keeping only matches
121 from the manually curated portion of the Pfam database (2). v) And finally, annotations
122 were manually added to genes when experimental evidence from the literature supported
123 the existence of a protein or characterized its function.

124 Table S1. PBCV-1 genes and gene annotations

Gene	Start	End	%A T	nt	aa	Da	pI	Stage	Virionic	COG	PfamA	BLASTp	Swissprot	Product	TM prediction ¹		
															T	H	P
a001L	280	549	68	270	89	10191	9.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A002L	512	1063	66	552	183	21545	9.4	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
a002aR	1022	1177	66	156	51	5820	9.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A002bL	1174	1335	56	162	53	6037	11.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A002cR	1367	1513	14	147	48	5331	11.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a002dR	1408	1539	12	132	43	4606	12.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A003R	1792	2217	70	426	141	15974	8.5	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	1	1
a004L	1891	2127	58	237	78	8746	9.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a004aL	2221	2361	52	141	46	5788	10.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A005R	2288	3094	66	807	268	28726	6.1	Early	N/A	COG0666 [FOG: Ankyrin repeat] [1e-05]	PF00023.23 [Ankyrin repeat] [6.3e-11]	YP_001426468 [hypothetical protein FR483_N836L (Paramecium bursaria Chlorella virus FR483)] [2e-44]	Q54KA7 [RecName: FullAnkyrin repeat PH and SEC7 domain containing protein secG] [1e-29]	Unknown protein	0	0	0
a006L	2393	2611	60	219	72	8415	8.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A007/008L	3292	4701	84	1410	469	51172	9.2	Early	N/A	COG0666 [FOG: Ankyrin repeat] [6e-06]	PF00023.23 [Ankyrin repeat] [1.1e-07]	XP_749852 [Pfs NACHT and Ankyrin domain protein (Aspergillus fumigatus Af293)] [2e-55]	Q502K3 [RecName: FullSerine/threonine-protein phosphatase 6 regulatory ankyrin repeat subunit C ShortSerine/threonine-protein phosphatase 6 regulatory subunit ARS-C ShortPP6-ARS-C] [2e-48]	Unknown protein	0	0	0
A009R	4998	5735	84	738	245	27634	8.1	Early	N/A	N/A	PF07150.4 [Protein of unknown function (DUF1390)] [7.8e-89]	YP_001426165 [hypothetical protein FR483_N533L (Paramecium bursaria Chlorella virus FR483)] [2e-80]	N/A	Unknown protein	0	0	0
A010R	5768	6973	88	1206	401	44998	5.2	Late	Yes	N/A	PF04451.5 [Large eukaryotic DNA virus major capsid protein] [1.9e-50]	YP_001426987 [hypothetical protein ATCV1_Z506L (Acanthocystis turfacea Chlorella virus 1)] [1e-89]	A7U6E9 [RecName: Fullmajor capsid protein ShortMCP] [8e-45]	Capsid protein	0	0	0
a010aR	6837	7022	60	186	61	7342	11.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A011L	6970	8181	84	1212	403	45076	5.4	Late	Yes	N/A	PF04451.5 [Large eukaryotic	YP_001426987 [hypothetical protein	A7U6E9 [RecName: Fullmajor capsid	Capsid protein	0	0	0

												DNA virus major capsid protein [2.9e-61]	ATCV1_Z506L (Acanthocystis turfacea Chlorella virus 1) [1e-106]	protein ShortMCP] [2e-43]				
a012R	7897	8190	64	294	97	11450	8.3	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a013L	8230	8571	58	342	113	12408	9.7	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A014R	8255	12364	40	4110	1369	141382	6.3	Late	Yes	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
a015L	8863	9411	48	549	182	20131	4.0	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	3	0
a016L	9808	10500	64	693	230	25288	4.4	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	3	4	3
a017L	11371	11571	54	201	66	7325	4.8	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A018L	12367	16374	80	4008	1335	####	4.9	Late	Yes	N/A	PF06598.4 [Chlorovirus glycoprotein repeat] [1.2e-11]	N/A	N/A	N/A	Unknown protein	0	0	0
a019R	12869	13519	60	651	216	23794	5.7	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	3	0
a020R	13889	14143	62	255	84	9524	10.8	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	1
a021R	14165	14632	68	468	155	17744	3.9	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	2	0
a022R	14702	15031	58	330	109	12127	4.6	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	2	0
a023R	15482	15706	70	225	74	8102	6.6	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	2
a024R	15770	16132	64	363	120	13384	3.7	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	1
A025/027/029L	16432	20511	78	4080	1359	####	4.4	Late	Yes	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
a026R	16916	17512	72	597	198	21885	5.1	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	5	5	5
a030R	18185	18721	72	537	178	20284	4.2	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	3	2
a031R	18755	19030	54	276	91	10478	4.1	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	1
a032R	19232	19810	68	579	192	21498	5.2	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	5	3
a033R	20189	20428	66	240	79	8651	10.4	N/A	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A034R	20572	21498	18	927	308	35163	10.4	Late	Yes	N/A	PF00069.18 [Protein kinase domain] [1.4e-07]	N/A	N/A	N/A	Protein kinase	0	1	0
A035L	21500	23260	88	1761	586	65606	8.9	Late	Yes	N/A	N/A	YP_001426757 [Hypothetical protein ATCV1_Z276L (Acanthocystis turfacea Chlorella virus 1)] [1e-10]	N/A	N/A	Unknown protein	0	1	0

a036R	22696	2303 7	56	342	113	12942	8.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A037L	23288	2360 5	68	318	105	12822	9.8	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
a038R	23584	2382 3	68	240	79	9448	10.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A039L	23623	2407 8	80	456	151	17102	4.1	Early	N/A	COG5201 [SCF ubiquitin ligase SKP1 component] [6e-16]	PF03931.8 [Skp1 family tetramerisation domain] [5.5e-19] / PF01466.12 [Skp1 family dimerisation domain] [6.8e-31]	XP_001753031 [predicted protein (Physcomitrella patens subsp. patens)] [2e-27]	Q9LNT9 [RecName: FullSKP1-like protein 4 ShortAtSK4] [9e-26]	Unknown protein	0	1	0
a040L	24087	2447 6	56	390	129	12234	3.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	0
A041R	24148	2538 6	34	1239	412	44315	10.8	Late	Yes	N/A	N/A	YP_001426427 [hypothetical protein FR483_N795L (Paramecium bursaria Chlorella virus FR483)] [1e-116]	N/A	Unknown protein	0	1	0
a042L	24238	2450 4	32	267	88	11031	12.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a043R	24798	2502 5	64	228	75	9172	9.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a044aL	25328	2549 2	66	165	54	6518	12.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A044L	25383	2718 2	84	1800	599	67849	6.4	Early	N/A	COG0464 [ATPases of the AAA+ class] [8e-06]	PF00004.22 [ATPase family associated with various cellular activities (AAA)] [1.7e-10]	XP_810119 [hypothetical protein (Trypanosoma cruzi strain CL Brener)] [6e-21]	Q5UR45 [RecName: FullPutative AAA family ATPase L572] [6e-17]	Unknown protein	0	0	0
a045R	26328	2654 0	52	213	70	8446	11.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a046R	26720	2722 0	68	501	166	19641	11.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a047L	26830	2703 3	64	204	67	7734	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a047aL	27217	2738 7	58	171	56	6491	4.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A048R	27248	2761 9	18	372	123	14017	7.6	Early-Late	N/A	N/A	N/A	YP_001425805 [hypothetical protein FR483_N173R (Paramecium bursaria Chlorella virus FR483)] [2e-19]	N/A	Unknown protein	0	0	0
A049L	27610	2826 9	70	660	219	25062	8.0	Early-Late	N/A	COG0584 [Glycerophosphoryl diester phosphodiesterase] [5e-06]	PF03009.10 [Glycerophosphoryl diester phosphodiesterase family] [3.8e-43]	YP_001426558 [hypothetical protein ATCV1_Z077L (Acanthocystis turfacea Chlorella virus 1)] [5e-48]	O07592 [RecName: FullPutative glycerophosphoryl diester phosphodiesterase yhdW ShortGlycerophosphodiester phosphodiesterase] [3e-23]	Unknown protein	0	0	0

A050L	28286	2871 1	84	426	141	16481	9.9	Early	N/A	N/A	PF03013.7 [Pyrimidine dimer DNA glycosylase] [9.7e-45]	YP_001426261 [hypothetical protein FR483_N629L (Paramecium bursaria Chlorella virus FR483)] [2e-49]	P04418 [RecName: FullEndonuclease V] [2e-21]	Pyrimidine dimer- specific glycosylase	0	0	0
A050aL	28821	2896 1	84	141	46	5443	8.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A051L	28984	2959 5	72	612	203	22804	8.6	Late	Yes	N/A	N/A	YP_001427069 [hypothetical protein ATCV1_Z588L (Acanthocystis turfacea Chlorella virus 1)] [2e- 49]	N/A	Unknown protein	1	2	1
a052R	29167	2941 8	62	252	83	10100	9.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A053R	29684	3077 5	62	1092	363	40175	6.4	Early	N/A	COG1052 [Lactate dehydrogenase and related dehydrogenases] [1e-05]	PF00389.23 [D- isomer specific 2- hydroxyacid dehydrogenase catalytic domain] [1.4e-34] / PF02826.12 [D- isomer specific 2- hydroxyacid dehydrogenase NAD binding domain] [3.5e-56]	YP_001426776 [hypothetical protein ATCV1_Z295L (Acanthocystis turfacea Chlorella virus 1)] [1e- 124]	P52643 [RecName: FullD-lactate dehydrogenase ShortD-LDH AltName: FullFermentative lactate dehydrogenase] [7e- 67]	Unknown protein	0	0	0
a054L	29800	3012 3	56	324	107	11750	10.2	N/A	N/A	N/A	N/A	YP_001426779 [hypothetical protein ATCV1_z298R (Acanthocystis turfacea Chlorella virus 1)] [9e- 05]	N/A	Hypothetical protein	0	0	0
a055L	30008	3040 3	62	396	131	15734	8.3	N/A	N/A	N/A	N/A	YP_001426778 [hypothetical protein ATCV1_z297R (Acanthocystis turfacea Chlorella virus 1)] [6e- 18]	N/A	Hypothetical protein	0	0	0
a056L	30289	3060 3	58	315	104	11511	11.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A058L	30976	3162 3	58	648	215	24718	4.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	2	0
a057R	31034	3144 7	74	414	137	14937	11.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A057aR	31650	3266 6	38	1017	338	37918	4.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a059L	31779	3200 3	52	225	74	8442	8.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A060L	32796	3350 0	86	705	234	27189	4.9	Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A061L	33529	3415 8	74	630	209	24425	4.7	Late	N/A	N/A	PF05711.4 [Macrocin-O- methyltransferase (TyfF)] [8.9e-20]	YP_003300173 [macrocin-O- methyltransferase domain-containing protein (Thermomonospora curvata DSM 43183)] [9e-31]	N/A	Unknown protein	0	0	0

a062R	33808	34179	60	372	123	14812	9.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
a062aR	34016	34198	54	183	60	6727	8.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
A063L	34191	34889	96	699	232	27050	6.0	Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A064R	34956	36872	24	1917	638	74913	4.9	Early-Late	N/A	N/A	PF05637.5 [galactosyl transferase GMA12/MNN10 family] [1.2e-10]	ZP_04834875 [conserved hypothetical protein (Prevotella melaninogenica ATCC 25845)] [6e-15]	N/A	UDP-Glucose glycosyltransferase	0	0	0
a065L	35265	35510	60	246	81	9861	7.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a066L	36607	36945	70	339	112	13228	10.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	0	1
a066aL	36851	36976	58	126	41	4846	8.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A067R	37043	37972	62	930	309	33211	3.6	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	1	0
a068L	37375	37677	66	303	100	9518	3.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a069L	37433	37873	56	441	146	15049	11.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	0
a070L	37798	38064	66	267	88	10265	9.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A071R	38003	39067	80	1065	354	41967	7.4	Early	N/A	N/A	N/A	ZP_06054798 [TPR repeat-containing protein (alpha proteobacterium HIMB114)] [8e-16]	N/A	Unknown protein	0	0	0
a072L	38164	38436	64	273	90	10264	11.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
a073L	38417	38626	56	210	69	8105	7.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a074L	38660	38863	74	204	67	8104	6.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A075L	39078	39920	80	843	280	33254	5.4	Early-Late	N/A	N/A	PF03016.8 [Exostosin family] [5.5e-09]	YP_001425644 [hypothetical protein FR483_N012L (Paramecium bursaria Chlorella virus FR483)] [3e-58]	N/A	Unknown protein	0	0	0
a075aR	39784	39933	60	150	49	5568	9.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A075bL	39950	40138	80	189	62	7489	4.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A075cR	40014	40172	70	159	52	5931	9.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
A076L	40190	40501	50	312	103	12094	11.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A077L	40470	40745	76	276	91	10180	5.0	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	1	1	1
A078R	40863	41759	74	897	298	33179	5.7	Early	N/A	COG0388 [Predicted amidohydrolase]	PF00795.15 [Carbon-nitrogen hydrolase] [5.5e-	YP_001425727 [hypothetical protein FR483_N095L	Q3HVN1 [RecName: FullIN-carbamoylputrescine	N-carbamoylputrescine amidohydrolase	0	2	0

										[7e-06]	34]	(Paramecium bursaria Chlorella virus FR483) [1e-113]	amidase] [2e-73]				
A078aL	41756	4187 8	92	123	40	4848	10.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A078bR	42007	4218 3	52	177	58	6943	10.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a078cL	42431	4256 5	62	135	44	5301	9.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	0
A079R	42467	4319 5	80	729	242	27092	9.3	Early	N/A	N/A	PF07150.4 [Protein of unknown function (DUF1390)] [5.4e-78]	YP_001425662 [hypothetical protein FR483_N030R (Paramecium bursaria Chlorella virus FR483)] [5e-29]	N/A	Unknown protein	0	0	0
a080L	42638	4304 5	58	408	135	16179	10.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	2	0
A081L	43192	4376 1	76	570	189	21916	9.0	Early	N/A	N/A	N/A	YP_001425639 [hypothetical protein FR483_N007L (Paramecium bursaria Chlorella virus FR483)] [5e-23]	N/A	Unknown protein	0	0	0
a082R	43760	4397 5	60	216	71	8434	6.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a083R	43791	4406 3	4	273	90	9976	10.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A084L	43811	4437 7	76	567	188	22309	7.0	Early	N/A	N/A	N/A	YP_001425642 [hypothetical protein FR483_N010L (Paramecium bursaria Chlorella virus FR483)] [7e-11]	N/A	Unknown protein	0	0	0
a084aL	44458	4458 0	46	123	40	4742	9.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A085R	44498	4522 6	66	729	242	27812	7.8	Late	Yes	N/A	PF03171.13 [2OG-Fe(II) oxygenase superfamily] [3.5e-11]	YP_001426050 [hypothetical protein FR483_N418R (Paramecium bursaria Chlorella virus FR483)] [7e-56]	Q5UP57 [RecName: FullPutative prolyl 4- hydroxylase Short4- PH AltName: FullProcollagen- proline2-oxoglutarate- 4-dioxygenase] [2e- 14]	Prolyl 4- hydroxylase	1	1	1
a086L	44953	4519 5	60	243	80	8487	10.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a086aL	45101	4522 9	54	129	42	5047	10.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A087R	45223	4662 3	32	1401	466	53644	8.5	Early	N/A	N/A	PF01844.16 [HNH endonuclease] [6.1e-13]	YP_001426447 [hypothetical protein FR483_N815L (Paramecium bursaria Chlorella virus FR483)] [4e-07]	N/A	Unknown protein	0	1	0
a086bL	46489	4663 2	72	144	47	5602	9.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a086cL	46508	4663 9	14	132	43	4980	4.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A088R	46755	4752 8	72	774	257	29442	5.0	Early	N/A	N/A	N/A	YP_001426459 [hypothetical protein]	N/A	Unknown protein	0	0	0

												FR483_N827R (Paramecium bursaria Chlorella virus FR483) [7e-07]						
A089R	47628	4783 4	78	207	68	7634	10.6	Early- Late	N/A	N/A	N/A	YP_001426713 [hypothetical protein ATCV1_Z232L (Acanthocystis turfacea Chlorella virus 1)] [2e- 14]	N/A	Unknown protein	0	0	0	
a089aL	47826	4797 2	76	147	48	5538	8.5	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0		
A090R	47851	4831 5	70	465	154	18442	4.9	Early- Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0	
A092/093L	48326	4962 7	72	1302	433	49577	10.7	Early- Late	Yes	N/A	PF08789.3 [PBCV-specific basic adaptor domain] [1.2e-15]	YP_001427042 [hypothetical protein ATCV1_Z561L (Acanthocystis turfacea Chlorella virus 1)] [2e- 90]	N/A	Unknown protein	0	0	0	
a092/093a L	48547	4873 2	68	186	61	7472	11.8	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0		
A094L	49662	5075 6	74	1095	364	40734	4.3	Early	N/A	COG2273 [Beta- glucanase/Beta- glucan synthetase] [4e- 06]	PF00722.14 [Glycosyl hydrolases family 16] [7.5e-13]	YP_001425751 [hypothetical protein FR483_N119L (Paramecium bursaria Chlorella virus FR483)] [1e-110]	P23903 [RecName: FullGlucan endo-13- beta-glucosidase A1 AltName: Full(1->3)- beta-glucan endohydrolase AltName: Full(1->3)- beta-glucanase A1 Flags: Precursor] [1e- 22]	beta-1-3-glucanase	1	1	1	
a095R	50014	5023 2	52	219	72	7701	10.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
a096R	50346	5054 9	68	204	67	7873	12.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0	
a097R	50678	5089 3	68	216	71	8581	9.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1	
A098R	50886	5259 2	76	1707	568	65159	8.2	Early	N/A	COG1215 [Glycosyltransfer ases probably involved in cell wall biogenesis] [4e-06]	PF03142.8 [Chitin synthase] [4.4e-09]	YP_001425756 [hypothetical protein FR483_N124R (Paramecium bursaria Chlorella virus FR483)] [0.0]	B1WB39 [RecName: FullHyaluronan synthase 1 AltName: FullHyaluronate synthase 1 AltName: FullHyaluronic acid synthase 1 ShortHA synthase 1] [1e-56]	Hyaluronan synthase	7	8	7	
a099L	52152	5239 4	62	243	80	8644	9.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
a099aL	52539	5268 2	60	144	47	5250	6.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
A100R	52691	5447 8	60	1788	595	65408	6.1	Early	N/A	COG0449 [Glucosamine 6- phosphate synthetase contains amidotransferase and phosphosugar isomerase domains] [8e-07]	PF00310.14 [Glutamine amidotransferase s class-II] [5.6e- 15] / PF01380.15 [SIS domain] [2.2e-26]	YP_001425667 [hypothetical protein FR483_N035R (Paramecium bursaria Chlorella virus FR483)] [0.0]	Q7WE36 [RecName: FullGlucosamine-- fructose-6-phosphate aminotransferase (isomerizing)] [1e- 153]	Glutamine:fructose -6-phosphate amidotransferase	0	0	0	

a101L	52925	5318 8	54	264	87	10061	8.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a102L	54468	5497 4	56	507	168	19199	11.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	1
A103R	54622	5561 4	84	993	330	37832	8.4	Early	N/A	COG5226 [mRNA capping enzyme guanylyltransferase (alpha) subunit] [4e-13]	PF01331.12 [mRNA capping enzyme catalytic domain] [2.1e-50] / PF03919.8 [mRNA capping enzyme C-terminal domain] [7.8e-15]	YP_001427200 [hypothetical protein ATCV1_Z719L (Acanthocystis turfacea Chlorella virus 1)] [2e-90]	P78587 [RecName: FullmRNA-capping enzyme subunit alpha AltName: FullmRNA guanylyltransferase AltName: FullGTP--RNA guanylyltransferase ShortGTase] [7e-16]	mRNA guanylyltransferase	0	3	0
a104L	55054	5534 4	58	291	96	10912	7.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A105L	55626	5648 0	84	855	284	33063	6.8	Early	N/A	N/A	PF00443.22 [Ubiquitin carboxyl-terminal hydrolase] [5.7e-25]	YP_001425764 [hypothetical protein FR483_N132L (Paramecium bursaria Chlorella virus FR483)] [8e-70]	P40818 [RecName: FullUbiquitin carboxyl-terminal hydrolase 8 AltName: FullUbiquitin thioesterase 8 AltName: FullUbiquitin-specific-processing protease 8 AltName: FullDeubiquitinating enzyme 8 AltName: FullUbiquitin isopeptidase Y ShorthUBPy] [3e-09]	Unknown protein	0	0	0
a106R	55770	5616 8	56	399	132	15709	9.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A107L	56516	5738 8	80	873	290	32511	6.2	Early	N/A	N/A	PF00382.12 [Transcription factor TFIIIB repeat] [4.6e-06]	YP_001425767 [hypothetical protein FR483_N135L (Paramecium bursaria Chlorella virus FR483)] [2e-57]	Q9SS44 [RecName: FullTranscription initiation factor IIB-2 AltName: FullGeneral transcription factor TFIIIB-2 ShortATFIIB2] [4e-10]	Unknown protein	0	0	0
a108R	56933	5714 8	60	216	71	8190	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A108aR	57212	5740 0	52	189	62	7278	9.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A108bL	57471	5797 7	84	507	168	19692	7.1	N/A	N/A	N/A	PF01541.17 [GIY-YIG catalytic domain] [1.9e-06]	YP_001427194 [hypothetical protein ATCV1_Z713L (Acanthocystis turfacea Chlorella virus 1)] [1e-54]	N/A	Hypothetical protein	0	0	0
a110L	58082	5839 0	50	309	102	11775	10.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A111/114R	58098	6068 0	80	2583	860	98045	6.6	Early	N/A	N/A	PF00852.12 [Glycosyltransferase family 10 (fucosyltransferase)] [1.4e-06]	YP_001426104 [hypothetical protein FR483_N472R (Paramecium bursaria Chlorella virus FR483)] [0.0]	N/A	Unknown protein	0	0	0
a112L	58775	5906 8	42	294	97	10605	9.2	N/A	N/A	N/A	N/A	YP_001426602 [hypothetical protein ATCV1_z121L (Acanthocystis turfacea	N/A	Hypothetical protein	1	2	1

a126R	66620	66820	24	201	66	7964	11.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	2	2
A127R	67154	67891	14	738	245	27126	10.1	Late	Yes	N/A	N/A	YP_001425800 [hypothetical protein FR483_N168R (Paramecium bursaria Chlorella virus FR483)] [1e-76]	N/A	Unknown protein	0	0	0
a128L	67510	67770	56	261	86	9678	4.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A129R	67952	69028	82	1077	358	40533	8.4	Early-Late	N/A	N/A	N/A	YP_001425778 [hypothetical protein FR483_N146R (Paramecium bursaria Chlorella virus FR483)] [8e-69]	N/A	Unknown protein	0	0	0
A130R	69049	69366	84	318	105	11644	10.0	Early-Late	N/A	N/A	N/A	YP_001426755 [hypothetical protein ATCV1_Z274R (Acanthocystis turfacea Chlorella virus 1)] [3e-24]	N/A	Unknown protein	4	3	3
A130aR	69261	69404	62	144	47	5585	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a130bR	69344	69475	58	132	43	5215	10.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A131L	69359	69769	68	411	136	16124	4.6	Early	N/A	N/A	N/A	YP_001426009 [hypothetical protein FR483_N377L (Paramecium bursaria Chlorella virus FR483)] [4e-06]	N/A	Unknown protein	0	0	0
a132R	69533	69805	54	273	90	10395	5.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
A133R	69960	70583	86	624	207	23480	9.6	Early	N/A	N/A	PF11264.1 [Thylakoid formation protein] [4.8e-52]	XP_001693657 [inositol phosphatase-like protein (Chlamydomonas reinhardtii)] [1e-24]	Q84PB7 [RecName: FullProtein THYLAKOID FORMATION1 chloroplastic Flags: Precursor] [5e-23]	Unknown protein	0	1	0
A134L	70558	71055	92	498	165	19545	9.9	Early-Late	N/A	N/A	PF01541.17 [GIY-YIG catalytic domain] [1.8e-11]	YP_001425818 [hypothetical protein FR483_N186L (Paramecium bursaria Chlorella virus FR483)] [2e-23]	N/A	Unknown protein	0	0	0
A135L	71055	71603	74	549	182	20728	8.4	Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	5	5	5
A136R	71134	71574	30	441	146	16367	11.5	N/A	Yes	N/A	N/A	YP_001425749 [hypothetical protein FR483_N117R (Paramecium bursaria Chlorella virus FR483)] [6e-07]	N/A	Unknown protein	0	0	0
A137R	71625	71843	44	219	72	8777	10.9	Early	N/A	N/A	N/A	YP_001427214 [hypothetical protein ATCV1_z733L (Acanthocystis turfacea Chlorella virus 1)] [2e-07]	N/A	Unknown protein	0	0	0
A138R	71880	72701	84	822	273	31471	4.4	Early-Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0

a138aR	72604	7274 7	72	144	47	5766	9.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A139L	72698	7315 3	60	456	151	17701	8.4	Late	Yes	N/A	N/A	YP_001425821 [hypothetical protein FR483_N189L (Paramecium bursaria Chlorella virus FR483)] [6e-24]	N/A	Unknown protein	2	2	2
A140/145R	73107	7649 0	90	3384	1127	####	11.0	Early- Late	Yes	N/A	N/A	YP_001425827 [hypothetical protein FR483_N195R (Paramecium bursaria Chlorella virus FR483)] [0.0]	N/A	Unknown protein	0	1	0
a144L	74244	7574 0	64	1497	498	57630	8.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	4	4
a142L	74474	7487 5	48	402	133	13511	8.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	4	4	4
a146L	75752	7600 6	68	255	84	8946	3.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
a147L	75882	7665 5	56	774	257	29267	8.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	5	1
A148R	76540	7687 2	82	333	110	11990	9.8	Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
a149L	76642	7705 2	62	411	136	16083	11.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A150L	76856	7731 4	74	459	152	17089	4.1	Early	N/A	N/A	N/A	YP_001425837 [hypothetical protein FR483_N205L (Paramecium bursaria Chlorella virus FR483)] [4e-11]	N/A	Unknown protein	0	0	0
A151R	77397	7780 4	52	408	135	15099	8.3	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
a152L	77822	7809 4	58	273	90	10135	11.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
A153R	77880	7925 9	80	1380	459	51301	9.2	Early- Late	N/A	COG1061 [DNA or RNA helicases of superfamily II] [2e-06]	PF04851.8 [Type III restriction enzyme res subunit] [5.5e-15]	YP_001425838 [hypothetical protein FR483_N206R (Paramecium bursaria Chlorella virus FR483)] [1e-176]	Q5UQ46 [RecName: FullPutative ATP- dependent RNA helicase L396] [1e- 46]	Unknown protein	0	0	0
A154L	79256	8029 9	70	1044	347	38950	8.4	Early- Late	N/A	N/A	N/A	YP_001425778 [hypothetical protein FR483_N146R (Paramecium bursaria Chlorella virus FR483)] [5e-65]	N/A	Unknown protein	0	0	0
a155R	79412	7969 6	52	285	94	11209	10.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a156L	79885	8021 7	52	333	110	12610	11.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A157L	80393	8072 5	80	333	110	12328	3.9	Early- Late	Yes	N/A	N/A	YP_001425841 [hypothetical protein FR483_N209L (Paramecium bursaria Chlorella virus FR483)] [2e-16]	N/A	Unknown protein	1	1	1

A158L	80770	81084	68	315	104	12162	3.3	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A159R	81104	81436	36	333	110	12869	8.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
a160L	81257	81751	68	495	164	17774	10.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	4	4	4
A161R	81345	81716	58	372	123	13383	11.0	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A162L	81717	82952	78	1236	411	46696	4.4	Early-Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	3	3	3
a162aR	82831	82959	68	129	42	4676	7.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A163R	82995	84296	12	1302	433	48956	7.0	Early-Late	N/A	N/A	PF00060.19 [Ligand-gated ion channel] [1.8e-09]	N/A	N/A	Unknown protein	3	3	3
a164L	83873	84277	80	405	134	15629	9.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	3	1
A164aR	84314	84493	14	180	59	7094	5.8	N/A	Yes	N/A	N/A	N/A	N/A	Unknown protein	1	0	0
A165L	84486	84935	60	450	149	17251	6.8	Early-Late	N/A	N/A	N/A	YP_001425846 [hypothetical protein FR483_N214L (Paramecium bursaria Chlorella virus FR483)] [7e-12]	N/A	Unknown protein	0	0	1
A165aL	84835	85326	70	492	163	19024	10.1	N/A	Yes	N/A	N/A	YP_001425848 [hypothetical protein FR483_N216L (Paramecium bursaria Chlorella virus FR483)] [3e-18]	N/A	Unknown protein	0	0	0
A166R	85431	86237	80	807	268	31183	4.7	Early-Late	N/A	N/A	PF09588.3 [YqaJ-like viral recombinase domain] [7.9e-24]	YP_001425850 [hypothetical protein FR483_N218R (Paramecium bursaria Chlorella virus FR483)] [1e-81]	Q5UQV1 [RecName: FullUncharacterized protein R354] [5e-12]	Unknown protein	0	0	0
a167L	85677	85880	64	204	67	8077	8.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a167aR	86128	86283	68	156	51	6043	10.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A168R	86276	86776	86	501	166	18317	4.6	Late	Yes	N/A	N/A	YP_001427100 [hypothetical protein ATCV1_Z619L (Acanthocystis turfacea Chlorella virus 1)] [3e-19]	N/A	Unknown protein	1	1	1
A169R	86904	87875	74	972	323	36836	5.8	Early-Late	N/A	COG0078 [Ornithine carbamoyltransferase] [5e-07]	PF02729.14 [Aspartate/ornithine carbamoyltransferase carbamoyl-P binding domain] [2.4e-44] / PF00185.17 [Aspartate/ornithine carbamoyltransferase Asp/Om	NP_001148757 [LOC100282373 (Zea mays)] [7e-78]	Q43087 [RecName: FullAspartate carbamoyltransferase 2 chloroplastic AltName: FullAspartate transcarbamylase 2 ShortATCase 2 Flags: Precursor] [5e-79]	Aspartate transcarbamylase	0	0	0

A185R	94548	9739 0	40	2843	913	####	8.3	Early	N/A	COG0417 [DNA polymerase elongation subunit (family B)] [1e-07]	PF03104.12 [DNA polymerase family B exonuclease domain] [1.3e-70] / PF00136.14 [DNA polymerase family B] [1.8e-118]	YP_001425655 [hypothetical protein FR483_N023L (Paramecium bursaria Chlorella virus FR483)] [0.0]	P30320 [RecName: FullDNA polymerase] [0.0]	Unknown protein	0	0	0
a186L	94966	9522 0	46	255	84	9946	5.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a187L	95295	9603 8	58	744	247	28922	10.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a188L	96724	9695 7	56	234	77	8398	11.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A188aR	96944	9739 0	58	447	148	17326	10.0	N/A	Yes	COG0417 [DNA polymerase elongation subunit (family B)] [3e-07]	PF00136.14 [DNA polymerase family B] [6.5e-17]	YP_001427279 [hypothetical protein ATCV1_Z798L (Acanthocystis turfacea Chlorella virus 1)] [8e-60]	P30320 [RecName: FullDNA polymerase] [2e-72]	Unknown protein	0	0	0
a188bR	97258	9739 8	72	141	46	5774	11.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A189/192R	97433	1E+0 5	28	3900	1299	143575	11.4	Late	Yes	N/A	N/A	YP_001427275 [hypothetical protein ATCV1_Z794L (Acanthocystis turfacea Chlorella virus 1)] [1e-157]	N/A	Unknown protein	0	0	0
a190L	97525	9774 3	52	219	72	7634	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A193L	1E+05	1E+0 5	72	789	262	29627	4.7	Early-Late	N/A	COG0592 [DNA polymerase sliding clamp subunit (PCNA homolog)] [7e-06]	PF00705.11 [Proliferating cell nuclear antigen N-terminal domain] [6.4e-26] / PF02747.8 [Proliferating cell nuclear antigen C-terminal domain] [1.2e-21]	YP_001425651 [hypothetical protein FR483_N019R (Paramecium bursaria Chlorella virus FR483)] [1e-103]	O82134 [RecName: FullProliferating cell nuclear antigen] [7e-36]	Unknown protein	0	0	0
a194R	1E+05	1E+0 5	56	249	82	9602	10.7	N/A	N/A	N/A	N/A	YP_001425650 [hypothetical protein FR483_n018L (Paramecium bursaria Chlorella virus FR483)] [1e-12]	N/A	Hypothetical protein	0	0	0
a195R	1E+05	1E+0 5	66	249	82	9700	8.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A196L	1E+05	1E+0 5	90	459	152	17456	8.4	Late	Yes	N/A	N/A	YP_001425648 [hypothetical protein FR483_N016R (Paramecium bursaria Chlorella virus FR483)] [2e-44]	N/A	Unknown protein	3	3	1
a197R	1E+05	1E+0 5	62	288	95	11090	8.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a198R	1E+05	1E+0 5	62	225	74	8873	10.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A199R	1E+05	1E+0 5	6	306	101	10904	10.1	Late	N/A	N/A	N/A	YP_001427266 [hypothetical protein]	N/A	Unknown protein	2	3	2

												ATCV1_Z785L (Acanthocystis turfacea Chlorella virus 1) [1e- 12]						
A200R	1E+05	1E+0 5	76	357	118	13165	9.3	Early	N/A	N/A	PF00383.15 [Cytidine and deoxycytidylate deaminase zinc- binding region] [8.4e-13]	YP_001425646 [hypothetical protein FR483_N014L (Paramecium bursaria Chlorella virus FR483)] [4e-39]	N/A	Unknown protein	0	0	0	
a200aR	1E+05	1E+0 5	52	165	54	6424	8.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
A201L	1E+05	1E+0 5	74	285	94	10005	10.7	Early- Late	Yes	N/A	N/A	YP_001425713 [hypothetical protein FR483_N081L (Paramecium bursaria Chlorella virus FR483)] [3e-22]	N/A	Unknown protein	2	2	2	
A201aL	1E+05	1E+0 5	18	177	58	6787	8.8	N/A	Yes	N/A	N/A	N/A	N/A	Unknown protein	0	0	0	
A202L	1E+05	1E+0 5	90	342	113	12232	5.0	Early- Late	Yes	N/A	N/A	YP_001425949 [hypothetical protein FR483_N317R (Paramecium bursaria Chlorella virus FR483)] [2e-25]	N/A	Unknown protein	0	0	0	
A203R	1E+05	1E+0 5	26	651	216	24011	6.0	Late	Yes	N/A	N/A	YP_001425947 [hypothetical protein FR483_N315L (Paramecium bursaria Chlorella virus FR483)] [2e-57]	N/A	Unknown protein	1	2	0	
a204L	1E+05	1E+0 5	62	240	79	8029	10.5	N/A	N/A	N/A	N/A	YP_001425948 [hypothetical protein FR483_n316R (Paramecium bursaria Chlorella virus FR483)] [9e-15]	N/A	Hypothetical protein	0	0	0	
a204aL	1E+05	1E+0 5	58	162	53	6061	4.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1	
A205R	1E+05	1E+0 5	80	621	206	22452	12.1	Late	Yes	N/A	PF08789.3 [PBCV-specific basic adaptor domain] [4.2e-16]	YP_001425945 [hypothetical protein FR483_N313L (Paramecium bursaria Chlorella virus FR483)] [6e-29]	N/A	Unknown protein	0	0	0	
a206L	1E+05	1E+0 5	64	258	85	10043	11.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
A207R	1E+05	1E+0 5	86	1119	372	41969	4.7	Early- Late	N/A	COG1166 [Arginine decarboxylase (spermidine biosynthesis)] [1e-06]	PF02784.9 [Pyridoxal- dependent decarboxylase pyridoxal binding domain] [3.3e-61] / PF00278.15 [Pyridoxal- dependent decarboxylase C- terminal sheet domain] [5.5e-23]	YP_001425944 [hypothetical protein FR483_N312L (Paramecium bursaria Chlorella virus FR483)] [1e-139]	P27117 [RecName: FullOrnithine decarboxylase ShortODC] [2e-74]	Arginine/Omithine decarboxylase	0	0	0	
A208R	1E+05	1E+0 5	76	936	311	35364	4.2	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0	

a210L	1E+05	1E+05	64	492	163	16631	5.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
a211R	1E+05	1E+05	56	324	107	12159	12.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	2
A212R	1E+05	1E+05	78	168	55	6345	12.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A213L	1E+05	1E+05	84	447	148	16483	4.5	Early-Late	Yes	N/A	N/A	YP_001425936 [hypothetical protein FR483_N304R (Paramecium bursaria Chlorella virus FR483)] [1e-30]	N/A	Unknown protein	1	1	1
a213aL	1E+05	1E+05	56	147	48	5567	12.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A214L	1E+05	1E+05	76	408	135	15424	7.1	Early	N/A	N/A	N/A	YP_001425935 [hypothetical protein FR483_N303R (Paramecium bursaria Chlorella virus FR483)] [9e-21]	N/A	Unknown protein	0	0	0
a214aL	1E+05	1E+05	46	144	47	5815	10.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	0	1
A215L	1E+05	1E+05	82	966	321	35707	10.1	Late	N/A	N/A	N/A	YP_001425925 [hypothetical protein FR483_N293R (Paramecium bursaria Chlorella virus FR483)] [1e-115]	N/A	Alkaline alginate lyase / 1-4 polyglucuronic acid lyase / polysaccharide lyase family 14 / vAL-1	0	0	0
a216R	1E+05	1E+05	62	324	107	11708	6.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	2
A217L	1E+05	1E+05	70	1185	394	45248	9.9	Early-Late	Yes	N/A	N/A	YP_001425921 [hypothetical protein FR483_N289R (Paramecium bursaria Chlorella virus FR483)] [1e-164]	N/A	Unknown protein	0	0	1
a218L	1E+05	1E+05	18	366	121	14878	10.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A219/222/226R	1E+05	1E+05	58	2034	677	77797	7.0	Early	Yes	COG1215 [Glycosyltransferases probably involved in cell wall biogenesis] [4e-06]	N/A	YP_001426347 [hypothetical protein FR483_N715L (Paramecium bursaria Chlorella virus FR483)] [0.0]	P58932 [RecName: FullCellulose synthase catalytic subunit (UDP-forming)] [6e-07]	Unknown protein	9	8	10
a220L	1E+05	1E+05	50	390	129	15951	8.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	0	1
a223R	1E+05	1E+05	60	213	70	7560	11.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a223aL	1E+05	1E+05	64	279	92	10021	8.5	N/A	N/A	N/A	N/A	YP_001426907 [hypothetical protein ATCV1_2426L (Acanthocystis turfacea Chlorella virus 1)] [4e-12]	N/A	Hypothetical protein	0	0	0
a224L	1E+05	1E+05	54	267	88	9977	10.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a225L	1E+05	1E+05	66	273	90	11049	6.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0

A227L	1E+05	1E+05	80	414	137	15689	10.0	Late	Yes	N/A	N/A	YP_001425876 [hypothetical protein FR483_N244R (Paramecium bursaria Chlorella virus FR483)] [6e-41]	N/A	Unknown protein	0	0	0
a228R	1E+05	1E+05	60	231	76	8736	9.7	N/A	N/A	N/A	N/A	YP_001425877 [hypothetical protein FR483_n245L (Paramecium bursaria Chlorella virus FR483)] [4e-11]	N/A	Hypothetical protein	0	1	0
a228aR	1E+05	1E+05	64	135	44	5254	10.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A229L	1E+05	1E+05	82	234	77	8329	8.5	Early-Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	2	2	2
A230R	1E+05	1E+05	36	591	196	22055	8.4	Late	Yes	N/A	N/A	YP_001425874 [hypothetical protein FR483_N242L (Paramecium bursaria Chlorella virus FR483)] [8e-34]	N/A	Unknown protein	4	4	4
A231L	1E+05	1E+05	80	1152	383	43644	9.9	Early-Late	Yes	N/A	N/A	YP_001426097 [hypothetical protein FR483_N465R (Paramecium bursaria Chlorella virus FR483)] [1e-137]	N/A	Unknown protein	1	0	0
a232R	1E+05	1E+05	60	264	87	10420	9.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a232aR	1E+05	1E+05	52	135	44	5103	10.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A233R	1E+05	1E+05	42	339	112	13599	9.3	Late	N/A	N/A	N/A	YP_001425872 [hypothetical protein FR483_N240L (Paramecium bursaria Chlorella virus FR483)] [1e-14]	N/A	Unknown protein	0	0	0
A234L	1E+05	1E+05	72	327	108	12388	10.4	Early-Late	N/A	N/A	N/A	YP_001425871 [hypothetical protein FR483_N239R (Paramecium bursaria Chlorella virus FR483)] [6e-21]	N/A	Unknown protein	0	0	0
a235R	1E+05	1E+05	54	252	83	9647	7.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a236L	1E+05	1E+05	68	264	87	10307	8.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	1	2
A237R	1E+05	1E+05	22	1557	518	58565	9.5	Late	Yes	COG5310 [Homospermidine synthase] [4e-23]	PF03435.11 [Saccharopine dehydrogenase] [1.2e-96]	YP_001425864 [hypothetical protein FR483_N232L (Paramecium bursaria Chlorella virus FR483)] [0.0]	Q98H64 [RecName: FullHomospermidine synthase ShortHSS] [2e-54]	Homospermidine synthase	0	0	0
a238L	1E+05	1E+05	46	345	114	12804	9.4	N/A	N/A	N/A	N/A	YP_001425865 [hypothetical protein FR483_n233R (Paramecium bursaria Chlorella virus FR483)] [3e-17]	N/A	Hypothetical protein	0	0	0
A239L	1E+05	1E+05	72	447	148	17052	5.0	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0

a240L	1E+05	1E+05	48	198	65	7734	12.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A240aL	1E+05	1E+05	72	186	61	7125	10.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	0
a240bR	1E+05	1E+05	68	141	46	5363	9.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
a240cL	1E+05	1E+05	52	165	54	6440	9.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a240dL	1E+05	1E+05	60	147	48	5633	6.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A241R	1E+05	1E+05	78	2178	725	82753	6.6	Early	N/A	COG1201 [Lhr-like helicases] [5e-06]	PF00270.22 [DEAD/DEAH box helicase] [7.3e-19] / PF08148.5 [DSHCT (NUC185) domain] [1.4e-15]	YP_001425859 [hypothetical protein FR483_N227L (Paramecium bursaria Chlorella virus FR483)] [0.0]	Q15477 [RecName: FullHelicase SK12W AltName: FullHelicase-like protein ShortHLP] [7e-87]	Unknown protein	0	0	0
a242L	1E+05	1E+05	62	354	117	13745	6.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	3	0
A243R	1E+05	1E+05	92	909	302	36101	10.3	Early-Late	N/A	N/A	N/A	YP_001425731 [hypothetical protein FR483_N099R (Paramecium bursaria Chlorella virus FR483)] [1e-134]	N/A	Unknown protein	0	0	0
a244L	1E+05	1E+05	44	234	77	8398	9.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A245R	1E+05	1E+05	80	564	187	19748	9.3	Late	Yes	COG2032 [Cu/Zn superoxide dismutase] [9e-07]	PF00080.13 [Copper/zinc superoxide dismutase (SODC)] [3.1e-49]	YP_001426671 [hypothetical protein ATCV1_Z190L (Acanthocystis turfacea Chlorella virus 1)] [1e-68]	Q751L8 [RecName: FullSuperoxide dismutase (Cu-Zn)] [8e-41]	Cu/Zn superoxide dismutase	1	1	0
A246R	1E+05	1E+05	82	342	113	12017	11.5	Late	Yes	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A246aR	1E+05	1E+05	56	459	152	16618	8.0	N/A	N/A	N/A	PF00967.10 [Barwin family] [3e-47]	NP_187123 [PR4 (PATHOGENESIS-RELATED 4) chitin binding (Arabidopsis thaliana)] [8e-10]	P43082 [RecName: FullHevein-like protein Flags: Precursor] [6e-11]	Hypothetical protein	0	0	0
A247R	1E+05	1E+05	84	1152	383	44592	6.1	Early-Late	N/A	COG0666 [FOG: Ankyrin repeat] [1e-09]	PF00023.23 [Ankyrin repeat] [8.6e-06]	YP_001958455 [hypothetical protein Aasi_1435 (Candidatus Amoebophilus asiaticus 5a2)] [9e-26]	Q4UMH6 [RecName: FullPutative ankyrin repeat protein RF_0381] [2e-20]	Unknown protein	0	0	0
A248R	1E+05	1E+05	70	927	308	35564	8.0	Early	N/A	COG0515 [Serine/threonine protein kinase] [1e-05]	PF00069.18 [Protein kinase domain] [2.4e-52]	YP_001426416 [hypothetical protein FR483_N784R (Paramecium bursaria Chlorella virus FR483)] [3e-61]	Q1ZXH2 [RecName: FullProbable serine/threonine-protein kinase fhkB AltName: FullForkhead-associated kinase protein B] [4e-24]	Protein kinase	0	0	0
a249L	1E+05	1E+05	70	309	102	11134	10.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A250R	1E+05	1E+05	82	285	94	10620	8.8	Early-Late	N/A	N/A	PF07885.9 [ion channel] [3.2e-11]	YP_001427066 [hypothetical protein ATCV1_Z585R (Acanthocystis turfacea Chlorella virus 1)] [4e-	N/A	Potassium ion channel protein (Kcv)	2	2	2

A261R	1E+05	1E+05	88	618	205	25196	8.7	Early-Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A262/263L	1E+05	1E+05	64	771	256	29470	9.6	N/A	Yes	N/A	N/A	YP_001425738 [hypothetical protein FR483_N106L (Paramecium bursaria Chlorella virus FR483)] [4e-46]	N/A	Unknown protein	2	3	2
a264R	1E+05	1E+05	84	204	67	7641	9.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	1
A265L	1E+05	1E+05	90	750	249	27947	9.3	Early-Late	N/A	N/A	PF04848.6 [Poxvirus A22 protein] [8.6e-11]	YP_001426640 [hypothetical protein ATCV1_Z159R (Acanthocystis turfatea Chlorella virus 1)] [5e-73]	N/A	Unknown protein	0	0	0
a266R	1E+05	1E+05	68	252	83	9838	9.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a266aR	1E+05	1E+05	58	126	41	4771	9.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A267L	1E+05	1E+05	80	945	314	36686	8.2	Early	N/A	N/A	N/A	YP_142777 [hypothetical protein MIM1_R423 (Acanthamoeba polyphaga mimivirus)] [2e-15]	Q5UQL9 [Uncharacterized protein R423] [2e-16]	Unknown protein	0	0	0
a268R	1E+05	1E+05	50	216	71	8208	9.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a269R	1E+05	1E+05	62	207	68	8054	8.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A271L	1E+05	1E+05	74	825	274	31114	7.1	Early-Late	Yes	COG2267 [Lysophospholipase] [1e-07]	PF12146.1 [Putative lysophospholipase] [1.9e-10]	YP_001426877 [hypothetical protein ATCV1_Z396R (Acanthocystis turfatea Chlorella virus 1)] [5e-51]	Q55EQ3 [RecName: FullUncharacterized abhydrolase domain-containing protein DDB_G0269086] [6e-08]	Unknown protein	0	3	0
a270R	1E+05	1E+05	80	282	93	10237	10.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	2	2
a272R	1E+05	1E+05	54	204	67	8074	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a272aR	1E+05	1E+05	58	240	79	9104	6.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A273L	1E+05	1E+05	88	417	138	15713	9.9	Late	Yes	N/A	PF03713.6 [Domain of unknown function (DUF305)] [6.8e-13]	YP_142507 [hypothetical protein MIM1_L153 (Acanthamoeba polyphaga mimivirus)] [4e-16]	Q5UPL7 [RecName: FullUncharacterized protein L153] [2e-17]	Unknown protein	3	3	3
A274R	1E+05	1E+05	16	792	263	31386	4.4	Early-Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A275R	1E+05	1E+05	80	759	252	28394	9.1	Early	N/A	N/A	PF07150.4 [Protein of unknown function (DUF1390)] [8.6e-79]	YP_001425662 [hypothetical protein FR483_N030R (Paramecium bursaria Chlorella virus FR483)] [2e-26]	N/A	Unknown protein	0	0	0
a276L	1E+05	1E+05	60	285	94	11687	10.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	2

A277L	1E+05	1E+05	70	912	303	34378	8.5	Late	N/A	COG0515 [Serine/threonine protein kinase] [1e-05]	PF00069.18 [Protein kinase domain] [2.2e-42]	YP_001426416 [hypothetical protein FR483_N784R (Paramecium bursaria Chlorella virus FR483)] [2e-28]	Q1ZXC8 [RecName: FullProbable serine/threonine- protein kinase pXi] [7e-23]	Protein kinase	0	0	0
A278L	1E+05	1E+05	80	1833	610	69231	10.8	Late	Yes	N/A	PF00069.18 [Protein kinase domain] [1.2e-07] / PF08789.3 [PBCV-specific basic adaptor domain] [7.5e-10]	YP_001427189 [hypothetical protein ATCV1_Z708L (Acanthocystis turfacea Chlorella virus 1)] [1e- 130]	Q99078 [RecName: FullDual specificity protein kinase FUZ7] [5e-07]	Protein kinase	0	1	0
a279R	1E+05	1E+05	68	219	72	8802	10.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	1
a280R	1E+05	1E+05	56	282	93	10103	3.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a281R	1E+05	1E+05	52	720	239	25998	4.6	N/A	N/A	N/A	N/A	YP_001426034 [hypothetical protein FR483_n402R (Paramecium bursaria Chlorella virus FR483)] [5e-25]	N/A	Hypothetical protein	0	6	0
A282L	1E+05	1E+05	60	1710	569	63371	10.8	Late	Yes	N/A	PF00069.18 [Protein kinase domain] [4.8e-10] / PF08789.3 [PBCV-specific basic adaptor domain] [1.3e-17]	YP_001427189 [hypothetical protein ATCV1_Z708L (Acanthocystis turfacea Chlorella virus 1)] [1e- 129]	N/A	Protein kinase	0	1	0
a283L	1E+05	1E+05	52	297	98	11412	12.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A284L	1E+05	1E+05	82	840	279	30766	9.2	Early- Late	Yes	COG3049 [Penicillin V acylase and related amidases] [1e-06]	PF02275.11 [Linear amide C- N hydrolases choloylglycine hydrolase family] [1e-49]	YP_001425725 [hypothetical protein FR483_N093R (Paramecium bursaria Chlorella virus FR483)] [4e-91]	P54948 [RecName: FullUncharacterized protein yxel] [1e-29]	Amidase	0	0	0
a285R	1E+05	1E+05	64	216	71	7946	12.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A286R	1E+05	1E+05	30	1137	378	43042	9.6	Late	Yes	N/A	N/A	YP_001425721 [hypothetical protein FR483_N089L (Paramecium bursaria Chlorella virus FR483)] [2e-92]	N/A	Unknown protein	0	0	0
a286aR	1E+05	1E+05	76	174	57	6925	9.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
a286bL	1E+05	1E+05	52	174	57	6719	9.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
A287R	1E+05	1E+05	74	807	268	31349	9.4	Early- Late	Yes	N/A	PF01541.17 [GIY-YIG catalytic domain] [4.2e-11] / PF07453.6 [NUMOD1 domain] [8.6e-11]	YP_001426191 [hypothetical protein FR483_N559R (Paramecium bursaria Chlorella virus FR483)] [6e-50]	N/A	Unknown protein	0	0	0
a288L	1E+05	1E+05	62	267	88	9946	9.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A289L	1E+05	1E+05	74	852	283	32038	7.3	Early- Late	N/A	COG0515 [Serine/threonine protein kinase] [9e-06]	PF00069.18 [Protein kinase domain] [5.5e-50]	YP_001426180 [hypothetical protein FR483_N548R (Paramecium bursaria	Q54DF2 [RecName: FullProbable serine/threonine- protein kinase MARK-	Protein kinase	0	0	0

												Chlorella virus FR483]] [2e-31]	A] [1e-23]				
a290R	1E+05	1E+05	48	408	135	15384	7.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	3	2
a291R	1E+05	1E+05	68	270	89	10691	8.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	1
A292L	1E+05	2E+05	80	987	328	36771	9.3	Late	N/A	N/A	PF01374.11 [Glycosyl hydrolase family 46] [6.1e-97]	YP_001426685 [hypothetical protein ATCV1_Z204R (Acanthocystis turfacea Chlorella virus 1)] [1e- 114]	O07921 [RecName: FullChitosanase Flags: Precursor] [9e- 17]	Chitosanase	0	0	0
a293R	1E+05	1E+05	56	507	168	18889	11.7	N/A	N/A	N/A	N/A	YP_001426689 [hypothetical protein ATCV1_z208L (Acanthocystis turfacea Chlorella virus 1)] [4e- 14]	N/A	Hypothetical protein	1	1	1
a294R	1E+05	2E+05	54	225	74	7580	8.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A295L	2E+05	2E+05	84	954	317	35626	7.9	Early- Late	Yes	COG1088 [dTDP- D-glucose 4e- dehydratase] [8e- 06]	PF01370.14 [NAD dependent epimerase/dehyd ratase family] [1.6e-68]	YP_001426763 [hypothetical protein ATCV1_Z282L (Acanthocystis turfacea Chlorella virus 1)] [1e- 119]	Q9LMU0 [RecName: FullPutative GDP-L- fucose synthase 2 AltName: FullGDP-4- keto-6-deoxy-D- mannose-3 5- epimerase-4- reductase 2 ShortAtGER2] [1e- 100]	Fucose synthetase	0	0	0
A296R	2E+05	2E+05	14	474	157	17393	12.2	Late	Yes	N/A	N/A	YP_001426690 [hypothetical protein ATCV1_Z209R (Acanthocystis turfacea Chlorella virus 1)] [3e- 16]	N/A	Unknown protein	0	1	1
A297L	2E+05	2E+05	88	534	177	20931	9.5	Late	N/A	N/A	N/A	YP_001425724 [hypothetical protein FR483_N092L (Paramecium bursaria Chlorella virus FR483)] [5e-47]	N/A	Unknown protein	0	0	0
a297aL	2E+05	2E+05	40	129	42	5180	11.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A298L	2E+05	2E+05	72	678	225	25489	5.1	Early	N/A	N/A	N/A	YP_001425716 [hypothetical protein FR483_N084R (Paramecium bursaria Chlorella virus FR483)] [4e-36]	N/A	Unknown protein	0	0	0
a299R	2E+05	2E+05	48	519	172	18677	11.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	3	2
a300R	2E+05	2E+05	58	231	76	9009	9.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A301L	2E+05	2E+05	80	726	241	27394	4.5	Early	N/A	N/A	N/A	YP_001425714 [hypothetical protein FR483_N082R (Paramecium bursaria Chlorella virus FR483)] [8e-11]	N/A	Unknown protein	0	0	0
a302R	2E+05	2E+05	50	423	140	16434	7.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	1

a303L	2E+05	2E+05	68	339	112	13416	11.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
A304R	2E+05	2E+05	8	237	78	9490	5.8	Late	Yes	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A305L	2E+05	2E+05	66	615	204	22910	10.7	Late	Yes	N/A	PF00782.13 [Dual specificity phosphatase catalytic domain] [3.6e-33]	YP_001425950 [hypothetical protein FR483_N318L (Paramecium bursaria Chlorella virus FR483)] [9e-64]	O6BY84 [RecName: FullDual specificity protein phosphatase 16 AllName: FullMitogen-activated protein kinase phosphatase 7 ShortMAP kinase phosphatase 7 ShortMKP-7] [7e-12]	Protein phosphatase	0	0	0
A306L	2E+05	2E+05	82	261	86	10359	4.0	Early-Late	N/A	N/A	N/A	YP_001426702 [hypothetical protein ATCV1_Z221L (Acanthocystis turfacea Chlorella virus 1)] [9e-23]	N/A	Unknown protein	0	0	0
a307R	2E+05	2E+05	30	237	78	9139	9.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
a307aR	2E+05	2E+05	68	243	80	9482	9.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
A308L	2E+05	2E+05	72	360	119	13562	4.0	Early	N/A	N/A	N/A	YP_001425955 [hypothetical protein FR483_N323L (Paramecium bursaria Chlorella virus FR483)] [6e-10]	N/A	Unknown protein	0	0	0
a309R	2E+05	2E+05	64	129	42	4922	7.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
A310L	2E+05	2E+05	88	513	170	18268	8.5	Late	Yes	N/A	N/A	YP_001426703 [hypothetical protein ATCV1_Z222L (Acanthocystis turfacea Chlorella virus 1)] [3e-52]	N/A	Unknown protein	0	0	0
a311R	2E+05	2E+05	54	252	83	8714	6.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A312L	2E+05	2E+05	66	717	238	26612	5.0	Early-Late	N/A	N/A	N/A	YP_001425959 [hypothetical protein FR483_N327L (Paramecium bursaria Chlorella virus FR483)] [4e-94]	N/A	Unknown protein	0	0	0
a312aR	2E+05	2E+05	42	159	52	6074	4.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
A313L	2E+05	2E+05	78	216	71	7838	9.8	Late	N/A	N/A	N/A	YP_001425961 [hypothetical protein FR483_N329L (Paramecium bursaria Chlorella virus FR483)] [9e-06]	N/A	Unknown protein	1	1	1
a313aR	2E+05	2E+05	64	162	53	6275	9.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A314R	2E+05	2E+05	68	243	80	9114	6.7	Late	Yes	N/A	N/A	YP_001425962 [hypothetical protein FR483_N330R (Paramecium bursaria Chlorella virus FR483)]	N/A	Unknown protein	1	1	1

A328L	2E+05	2E+05	88	1068	355	42024	9.2	N/A	N/A	N/A	N/A	YP_001425978 [hypothetical protein FR483_N346L (Paramecium bursaria Chlorella virus FR483)] [3e-69]	N/A	Hypothetical protein	0	0	0
A329R	2E+05	2E+05	64	291	96	10756	4.8	Late	N/A	N/A	N/A	YP_001426735 [hypothetical protein ATCV1_Z254R (Acanthocystis turfacea Chlorella virus 1)] [3e- 13]	N/A	Unknown protein	0	0	0
a329aL	2E+05	2E+05	36	135	44	4997	8.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A329bR	2E+05	2E+05	80	186	61	6905	10.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	1	1
a329cR	2E+05	2E+05	74	186	61	6892	9.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	1
A330R	2E+05	2E+05	84	1299	432	49958	7.5	Early- Late	N/A	COG0666 [FOG: Ankyrin repeat] [9e-06]	PF00023.23 [Ankyrin repeat] [1.3e-07]	XP_001321901 [ankyrin repeat protein (Trichomonas vaginalis G3)] [7e-31]	Q12955 [RecName: FullAnkyrin-3 ShortANK-3 AllName: FullAnkyrin-G] [1e-26]	Unknown protein	0	0	0
a331L	2E+05	2E+05	62	204	67	7616	9.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	1
A333L	2E+05	2E+05	92	1173	390	43025	8.1	Early- Late	N/A	N/A	PF03067.8 [Chitin binding domain] [2.1e-24] / PF01607.17 [Chitin binding Peritrophin-A domain] [1.7e-07]	YP_001426322 [hypothetical protein FR483_N690R (Paramecium bursaria Chlorella virus FR483)] [2e-85]	N/A	Unknown protein	1	2	1
a334L	2E+05	2E+05	54	270	89	10302	12.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a335R	2E+05	2E+05	58	225	74	7939	10.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a336R	2E+05	2E+05	60	258	85	9750	8.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A337L	2E+05	2E+05	76	609	202	23000	8.8	Early- Late	N/A	N/A	N/A	YP_001426717 [hypothetical protein ATCV1_Z236L (Acanthocystis turfacea Chlorella virus 1)] [5e- 16]	N/A	Unknown protein	0	0	0
a340R	2E+05	2E+05	62	279	92	11149	8.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A339L	2E+05	2E+05	76	186	61	7372	11.1	Early- Late	Yes	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A341L	2E+05	2E+05	76	408	135	15341	4.0	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
a341aR	2E+05	2E+05	52	198	65	7269	4.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a341bR	2E+05	2E+05	8	159	52	5765	8.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A342L	2E+05	2E+05	74	1731	576	63813	9.2	Early- Late	Yes	N/A	N/A	YP_001425991 [hypothetical protein FR483_N359L (Paramecium bursaria Chlorella virus FR483)] [0.0]	N/A	Unknown protein	1	1	1

a343R	2E+05	2E+05	56	261	86	10111	8.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a344R	2E+05	2E+05	44	348	115	13071	10.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a345L	2E+05	2E+05	54	291	96	11083	10.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a346L	2E+05	2E+05	48	228	75	8680	9.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	2
a347L	2E+05	2E+05	40	213	70	8435	7.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A348R	2E+05	2E+05	66	480	159	18321	8.9	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
a348aR	2E+05	2E+05	52	156	51	5768	9.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
A349L	2E+05	2E+05	66	549	182	21077	10.0	Early-Late	Yes	N/A	N/A	YP_001426595 [hypothetical protein ATCV1_Z114L (Acanthocystis turfacea Chlorella virus 1)] [2e-31]	N/A	Unknown protein	0	1	0
A350R	2E+05	2E+05	64	369	122	14676	9.7	N/A	Yes	N/A	PF12239.1 [Protein of unknown function (DUF3605)] [4.4e-23]	YP_001426000 [hypothetical protein FR483_n368R (Paramecium bursaria Chlorella virus FR483)] [6e-30]	N/A	Unknown protein	0	0	0
a350aR	2E+05	2E+05	52	135	44	5299	11.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A351L	2E+05	2E+05	80	1077	358	41335	9.6	Early	N/A	N/A	PF01541.17 [GTY-YIG catalytic domain] [3.6e-08]	YP_001426752 [hypothetical protein ATCV1_Z271R (Acanthocystis turfacea Chlorella virus 1)] [3e-84]	N/A	Unknown protein	0	1	0
a351aL	2E+05	2E+05	36	132	43	5148	8.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A352L	2E+05	2E+05	90	624	207	23310	3.6	Late	Yes	N/A	N/A	YP_001426751 [hypothetical protein ATCV1_Z270R (Acanthocystis turfacea Chlorella virus 1)] [2e-65]	Q5UQF7 [RecName: FullUncharacterized protein R489 Flags: Precursor] [1e-05]	Unknown protein	0	1	1
a353R	2E+05	2E+05	48	207	68	9017	12.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A354R	2E+05	2E+05	88	1038	345	40103	9.6	Early-Late	N/A	N/A	N/A	YP_001427263 [hypothetical protein ATCV1_Z782L (Acanthocystis turfacea Chlorella virus 1)] [4e-72]	N/A	Unknown protein	0	0	0
a355L	2E+05	2E+05	64	219	72	8354	7.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A356R	2E+05	2E+05	68	324	107	12512	10.5	N/A	Yes	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A357L	2E+05	2E+05	82	987	328	36796	4.0	Early-Late	N/A	N/A	N/A	YP_001426005 [hypothetical protein FR483_N373L (Paramecium bursaria Chlorella virus FR483)]	N/A	Unknown protein	0	0	0

a395bL	2E+05	2E+05	64	138	45	5372	9.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A396L	2E+05	2E+05	82	459	152	18281	7.9	Early-Late	N/A	N/A	N/A	YP_001426451 [hypothetical protein FR483_N819R (Paramecium bursaria Chlorella virus FR483)] [3e-13]	N/A	Unknown protein	1	1	1
A397R	2E+05	2E+05	86	459	152	18036	4.4	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A398L	2E+05	2E+05	60	357	118	12987	9.9	Late	Yes	N/A	N/A	YP_001426210 [hypothetical protein FR483_N578R (Paramecium bursaria Chlorella virus FR483)] [1e-35]	N/A	Unknown protein	2	3	3
a398aR	2E+05	2E+05	54	180	59	6906	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A399R	2E+05	2E+05	64	585	194	21832	8.9	Early	N/A	N/A	PF00075.17 [RNase H] [6.7e-19]	YP_001426207 [hypothetical protein FR483_N575L (Paramecium bursaria Chlorella virus FR483)] [9e-37]	N/A	Unknown protein	0	0	0
A400R	2E+05	2E+05	78	357	118	13634	9.5	Early-Late	Yes	N/A	N/A	YP_001426835 [hypothetical protein ATCV1_z354L (Acanthocystis turfacea Chlorella virus 1)] [3e-20]	N/A	Unknown protein	0	0	0
A401R	2E+05	2E+05	78	834	277	31433	6.8	Early-Late	N/A	N/A	N/A	YP_001426204 [hypothetical protein FR483_N572L (Paramecium bursaria Chlorella virus FR483)] [2e-95]	N/A	Unknown protein	0	1	1
A402R	2E+05	2E+05	78	684	227	25380	7.5	Late	N/A	N/A	N/A	YP_001426843 [hypothetical protein ATCV1_Z362R (Acanthocystis turfacea Chlorella virus 1)] [6e-67]	N/A	Unknown protein	0	0	0
A403R	2E+05	2E+05	78	282	93	10830	10.0	Early-Late	N/A	N/A	N/A	YP_001426844 [hypothetical protein ATCV1_Z363R (Acanthocystis turfacea Chlorella virus 1)] [7e-20]	N/A	Unknown protein	0	0	0
A404R	2E+05	2E+05	88	576	191	22111	2.9	Late	N/A	N/A	N/A	YP_001426199 [hypothetical protein FR483_N567L (Paramecium bursaria Chlorella virus FR483)] [2e-13]	N/A	Unknown protein	1	1	1
A404aL	2E+05	2E+05	84	159	52	5910	4.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A405R	2E+05	2E+05	6	1491	496	53502	10.3	Late	Yes	N/A	N/A	YP_001426197 [hypothetical protein FR483_N565L (Paramecium bursaria Chlorella virus FR483)] [1e-42]	N/A	Unknown protein	1	2	1

a406L	2E+05	2E+05	70	201	66	7386	9.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A407L	2E+05	2E+05	80	633	210	23382	8.9	Late	Yes	N/A	N/A	YP_001426955 [hypothetical protein ATCV1_Z474R (Acanthocystis turfacea Chlorella virus 1)] [2e-70]	N/A	Unknown protein	1	2	2
A408L	2E+05	2E+05	56	834	277	30772	4.5	Early-Late	N/A	N/A	N/A	YP_001426126 [hypothetical protein FR483_N494R (Paramecium bursaria Chlorella virus FR483)] [8e-48]	N/A	Unknown protein	0	0	0
a409R	2E+05	2E+05	44	249	82	9006	5.4	N/A	N/A	N/A	N/A	YP_001426951 [hypothetical protein ATCV1_z470L (Acanthocystis turfacea Chlorella virus 1)] [3e-06]	N/A	Hypothetical protein	1	2	1
A410L	2E+05	2E+05	72	333	110	13015	4.4	Early	N/A	N/A	N/A	YP_001426942 [hypothetical protein ATCV1_Z461L (Acanthocystis turfacea Chlorella virus 1)] [1e-26]	N/A	Unknown protein	0	0	0
A411R	2E+05	2E+05	78	513	170	19584	4.6	Late	N/A	N/A	N/A	YP_001426120 [hypothetical protein FR483_N488R (Paramecium bursaria Chlorella virus FR483)] [3e-09]	N/A	Unknown protein	0	0	0
A412R	2E+05	2E+05	74	540	179	20539	7.4	Early	N/A	N/A	N/A	YP_001426947 [hypothetical protein ATCV1_Z466R (Acanthocystis turfacea Chlorella virus 1)] [3e-47]	N/A	Unknown protein	0	0	0
A413L	2E+05	2E+05	80	735	244	26998	9.5	Late	Yes	N/A	N/A	YP_001426125 [hypothetical protein FR483_N493L (Paramecium bursaria Chlorella virus FR483)] [1e-29]	N/A	Unknown protein	2	2	2
A414R	2E+05	2E+05	48	282	93	10612	10.8	Late	Yes	N/A	N/A	YP_001426117 [hypothetical protein FR483_N485R (Paramecium bursaria Chlorella virus FR483)] [5e-11]	N/A	Unknown protein	2	2	2
a415L	2E+05	2E+05	66	210	69	7685	10.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
A416R	2E+05	2E+05	82	567	188	22361	6.5	Late	N/A	COG1428 [Deoxynucleoside kinases] [8e-07]	PF01712.12 [Deoxynucleoside kinase] [1.1e-36]	YP_001426070 [hypothetical protein FR483_N436R (Paramecium bursaria Chlorella virus FR483)] [6e-48]	Q197D1 [RecName: FullPutative kinase protein 029R] [4e-21]	Unknown protein	0	1	0
A417L	2E+05	2E+05	82	1290	429	48932	7.5	Late	N/A	N/A	N/A	YP_001426073 [hypothetical protein FR483_N441L (Paramecium bursaria Chlorella virus FR483)] [4e-52]	A6UWR5 [RecName: FullReplication factor C large subunit ShortRFC large subunit AltName: FullClamp loader large subunit] [1e-06]	Unknown protein	0	0	0

a418R	2E+05	2E+05	62	240	79	9726	10.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
a419R	2E+05	2E+05	74	213	70	7892	5.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A420L	2E+05	2E+05	82	213	70	7918	6.4	Late	Yes	N/A	N/A	YP_001426933 [hypothetical protein ATCV1_Z452R (Acanthocystis turfacea Chlorella virus 1)] [5e-21]	N/A	Unknown protein	1	1	1
A421R	2E+05	2E+05	10	297	98	11056	10.1	Late	Yes	N/A	N/A	YP_001426078 [hypothetical protein FR483_N446R (Paramecium bursaria Chlorella virus FR483)] [2e-16]	N/A	Unknown protein	1	1	1
A422R	2E+05	2E+05	82	993	330	37736	9.6	Early-Late	N/A	N/A	PF07463.4 [NUMOD4 motif] [5.7e-13] / PF01844.16 [HNH endonuclease] [1.4e-06]	YP_001426831 [hypothetical protein ATCV1_Z350L (Acanthocystis turfacea Chlorella virus 1)] [6e-15]	Q5UPT4 [RecName: FullUncharacterized HNH endonuclease L247] [1e-12]	Unknown protein	0	0	0
A422aR	2E+05	2E+05	74	192	63	7880	9.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	0
A423R	2E+05	2E+05	72	474	157	18458	6.5	Late	Yes	N/A	N/A	YP_001426081 [hypothetical protein FR483_N449R (Paramecium bursaria Chlorella virus FR483)] [3e-12]	N/A	Unknown protein	0	1	0
A424R	2E+05	2E+05	86	330	109	12637	11.3	Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
a425L	2E+05	2E+05	58	189	62	7087	4.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
A426R	2E+05	2E+05	52	345	114	13391	5.0	Late	N/A	N/A	N/A	YP_001426084 [hypothetical protein FR483_N452R (Paramecium bursaria Chlorella virus FR483)] [2e-17]	N/A	Unknown protein	0	0	0
A427L	2E+05	2E+05	76	360	119	13765	6.5	Early	N/A	N/A	PF00085.13 [Thioredoxin] [8.6e-12]	YP_001426085 [hypothetical protein FR483_N453L (Paramecium bursaria Chlorella virus FR483)] [5e-14]	N/A	Unknown protein	0	0	0
A428L	2E+05	2E+05	80	438	145	15912	3.8	Early-Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A429L	2E+05	2E+05	70	1431	476	55474	4.7	Early	N/A	N/A	N/A	YP_001426092 [hypothetical protein FR483_N460L (Paramecium bursaria Chlorella virus FR483)] [6e-50]	N/A	Unknown protein	0	0	0
A430L	2E+05	2E+05	76	1314	437	48165	7.5	Late	Yes	N/A	PF04451.5 [Large eukaryotic DNA virus major capsid protein] [2.6e-90]	YP_001426761 [hypothetical protein ATCV1_Z280L (Acanthocystis turfacea Chlorella virus 1)] [0.0]	A7U6E9 [RecName: FullMajor capsid protein ShortMCP] [5e-90]	Major capsid protein	0	0	0
A431L	2E+05	2E+05	64	201	66	7999	10.3	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0

A432R	2E+05	2E+05	14	474	157	17358	10.1	Late	N/A	N/A	N/A	YP_001426109 [hypothetical protein FR483_N477R (Paramecium bursaria Chlorella virus FR483)] [9e-27]	N/A	Unknown protein	0	0	0
a433R	2E+05	2E+05	56	342	113	13111	9.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a433aR	2E+05	2E+05	46	135	44	5413	10.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a434L	2E+05	2E+05	58	171	56	5966	9.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a434aR	2E+05	2E+05	62	189	62	6769	9.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A436L	2E+05	2E+05	60	192	63	6932	13.0	N/A	Yes	N/A	PF08789.3 [PBCV-specific basic adaptor domain] [1.5e-16]	YP_001426679 [hypothetical protein ATCV1_Z198L (Acanthocystis turfacea Chlorella virus 1)] [3e-06]	N/A	Unknown protein	0	0	0
A435R	2E+05	2E+05	30	192	63	6795	11.2	Early	N/A	N/A	N/A	YP_001426611 [hypothetical protein ATCV1_z130R (Acanthocystis turfacea Chlorella virus 1)] [1e-05]	N/A	Unknown protein	0	1	0
A437L	2E+05	2E+05	76	312	103	10876	11.0	Late	Yes	N/A	PF05854.4 [Non-histone chromosomal protein MC1] [5.9e-07]	YP_001426116 [hypothetical protein FR483_N484L (Paramecium bursaria Chlorella virus FR483)] [3e-30]	N/A	Unknown protein	0	1	0
A437aR	2E+05	2E+05	40	153	50	5255	10.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A438L	2E+05	2E+05	86	237	78	8988	10.7	Early-Late	Yes	COG0695 [Glutaredoxin and related proteins] [3e-06]	PF00462.17 [Glutaredoxin] [3e-10]	YP_001426068 [hypothetical protein FR483_N436R (Paramecium bursaria Chlorella virus FR483)] [8e-18]	Q1RHJ0 [RecName: Full Glutaredoxin-1] [3e-06]	Glutaredoxin	0	0	0
A439R	2E+05	2E+05	34	339	112	12855	8.3	N/A	N/A	N/A	N/A	YP_001426616 [hypothetical protein ATCV1_Z135R (Acanthocystis turfacea Chlorella virus 1)] [8e-28]	N/A	Hypothetical protein	3	3	2
A439aR	2E+05	2E+05	70	171	56	6279	6.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A440L	2E+05	2E+05	70	267	88	10112	11.1	Early	Yes	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A441L	2E+05	2E+05	72	414	137	15531	4.4	Early	N/A	N/A	N/A	YP_001426062 [hypothetical protein FR483_N430L (Paramecium bursaria Chlorella virus FR483)] [1e-27]	N/A	Unknown protein	0	0	0
a442R	2E+05	2E+05	10	396	131	15348	8.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A443R	2E+05	2E+05	74	927	308	34961	5.3	Early	Yes	N/A	N/A	YP_001426065 [hypothetical protein FR483_N433R]	N/A	Unknown protein	0	0	0

												(Paramecium bursaria Chlorella virus FR483)] [2e-50]						
A444L	2E+05	2E+05	84	315	104	11654	5.2	Late	N/A	N/A	N/A	YP_001426058 [hypothetical protein FR483_N426R (Paramecium bursaria Chlorella virus FR483)] [2e-06]	N/A	Unknown protein	0	0	0	
A445L	2E+05	2E+05	86	1389	462	52542	8.5	Early	N/A	COG0661 [Predicted unusual protein kinase] [5e-06]	PF03109.9 [ABC1 family] [1.3e-29]	YP_001426053 [hypothetical protein FR483_N421R (Paramecium bursaria Chlorella virus FR483)] [1e-168]	Q55884 [RecName: FullUncharacterized protein sll0095] [1e- 21]	Unknown protein	0	0	0	
a446R	2E+05	2E+05	44	306	101	11199	9.4	N/A	N/A	N/A	N/A	YP_001426569 [hypothetical protein ATCV1_z088L (Acanthocystis turfacea Chlorella virus 1)] [5e- 12]	N/A	Hypothetical protein	1	2	2	
a447R	2E+05	2E+05	46	300	99	10962	12.1	N/A	N/A	N/A	N/A	YP_001426568 [hypothetical protein ATCV1_z087L (Acanthocystis turfacea Chlorella virus 1)] [4e- 11]	N/A	Hypothetical protein	0	2	0	
A447aR	2E+05	2E+05	8	138	45	5187	4.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0	
A448L	2E+05	2E+05	76	321	106	12369	10.4	Late	Yes	N/A	PF00085.13 [Thioredoxin] [6.7e-16]	YP_001426562 [hypothetical protein ATCV1_Z081L (Acanthocystis turfacea Chlorella virus 1)] [1e- 31]	Q9XI01 [RecName: FullProbable protein disulfide-isomerase 1 ShortPDI 1 Flags: Precursor] [1e-07]	Protein disulphide isomerase with heme binding site	0	0	0	
a448aL	2E+05	2E+05	40	144	47	5352	9.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
A449R	2E+05	2E+05	70	582	193	22451	8.9	Early	N/A	N/A	PF02940.8 [mRNA capping enzyme beta chain] [3.1e-18]	YP_001426048 [hypothetical protein FR483_N416L (Paramecium bursaria Chlorella virus FR483)] [5e-47]	N/A	Unknown protein	0	0	0	
A450R	2E+05	2E+05	86	750	249	28143	9.4	Early	N/A	N/A	PF07150.4 [Protein of unknown function (DUF1390)] [2.5e-82]	YP_001426424 [hypothetical protein FR483_N792R (Paramecium bursaria Chlorella virus FR483)] [2e-29]	N/A	Unknown protein	0	0	0	
a451L	2E+05	2E+05	54	306	101	12174	10.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	2	0	
A452L	2E+05	2E+05	82	240	79	9447	4.2	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0	
a453R	2E+05	2E+05	10	270	89	10218	7.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
A454L	2E+05	2E+05	82	870	289	31194	4.7	Early- Late	Yes	N/A	N/A	YP_001426311 [hypothetical protein FR483_N679R (Paramecium bursaria Chlorella virus FR483)] [1e-102]	N/A	Unknown protein	1	1	0	
a455R	2E+05	2E+05	42	444	147	15433	7.9	N/A	N/A	N/A	N/A	YP_001426312 [hypothetical protein	N/A	Hypothetical protein	2	2	2	

												FR483_n680L (Paramecium bursaria Chlorella virus FR483) [5e-12]						
A456L	2E+05	2E+05	74	1965	654	75235	5.5	Early	Yes	COG3378 [Predicted ATPase] [3e-06]	PF08706.4 [D5 N terminal like] [3.9e-09]	YP_001426306 [hypothetical protein FR483_N674R (Paramecium bursaria Chlorella virus FR483) [0.0]	N/A	Unknown protein	0	0	0	
a457R	2E+05	2E+05	44	231	76	8771	12.0	N/A	N/A	N/A	N/A	YP_001426309 [hypothetical protein FR483_n677L (Paramecium bursaria Chlorella virus FR483) [2e-06]	N/A	Hypothetical protein	0	0	0	
a458L	2E+05	2E+05	60	252	83	9651	12.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
a459R	2E+05	2E+05	52	303	100	11291	10.5	N/A	N/A	N/A	N/A	YP_001426309 [hypothetical protein FR483_n677L (Paramecium bursaria Chlorella virus FR483) [2e-22]	N/A	Hypothetical protein	0	0	0	
a460R	2E+05	2E+05	56	237	78	9235	11.2	N/A	N/A	N/A	N/A	YP_001426548 [hypothetical protein ATCV1_z067L (Acanthocystis turfacea Chlorella virus 1)] [5e- 11]	N/A	Hypothetical protein	0	0	0	
A461R	2E+05	2E+05	54	234	77	8987	10.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
A462R	2E+05	2E+05	36	216	71	8338	9.7	Early- Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0	
a463L	2E+05	2E+05	54	576	191	22590	10.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
A464R	2E+05	2E+05	50	828	275	31170	6.7	Early	N/A	COG0571 [dsRNA-specific ribonuclease] [1e- 07]	PF00636.19 [RNase3 domain] [8.7e-21] / PF00035.18 [Double-stranded RNA binding motif] [1.2e-11]	YP_001426302 [hypothetical protein FR483_N670L (Paramecium bursaria Chlorella virus FR483) [9e-98]	Q8KAN7 [RecName: FullRibonuclease 3 AltName: FullRibonuclease III ShortRNase III] [5e- 32]	Rnase III	0	0	0	
A464aR	2E+05	2E+05	58	126	41	5049	11.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
A465R	2E+05	2E+05	74	357	118	13528	10.2	Early- Late	Yes	COG5054 [Mitochondrial sulfhydryl oxidase involved in the biogenesis of cytosolic Fe/S proteins] [4e-06]	PF04777.6 [Erv1 / Alr family] [3.5e- 22]	YP_001426300 [hypothetical protein FR483_N668R (Paramecium bursaria Chlorella virus FR483) [8e-42]	Q5UQV6 [RecName: FullProbable FAD- linked sulfhydryl oxidase R368] [3e- 12]	Unknown protein	0	0	0	
a466L	2E+05	2E+05	68	276	91	10518	8.9	N/A	N/A	N/A	N/A	YP_001426543 [hypothetical protein ATCV1_z062L (Acanthocystis turfacea Chlorella virus 1)] [5e- 15]	N/A	Hypothetical protein	0	0	0	
A467L	2E+05	2E+05	78	939	312	36694	7.5	Early- Late	N/A	N/A	N/A	YP_001426297 [hypothetical protein FR483_N665R (Paramecium bursaria Chlorella virus FR483)]	N/A	Unknown protein	0	0	0	

A505L	2E+05	2E+05	64	1455	484	54765	5.4	Early	N/A	N/A	N/A	YP_001426328 [hypothetical protein FR483_N696R (Paramecium bursaria Chlorella virus FR483)] [1e-127]	N/A	Unknown protein	0	0	0
a506R	2E+05	2E+05	50	213	70	7759	10.4	N/A	N/A	N/A	N/A	YP_001426333 [hypothetical protein FR483_n701L (Paramecium bursaria Chlorella virus FR483)] [9e-14]	N/A	Hypothetical protein	0	0	0
a507R	2E+05	2E+05	48	561	186	21950	10.5	N/A	N/A	N/A	N/A	YP_001426327 [hypothetical protein FR483_n695L (Paramecium bursaria Chlorella virus FR483)] [2e-10]	N/A	Hypothetical protein	0	0	0
a508R	2E+05	2E+05	54	234	77	8501	11.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
a509R	2E+05	2E+05	52	306	101	12606	10.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a510R	2E+05	2E+05	50	204	67	7609	10.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a511L	2E+05	2E+05	50	222	73	8589	9.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A512R	2E+05	2E+05	56	2415	804	89893	5.5	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
a513R	2E+05	2E+05	56	363	120	13489	12.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	3	3
a514L	2E+05	2E+05	62	435	144	15673	12.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	2	1
a515L	2E+05	2E+05	44	612	203	22883	10.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a516R	2E+05	2E+05	44	207	68	7990	12.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
A517L	2E+05	2E+05	86	1035	344	39045	8.8	Early	N/A	COG0270 [Site-specific DNA methylase] [9e-06]	PF00145.10 [C-5 cytosine-specific DNA methylase] [5.4e-40]	YP_001426001 [hypothetical protein FR483_N369L (Paramecium bursaria Chlorella virus FR483)] [1e-126]	P36216 [RecName: FullModification methylase CvJI ShortM.CvJI AltName: FullCytosine-specific methyltransferase CvJII] [9e-76]	Unknown protein	0	1	0
a518R	2E+05	2E+05	62	303	100	10825	9.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A519L	2E+05	2E+05	74	249	82	9332	9.8	Late	N/A	N/A	N/A	YP_001425920 [hypothetical protein FR483_N288R (Paramecium bursaria Chlorella virus FR483)] [9e-16]	N/A	Unknown protein	2	2	2
A520L	2E+05	2E+05	74	303	100	11674	10.7	Late	Yes	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A521L	2E+05	2E+05	70	630	209	23738	11.4	Early-Late	Yes	N/A	N/A	YP_001426325 [hypothetical protein FR483_N693R (Paramecium bursaria Chlorella virus FR483)]	N/A	Unknown protein	0	0	0

A533R	3E+05	3E+05	82	1125	374	40132	3.8	Early-Late	Yes	N/A	N/A	YP_001426965 [hypothetical protein ATCV1_z484R (Acanthocystis turfacea Chlorella virus 1)] [1e-117]	N/A	Unknown protein	0	0	0
A534R	3E+05	3E+05	56	318	105	11783	9.7	N/A	Yes	N/A	N/A	YP_001426148 [hypothetical protein FR483_N516R (Paramecium bursaria Chlorella virus FR483)] [3e-37]	N/A	Unknown protein	0	0	0
A535L	3E+05	3E+05	82	216	71	8210	4.7	Early-Late	Yes	N/A	N/A	YP_001426150 [hypothetical protein FR483_N518L (Paramecium bursaria Chlorella virus FR483)] [7e-11]	N/A	Unknown protein	0	0	0
A536L	3E+05	3E+05	80	222	73	8485	10.0	Early-Late	Yes	N/A	N/A	YP_001426152 [hypothetical protein FR483_N520L (Paramecium bursaria Chlorella virus FR483)] [6e-09]	N/A	Unknown protein	1	1	0
a536aL	3E+05	3E+05	48	177	58	6753	8.5	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	0	
A537L	3E+05	3E+05	78	798	265	31237	6.6	Early	N/A	N/A	N/A	YP_001426153 [hypothetical protein FR483_N521L (Paramecium bursaria Chlorella virus FR483)] [1e-18]	N/A	Unknown protein	0	0	0
a538L	3E+05	3E+05	68	198	65	7367	9.9	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0	
A539R	3E+05	3E+05	80	522	173	19839	10.1	Early	N/A	N/A	PF01541.17 [GIY-YIG catalytic domain] [1.1e-06]	YP_001426156 [hypothetical protein FR483_N524R (Paramecium bursaria Chlorella virus FR483)] [2e-46]	N/A	Unknown protein	0	0	0
a539aR	3E+05	3E+05	64	165	54	6339	9.3	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0	
A540L	3E+05	3E+05	82	3771	1256	127197	6.2	Late	Yes	N/A	N/A	YP_001426157 [hypothetical protein FR483_N525L (Paramecium bursaria Chlorella virus FR483)] [1e-173]	N/A	Unknown protein	0	0	0
a541R	3E+05	3E+05	52	291	96	10536	4.0	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0	
a542R	3E+05	3E+05	48	219	72	9007	9.1	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0	
A543L	3E+05	3E+05	58	168	55	5802	6.5	Late	N/A	N/A	N/A	N/A	Unknown protein	0	2	1	
A544R	3E+05	3E+05	52	897	298	34250	8.2	Late	N/A	COG1793 [ATP-dependent DNA ligase] [5e-11]	PF01068.14 [ATP-dependent DNA ligase domain] [1.3e-22]	YP_001426668 [hypothetical protein ATCV1_z187L (Acanthocystis turfacea Chlorella virus 1)] [1e-86]	P44121 [RecName: FullDNA ligase AltName: FullPolydeoxyribonucleotide synthase (ATP)] [5e-10]	ATP-dependent DNA ligase	0	0	0
a545L	3E+05	3E+05	72	222	73	8111	10.0	N/A	N/A	N/A	N/A	YP_001426667 [hypothetical protein ATCV1_z186R	N/A	Hypothetical protein	0	1	0

A558L	3E+05	3E+05	82	1203	400	45547	5.1	Early-Late	Yes	N/A	PF04451.5 [Large eukaryotic DNA virus major capsid protein] [6.6e-60]	YP_001426987 [hypothetical protein ATCV1_Z506L (Acanthocystis turfacea Chlorella virus 1)] [1e-156]	A7U6E9 [RecName: FullMajor capsid protein ShortMCP] [3e-47]	Capsid protein	0	0	0
a558aL	3E+05	3E+05	36	141	46	5526	5.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A559L	3E+05	3E+05	78	642	213	24034	10.2	Late	Yes	N/A	N/A	YP_001425898 [hypothetical protein FR483_N266R (Paramecium bursaria Chlorella virus FR483)] [1e-16]	N/A	Unknown protein	1	1	0
a560R	3E+05	3E+05	68	315	104	11524	8.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
A561L	3E+05	3E+05	90	1950	649	71004	9.9	Late	Yes	N/A	N/A	YP_001426992 [hypothetical protein ATCV1_Z511L (Acanthocystis turfacea Chlorella virus 1)] [4e-60]	N/A	Unknown protein	1	2	1
a562R	3E+05	3E+05	42	198	65	7690	11.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a563R	3E+05	3E+05	68	180	59	6418	10.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A564L	3E+05	3E+05	70	1056	351	39884	8.4	Early-Late	N/A	N/A	N/A	YP_001425778 [hypothetical protein FR483_N146R (Paramecium bursaria Chlorella virus FR483)] [5e-66]	N/A	Unknown protein	0	0	0
A565R	3E+05	3E+05	70	2013	670	73169	7.3	Early-Late	Yes	N/A	N/A	YP_001425891 [hypothetical protein FR483_N259L (Paramecium bursaria Chlorella virus FR483)] [2e-69]	N/A	Unknown protein	1	1	1
a566L	3E+05	3E+05	60	273	90	10337	7.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A567L	3E+05	3E+05	76	459	152	17418	10.1	Early-Late	Yes	N/A	N/A	YP_001425887 [hypothetical protein FR483_N255R (Paramecium bursaria Chlorella virus FR483)] [1e-12]	N/A	Unknown protein	0	0	0
A568L	3E+05	3E+05	72	540	179	20899	7.5	Early	N/A	N/A	N/A	YP_001426314 [hypothetical protein FR483_N682L (Paramecium bursaria Chlorella virus FR483)] [3e-10]	N/A	Unknown protein	1	1	2
a569R	3E+05	3E+05	66	255	84	9639	9.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	1
A570L	3E+05	3E+05	86	387	128	14556	4.8	Late	N/A	N/A	N/A	YP_001426316 [hypothetical protein FR483_N684L (Paramecium bursaria Chlorella virus FR483)] [9e-38]	N/A	Unknown protein	0	0	0
A571R	3E+05	3E+05	14	351	116	12972	12.0	Late	Yes	N/A	PF08789.3 [PBCV-specific basic adaptor]	YP_001426112 [hypothetical protein FR483_N480R]	N/A	Unknown protein	0	0	0

										topoisomerase (DNA gyrase/topo II topoisomerase IV) B subunit [2e-08]	[1.4e-22] / PF00521.13 [DNA gyrase/topoisomerase IV subunit A] [6.5e-124]	FR483_N549R (Paramecium bursaria Chlorella virus FR483) [0.0]	topoisomerase 2 AltName: FullDNA topoisomerase II [0.0]				
a584R	3E+05	3E+05	56	297	98	11326	11.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a585R	3E+05	3E+05	62	240	79	9304	9.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A586R	3E+05	3E+05	62	228	75	8567	11.8	N/A	Yes	N/A	N/A	YP_001426187 [hypothetical protein FR483_n555L (Paramecium bursaria Chlorella virus FR483)] [1e-11]	N/A	Unknown protein	0	0	0
a587R	3E+05	3E+05	60	345	114	12877	8.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a588R	3E+05	3E+05	42	351	116	13729	11.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a589L	3E+05	3E+05	56	231	76	8638	12.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A589aL	3E+05	3E+05	80	153	50	5921	8.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A590L	3E+05	3E+05	84	1047	348	42036	4.6	Early-Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A590aL	3E+05	3E+05	44	126	41	4735	12.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
a591L	3E+05	3E+05	68	168	55	6171	10.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A592R	3E+05	3E+05	82	210	69	7861	4.7	Early-Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A593R	3E+05	3E+05	74	900	299	32019	7.1	N/A	N/A	N/A	N/A	YP_001426230 [hypothetical protein FR483_N598R (Paramecium bursaria Chlorella virus FR483)] [3e-35]	N/A	Hypothetical protein	0	0	0
a594R	3E+05	3E+05	42	351	116	14065	12.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a595L	3E+05	3E+05	56	255	84	9195	7.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
A596R	3E+05	3E+05	78	429	142	15892	6.5	Early-Late	N/A	COG2131 [Deoxycytidylate deaminase] [9e-07]	PF00383.15 [Cytidine and deoxycytidylate deaminase zinc-binding region] [1.1e-26]	YP_001427097 [hypothetical protein ATCV1_2616L (Acanthocystis turfacea Chlorella virus 1)] [2e-56]	P33968 [RecName: FullUncharacterized deaminase in luxG 3region] [1e-19]	Unknown protein	0	0	0
a597L	3E+05	3E+05	54	297	98	10825	6.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A598L	3E+05	3E+05	84	1092	363	41558	6.9	Early-Late	Yes	COG0076 [Glutamate decarboxylase and related PLP-dependent proteins] [5e-06]	PF00282.12 [Pyridoxal-dependent decarboxylase conserved domain] [1.1e-17]	YP_001426232 [hypothetical protein FR483_N600L (Paramecium bursaria Chlorella virus FR483)] [1e-132]	P54772 [RecName: FullHistidine decarboxylase ShortHDC AltName: FullTOM92] [3e-53]	Unknown protein	0	0	0

a599R	3E+05	3E+05	62	540	179	21558	8.7	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a600R	3E+05	3E+05	52	249	82	9151	11.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
a600aR	3E+05	3E+05	58	141	46	5877	8.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A601R	3E+05	3E+05	52	306	101	11592	10.6	Early-Late	N/A	N/A	N/A	YP_001426767 [hypothetical protein ATCV1_z286L (Acanthocystis turfacea Chlorella virus 1)] [5e-19]	N/A	Unknown protein	0	2	2
A602L	3E+05	3E+05	60	606	201	23569	5.1	Early	N/A	N/A	N/A	YP_001426587 [hypothetical protein ATCV1_z106R (Acanthocystis turfacea Chlorella virus 1)] [3e-14]	N/A	Unknown protein	0	0	0
a603R	3E+05	3E+05	78	318	105	12141	4.6	N/A	N/A	N/A	N/A	YP_001426236 [hypothetical protein FR483_N604R (Paramecium bursaria Chlorella virus FR483)] [6e-20]	N/A	Hypothetical protein	0	0	0
A603aL	3E+05	3E+05	72	186	61	6697	8.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
a603bR	3E+05	3E+05	72	147	48	5317	4.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	0
A604L	3E+05	3E+05	80	405	134	15162	9.9	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	3	0
A605L	3E+05	3E+05	80	477	158	17769	10.9	Early-Late	Yes	N/A	N/A	N/A	N/A	Unknown protein	1	1	1
a606L	3E+05	3E+05	50	345	114	13257	10.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
A607R	3E+05	3E+05	12	1176	391	46171	7.2	Early-Late	N/A	N/A	N/A	YP_001426342 [hypothetical protein FR483_N710R (Paramecium bursaria Chlorella virus FR483)] [9e-77]	Q02357 [RecName: FullAnkyrin-1 ShortANK-1 AltName: FullErythrocyte ankyrin] [3e-06]	Unknown protein	0	0	0
A609L	3E+05	3E+05	82	1170	389	43459	4.7	Early	N/A	COG0677 [UDP-N-acetyl-D-mannosaminuronate dehydrogenase] [2e-06]	PF03721.7 [UDP-glucose/GDP-mannose dehydrogenase family NAD binding domain] [8.7e-41] / PF00984.12 [UDP-glucose/GDP-mannose dehydrogenase family central domain] [5.8e-23] / PF03720.8 [UDP-glucose/GDP-mannose dehydrogenase family UDP binding domain] [5e-08]	YP_001426344 [hypothetical protein FR483_N712L (Paramecium bursaria Chlorella virus FR483)] [1e-133]	O33952 [RecName: FullUDP-glucose 6-dehydrogenase ShortUDP-Glc dehydrogenase ShortUDP-GlcDH ShortUDPGDH] [1e-119]	UDP-glucose dehydrogenase	0	0	0

a610R	3E+05	3E+05	66	276	91	10725	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a611R	3E+05	3E+05	62	225	74	8698	8.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A612L	3E+05	3E+05	78	360	119	13587	8.7	Late	Yes	COG2940 [Proteins containing SET domain] [1e-06]	PF00856.21 [SET domain] [1.2e-12]	YP_001426351 [hypothetical protein FR483_N719L (Paramecium bursaria Chlorella virus FR483)] [3e-33]	Q0U3A4 [RecName: FullHistone-lysine N-methyltransferase SET9 AllName: FullSET domain protein 9] [4e-07]	Histone H3K27 methylase	0	0	0
a613R	3E+05	3E+05	54	264	87	10383	10.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A614L	3E+05	3E+05	86	1734	577	64733	11.2	Late	Yes	N/A	PF00069.18 [Protein kinase domain] [5.6e-11]	YP_001426352 [hypothetical protein FR483_N720L (Paramecium bursaria Chlorella virus FR483)] [1e-100]	N/A	Protein kinase	0	0	0
a615R	3E+05	3E+05	66	201	66	7112	9.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a616R	3E+05	3E+05	60	282	93	9978	7.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A617R	3E+05	3E+05	34	966	321	37586	9.9	Early-Late	Yes	N/A	N/A	YP_001426356 [hypothetical protein FR483_N724R (Paramecium bursaria Chlorella virus FR483)] [3e-82]	Q5UQJ6 [RecName: FullPutative serine/threonine-protein kinase R400] [7e-12]	Unknown protein	0	0	0
A618L	3E+05	3E+05	70	396	131	15223	3.8	Late	N/A	N/A	N/A	YP_001426357 [hypothetical protein FR483_N725L (Paramecium bursaria Chlorella virus FR483)] [3e-14]	N/A	Unknown protein	0	0	0
A619L	3E+05	3E+05	84	714	237	26943	4.5	Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A620L	3E+05	3E+05	86	252	83	9739	9.6	Late	N/A	N/A	N/A	YP_001426363 [hypothetical protein FR483_N731L (Paramecium bursaria Chlorella virus FR483)] [1e-16]	N/A	Unknown protein	2	2	2
a620aR	3E+05	3E+05	56	174	57	6932	9.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A621L	3E+05	3E+05	84	354	117	12935	9.5	Late	Yes	N/A	N/A	YP_001427169 [hypothetical protein ATCV1_z688R (Acanthocystis turfacea Chlorella virus 1)] [4e-35]	N/A	Unknown protein	2	2	2
a621aR	3E+05	3E+05	76	168	55	6081	4.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
a621bL	3E+05	3E+05	56	141	46	5224	10.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A622L	3E+05	3E+05	80	1563	520	58097	5.7	Late	Yes	N/A	PF04451.5 [Large eukaryotic DNA virus major capsid protein] [1.7e-66]	YP_001426369 [hypothetical protein FR483_N737L (Paramecium bursaria Chlorella virus FR483)] [0.0]	A7U6E9 [RecName: FullMajor capsid protein ShortMCP] [9e-57]	Capsid protein	0	0	0

A623L	3E+05	3E+05	72	204	67	7703	9.2	Early	N/A	N/A	PF01428.9 [AN1-like Zinc finger] [1.7e-12]	XP_002466323 [hypothetical protein SORBIDRAFT_01g005640 (Sorghum bicolor)] [2e-07]	Q852K6 [RecName: FullZinc finger A20 and AN1 domain-containing stress-associated protein 7 ShortOSAP7] [2e-08]	Unknown protein	0	0	0
A623aL	3E+05	3E+05	56	180	59	6781	4.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A624R	3E+05	3E+05	32	366	121	13570	9.3	Late	Yes	N/A	PF09945.2 [Predicted membrane protein (DUF2177)] [3.4e-26]	YP_001426373 [hypothetical protein FR483_N741R (Paramecium bursaria Chlorella virus FR483)] [2e-21]	N/A	Unknown protein	3	4	3
A625R	3E+05	3E+05	72	1299	432	49945	10.7	Late	Yes	COG0675 [Transposase and inactivated derivatives] [1e-06]	PF12323.1 [Helix-turn-helix domain] [1.4e-06] / PF07282.4 [Putative transposase DNA-binding domain] [6.7e-18]	ZP_03272524 [transposase IS605 OrfB family (Arthrospira maxima CS-326)] [2e-20]	Q5UPI0 [RecName: FullPutative transposase R104] [8e-19]	Unknown protein	0	0	0
a626L	3E+05	3E+05	48	225	74	8395	9.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a626aR	3E+05	3E+05	60	159	52	6068	11.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A627R	3E+05	3E+05	68	1338	445	49629	11.1	Late	Yes	N/A	N/A	YP_001426405 [hypothetical protein FR483_N773L (Paramecium bursaria Chlorella virus FR483)] [1e-135]	N/A	Unknown protein	1	3	0
a627aR	3E+05	3E+05	54	138	45	5526	12.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A628L	3E+05	3E+05	82	294	97	10499	4.5	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
A629R	3E+05	3E+05	74	2316	771	86292	7.5	Early-Late	Yes	COG0209 [Ribonucleotide reductase alpha subunit] [4e-11]	PF03477.9 [ATP cone domain] [8.5e-15] / PF00317.14 [Ribonucleotide reductase alpha domain] [7.9e-19] / PF02867.8 [Ribonucleotide reductase barrel domain] [2e-194]	YP_001426398 [hypothetical protein FR483_N766L (Paramecium bursaria Chlorella virus FR483)] [0.0]	Q03604 [RecName: FullRibonucleoside-diphosphate reductase large subunit AltName: FullRibonucleotide reductase large subunit] [0.0]	Unknown protein	0	0	0
a630R	3E+05	3E+05	50	240	79	9246	12.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A631L	3E+05	3E+05	42	270	89	10392	9.9	N/A	Yes	N/A	N/A	YP_001426402 [hypothetical protein FR483_n770R (Paramecium bursaria Chlorella virus FR483)] [3e-08]	N/A	Unknown protein	0	0	0
a632L	3E+05	3E+05	64	219	72	7985	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	2
A633R	3E+05	3E+05	76	363	120	13184	9.2	Late	N/A	N/A	N/A	YP_001426396 [hypothetical protein FR483_N764L	N/A	Unknown protein	3	3	3

A646aL	3E+05	3E+05	84	126	41	5031	10.0	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A647R	3E+05	3E+05	26	570	189	22158	8.3	Early	N/A	N/A	PF07150.4 [Protein of unknown function (DUF1390)] [5.7e-46]	YP_001426682 [hypothetical protein ATCV1_Z201L (Acanthocystis turfacea Chlorella virus 1)] [1e-25]	N/A	Unknown protein	0	0	0
a648L	3E+05	3E+05	54	210	69	8097	8.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A649R	3E+05	3E+05	68	780	259	28689	9.0	Early	N/A	N/A	PF07150.4 [Protein of unknown function (DUF1390)] [9.1e-85]	YP_001426165 [hypothetical protein FR483_N533L (Paramecium bursaria Chlorella virus FR483)] [6e-29]	N/A	Unknown protein	0	0	0
a650L	3E+05	3E+05	72	264	87	10200	12.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a650aL	3E+05	3E+05	10	123	40	4447	7.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
a650bL	3E+05	3E+05	72	177	58	6871	11.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a650cR	3E+05	3E+05	40	183	60	7437	9.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A651L	3E+05	3E+05	80	693	230	26344	9.6	Early-Late	N/A	N/A	PF01541.17 [GIY-YIG catalytic domain] [2.1e-06] / PF07453.6 [NUMOD1 domain] [1.3e-11]	YP_001426012 [hypothetical protein FR483_N380R (Paramecium bursaria Chlorella virus FR483)] [5e-51]	N/A	Unknown protein	0	0	0
a652R	3E+05	3E+05	38	228	75	8583	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	0
a653R	3E+05	3E+05	26	198	65	7565	9.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A654L	3E+05	3E+05	78	594	197	22397	6.1	Early	N/A	N/A	PF00583.17 [Acetyltransferase (GNAT) family] [6.8e-06]	YP_001426628 [hypothetical protein ATCV1_Z147L (Acanthocystis turfacea Chlorella virus 1)] [4e-53]	N/A	Unknown protein	0	0	0
A655L	3E+05	3E+05	66	321	106	12002	11.4	N/A	Yes	N/A	N/A	N/A	N/A	Unknown protein	0	1	0
A656L	3E+05	3E+05	82	723	240	26607	5.0	Early	N/A	N/A	PF01391.11 [Collagen triple helix repeat (20 copies)] [9.1e-11]	YP_001426629 [hypothetical protein ATCV1_Z148L (Acanthocystis turfacea Chlorella virus 1)] [2e-11]	N/A	Unknown protein	2	2	2
A658R	3E+05	3E+05	54	288	95	11136	8.1	Late	N/A	N/A	N/A	N/A	N/A	Unknown protein	1	2	1
A659L	3E+05	3E+05	86	579	192	21545	3.4	Late	N/A	N/A	N/A	YP_001427157 [hypothetical protein ATCV1_Z676R (Acanthocystis turfacea Chlorella virus 1)] [2e-12]	N/A	Unknown protein	1	1	1
a660R	3E+05	3E+05	58	333	110	13462	11.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0

a661R	3E+05	3E+05	58	309	102	11735	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
A662L	3E+05	3E+05	74	516	171	19589	10.8	Early-Late	N/A	N/A	PF04117.5 [Mpv17 / PMP22 family] [2.1e-10]	YP_001425817 [hypothetical protein FR483_N185L (Paramecium bursaria Chlorella virus FR483)] [1e-54]	Q54FR4 [RecName: FullPXMP2/4 family protein 4] [8e-08]	Unknown protein	2	4	2
a663R	3E+05	3E+05	52	240	79	9660	11.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A664L	3E+05	3E+05	80	465	154	17176	5.6	Early	N/A	N/A	N/A	YP_001425783 [hypothetical protein FR483_N151R (Paramecium bursaria Chlorella virus FR483)] [1e-42]	N/A	Unknown protein	0	0	0
A665L	3E+05	3E+05	50	516	171	19498	9.8	Early	N/A	N/A	N/A	YP_001425783 [hypothetical protein FR483_N151R (Paramecium bursaria Chlorella virus FR483)] [1e-26]	N/A	Unknown protein	0	0	0
A666L	3E+05	3E+05	74	2757	918	####	6.2	Early	N/A	COG1126 [ABC-type polar amino acid transport system ATPase component] [1e-05]	PF00005.20 [ABC transporter] [2.7e-15]	YP_001426365 [hypothetical protein FR483_N733R (Paramecium bursaria Chlorella virus FR483)] [0.0]	Q75EV6 [RecName: FullElongation factor 3 ShortEF-3] [0.0]	Unknown protein	0	0	0
a667R	3E+05	3E+05	52	291	96	10975	8.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	1	0
a668R	3E+05	3E+05	50	198	65	7392	6.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a669R	3E+05	3E+05	50	252	83	9142	10.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a670R	3E+05	3E+05	46	369	122	13755	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a671L	3E+05	3E+05	78	294	97	11430	9.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
A672R	3E+05	3E+05	80	636	211	23295	5.2	Early	N/A	COG0666 [FOG: Ankyrin repeat] [2e-06]	PF00023.23 [Ankyrin repeat] [2.1e-09]	YP_002840971 [Ankyrin (Sulfolobus islandicus Y.N.15.51)] [2e-26]	Q4UMH6 [RecName: FullPutative ankyrin repeat protein RF_0381] [2e-17]	Unknown protein	0	0	0
a673L	3E+05	3E+05	46	207	68	7488	4.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A674R	3E+05	3E+05	84	651	216	24902	8.1	Early	N/A	COG1351 [Predicted alternative thymidylate synthase] [5e-07]	PF02511.8 [Thymidylate synthase complementing protein] [6.2e-55]	YP_001427299 [hypothetical protein ATCV1_2818L (Acanthocystis turfacea Chlorella virus 1)] [3e-78]	P73053 [RecName: FullThymidylate synthase thyX ShortTS ShortTSase] [9e-57]	Thymidylate synthase X	0	0	0
a675L	3E+05	3E+05	54	369	122	14417	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	2	2	2
A676R	3E+05	3E+05	70	1125	374	42432	10.6	Late	Yes	N/A	PF08789.3 [PBCV-specific basic adaptor domain] [1.9e-17] / PF08793.3 [2-cysteine adaptor domain] [1.8e-15]	YP_001425796 [hypothetical protein FR483_N164R (Paramecium bursaria Chlorella virus FR483)] [6e-55]	N/A	Unknown protein	0	0	0

a677L	3E+05	3E+05	70	231	76	8171	4.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A678R	3E+05	3E+05	90	1158	385	41287	10.3	Late	Yes	N/A	N/A	YP_001426663 [hypothetical protein ATCV1_Z182R (Acanthocystis turfacea Chlorella virus 1)] [7e-75]	N/A	Unknown protein	0	3	0
a679L	3E+05	3E+05	64	285	94	10193	11.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a680R	3E+05	3E+05	56	264	87	10337	3.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a681R	3E+05	3E+05	28	225	74	9708	13.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A681aL	3E+05	3E+05	54	177	58	7012	10.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A682L	3E+05	3E+05	82	1107	368	39915	5.8	Early	N/A	COG0666 [FOG: Ankyrin repeat] [8e-06]	PF00023.23 [Ankyrin repeat] [4.3e-10]	XP_001649474 [ankyrin 23/unc44 (Aedes aegypti)] [2e-51]	Q54KA7 [RecName: FullAnkyrin repeat PH and SEC7 domain containing protein secG] [3e-46]	Unknown protein	0	0	0
a682aL	3E+05	3E+05	52	138	45	5452	10.9	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
a682bR	3E+05	3E+05	38	144	47	5490	9.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A683L	3E+05	3E+05	74	1104	367	41847	7.9	Early-Late	N/A	COG0270 [Site-specific DNA methylase] [1e-06]	PF00145.10 [C-5 cytosine-specific DNA methylase] [5.6e-66]	YP_001426001 [hypothetical protein FR483_N369L (Paramecium bursaria Chlorella virus FR483)] [4e-73]	P38216 [RecName: FullModification methylase CvJI ShortM.CvJI AltName: FullCytosine-specific methyltransferase CvJI] [0.0]	Unknown protein	0	0	0
a684R	3E+05	3E+05	54	321	106	12535	8.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a685R	3E+05	3E+05	50	243	80	8650	6.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A686L	3E+05	3E+05	68	489	162	18316	6.9	Early	Yes	N/A	N/A	N/A	N/A	Unknown protein	0	1	0
a686aL	3E+05	3E+05	78	189	62	7504	8.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A685bR	3E+05	3E+05	78	171	56	6326	9.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a685cL	3E+05	3E+05	62	135	44	4817	9.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A687R	3E+05	3E+05	66	228	75	8976	4.4	Early	N/A	N/A	N/A	N/A	N/A	Unknown protein	0	0	0
a688L	3E+05	3E+05	62	216	71	7957	12.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	0
A689L	3E+05	3E+05	70	426	141	15974	8.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	1	1
a690R	3E+05	3E+05	36	237	78	8746	9.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a690aL	3E+05	3E+05	78	126	41	4982	10.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1

A690bL	3E+05	3E+05	78	144	47	5751	7.6	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	1	1	1
a690cR	3E+05	3E+05	36	126	41	5159	10.5	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a690dR	3E+05	3E+05	40	132	43	4606	12.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A690eR	3E+05	3E+05	44	147	48	5331	11.2	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A690fR	3E+05	3E+05	0	162	53	6037	11.8	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a690gR	3E+05	3E+05	64	156	51	5820	9.3	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
A691R	3E+05	3E+05	66	552	183	21545	9.4	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0
a692R	3E+05	3E+05	68	270	89	10191	9.1	N/A	N/A	N/A	N/A	N/A	N/A	Hypothetical protein	0	0	0

125 1 – Transmembrane regions of the protein were predicted by TMHMM (T), HMMTOP (H), and Phobius (P) methods. For all the method
126 default parameters were used for the prediction. The number shown in the table is the number of helices predicted by the methods.

127 Table S2. PBCV-1 virionic proteome and mass spectral data from each method

		Proteomic Methods and Parameters						
Protein	Method number	Method 1				Method 2		
		Mascot score	Queries matched	Unique peptides	emPAI*	Unique peptides	Coverage fraction	Number spectra
A010R	1&2	3408	53	25	6.21	19	0.5736	238
A011L	1&2	8196	113	29	13.68	22	0.7047	463
A014R	1&2	160	4	3	0.07	3	0.0226	20
A025/027/029L	1&2	161	3	2	0.07	3	0.028	49
A034R	1&2	839	43	13	2.2	17	0.4838	232
A035L	1&2	5397	100	23	2.48	30	0.6058	1175
A041R	1&2	1479	32	13	1.72	15	0.4515	281
A051L	1&2	690	12	8	1.6	9	0.601	78
A085R	1&2	1597	24	12	3.8	11	0.5868	131
A092/093L	1&2	1016	25	16	1.62	11	0.3118	223
A121R	1&2	41	4	3	0.28	6	0.4327	70
A127R	1&2	511	17	8	0.77	9	0.3469	66
A136R	1&2	109	11	6	0.46	4	0.2877	126
A139L	1&2	354	11	4	1	8	0.4503	294
A140 145R	1&2	11632	241	46	3	58	0.5209	3522
A165aL	1&2	491	13	8	2.14	6	0.5337	134
A168R	1&2	1439	25	7	1.76	7	0.494	569
A171R	1&2	405	12	7	0.25	4	0.1809	13
A176L	1&2	298	5	2	0.9	4	0.3636	166
A189 192R	1&2	1870	58	29	0.5	30	0.3087	511
A203R	1&2	1811	37	10	4.46	7	0.5463	192
A205R	1&2	664	27	8	2.05	8	0.2427	302
A213L	1&2	767	12	8	4.4	5	0.5541	147
A217L	1&2	844	26	13	0.88	16	0.5025	174
A227L	1&2	519	17	5	1.19	4	0.365	128
A230R	1&2	228	6	4	0.76	5	0.2806	20
A231L	1&2	313	11	8	0.43	16	0.5561	139
A237R	1&2	3551	79	38	5.72	31	0.7297	724
A245R	1&2	151	5	4	0.59	4	0.369	19
A246aR	1&2	60	2	1	0.2	7	0.6842	57
A252R	1&2	462	17	14	1.39	13	0.4064	107
A262/263L	1&2	169	8	6	0.7	14	0.543	632
A278L	1&2	3106	89	35	3.6	25	0.4213	822
A282L	1&2	5342	128	34	4.83	25	0.4499	1228
A284L	1&2	1996	38	11	2.41	13	0.7312	311

A286R	1&2	1284	28	16	1.42	15	0.5661	206
A296R	1&2	334	10	3	0.7	2	0.1911	101
A305L	1&2	873	22	12	3.44	14	0.6373	200
A310L	1&2	976	10	3	1.33	5	0.4824	180
A314R	1&2	43	3	1	0.38	3	0.4875	51
A316R	1&2	224	4	2	0.14	11	0.3014	95
A320R	1&2	486	10	4	0.79	3	0.2446	102
A322L	1&2	420	9	6	1.53	11	0.5341	229
A342L	1&2	2337	52	22	1.87	23	0.7188	1093
A349L	1&2	2517	51	13	11.17	10	0.5	759
A352L	1&2	1496	31	10	3.38	8	0.4589	1536
A363R	1&2	4467	120	46	2.01	55	0.5589	1112
A375R	1&2	419	14	7	1.24	6	0.3584	117
A378L	1&2	868	64	11	1.57	8	0.2375	306
A383R	1&2	946	17	12	1.06	13	0.4609	157
A384dL	1&2	2041	37	15	0.98	21	0.4548	659
A398L	1&2	330	5	4	1.53	6	0.4237	184
A405R	1&2	219	9	7	0.35	16	0.3387	252
A407L	1&2	584	12	7	0.94	10	0.6714	335
A413L	1&2	144	7	4	0.42	3	0.1475	41
A414R	1&2	144	7	4	0.75	2	0.5054	54
A421R	1&2	63	1	1	0.31	6	0.5306	230
A430L	1&2	34014	531	33	13.06	23	0.5995	2982
A437L	1&2	1500	29	9	19.75	11	0.7864	722
A440L	1&2	22	4	2	nd	2	0.2386	12
A448L	1&2	221	5	4	1.06	7	0.6132	160
A454L	1&2	3358	79	14	3.54	12	0.7474	1027
A465R	1&2	94	4	2	0.56	7	0.4576	119
A480L	1&2	47	3	1	0.35	2	0.2473	56
A484L	1&2	34	2	2	0.18	9	0.5548	43
A488R	1&2	2030	46	19	4.68	12	0.4732	365
A497R	1&2	291	8	2	1.69	7	0.3973	93
A500L	1&2	705	15	8	0.77	13	0.3722	764
A521aL	1&2	972	19	10	3.53	7	0.4257	118
A521L	1&2	605	9	5	0.93	6	0.4306	66
A523R	1&2	2560	49	18	24.79	11	0.8304	762
A526R	1&2	431	13	8	1.55	9	0.8014	328
A527R	1&2	223	7	6	2.69	6	0.4242	73
A532L	1&2	82	4	2	0.4	4	0.519	430
A533R	1&2	200	3	3	0.35	3	0.0722	4
A534R	1&2	66	1	1	0.31	3	0.4857	30
A535L	1&2	260	3	1	0.42	2	0.4085	60
A536L	1&2	234	3	1	0.42	5	0.5342	331

A558L	1&2	2749	47	24	6.04	13	0.5025	133
A559L	1&2	599	14	7	1.84	9	0.3474	178
A561L	1&2	2045	69	24	1.44	19	0.4206	331
A565R	1&2	4321	70	26	2.49	33	0.6045	709
A572R	1&2	233	9	6	1.11	11	0.7293	125
A577L	1&2	54	2	2	0.22	3	0.2105	13
A579L	1&2	663	17	6	0.77	9	0.4706	101
A605L	1&2	72	6	3	0.42	8	0.5633	220
A614L	1&2	1895	51	26	2.59	25	0.5338	571
A622L	1&2	3903	71	29	4.19	18	0.5115	261
A627R	1&2	1547	28	15	1.77	19	0.5483	384
A643R	1&2	1337	50	26	3.22	12	0.2774	409
A644R	1&2	2024	40	16	17.33	10	0.5723	370
A676R	1&2	913	31	21	3.1	20	0.5561	302
A678R	1&2	768	16	10	1.51	13	0.4494	232
A157L	2					3	0.2273	25
A164aR	2					4	0.5424	38
A174L	2					2	0.1385	43
A196L	2					4	0.3289	57
A202L	2					4	0.4513	44
A350R	2					2	0.2459	13
A384bL	2					2	0.5	300
A400R	2					2	0.2288	5
A420L	2					2	0.3714	7
A423R	2					2	0.1465	7
A438L	2					2	0.3077	32
A502L	2					7	0.4105	215
A520L	2					4	0.5	30
A531L	2					2	0.5821	176
A532aL	2					3	0.32	35
A612L	2					2	0.2437	6
A018L	1	193	2	1	0.05			
A122/123cLz	1	123	16	2	0.02			
A122/123R	1	14	3	1	nd			
A137R	1	24	1	1	nd			
A172aL	1	18	6	1	nd			
A173L	1	16	3	1	0.1			
A188aR	1	17	7	4	0.19			
A201aL	1	18	8	1	nd			
A201L	1	53	1	1	0.35			
A219 222 226R	1	14	5	2	0.04			
A255R	1	35	7	2	0.2			
A256/257L	1	26	14	6	0.03			

A260aR	1	16	4	2	0.44
A271L	1	13	3	1	nd
A273L	1	133	3	2	0.22
A287R	1	13	9	5	0.11
A295L	1	14	2	1	nd
A304R	1	14	2	2	nd
A321R	1	38	1	1	0.26
A339L	1	17	5	1	0.48
A356R	1	19	3	1	nd
A436L	1	322	8	2	1.3
A443R	1	250	133	3	0.09
A456L	1	31	5	4	0.04
A476R	1	33	4	1	nd
A540L	1	199	3	3	0.17
A548L	1	20	14	5	0.06
A567L	1	18	1	1	0.19
A571R	1	28	7	3	0.26
A586R	1	15	1	1	nd
A598L	1	13	4	3	0.08
A617R	1	23	5	3	nd
A621L	1	165	3	1	0.26
A624R	1	103	1	1	0.25
A625R	1	15	5	3	0.07
A629R	1	14	10	4	0.04
A631L	1	17	3	1	nd
A655L	1	23	11	3	0.28
A686L	1	17	31	1	0.18

128

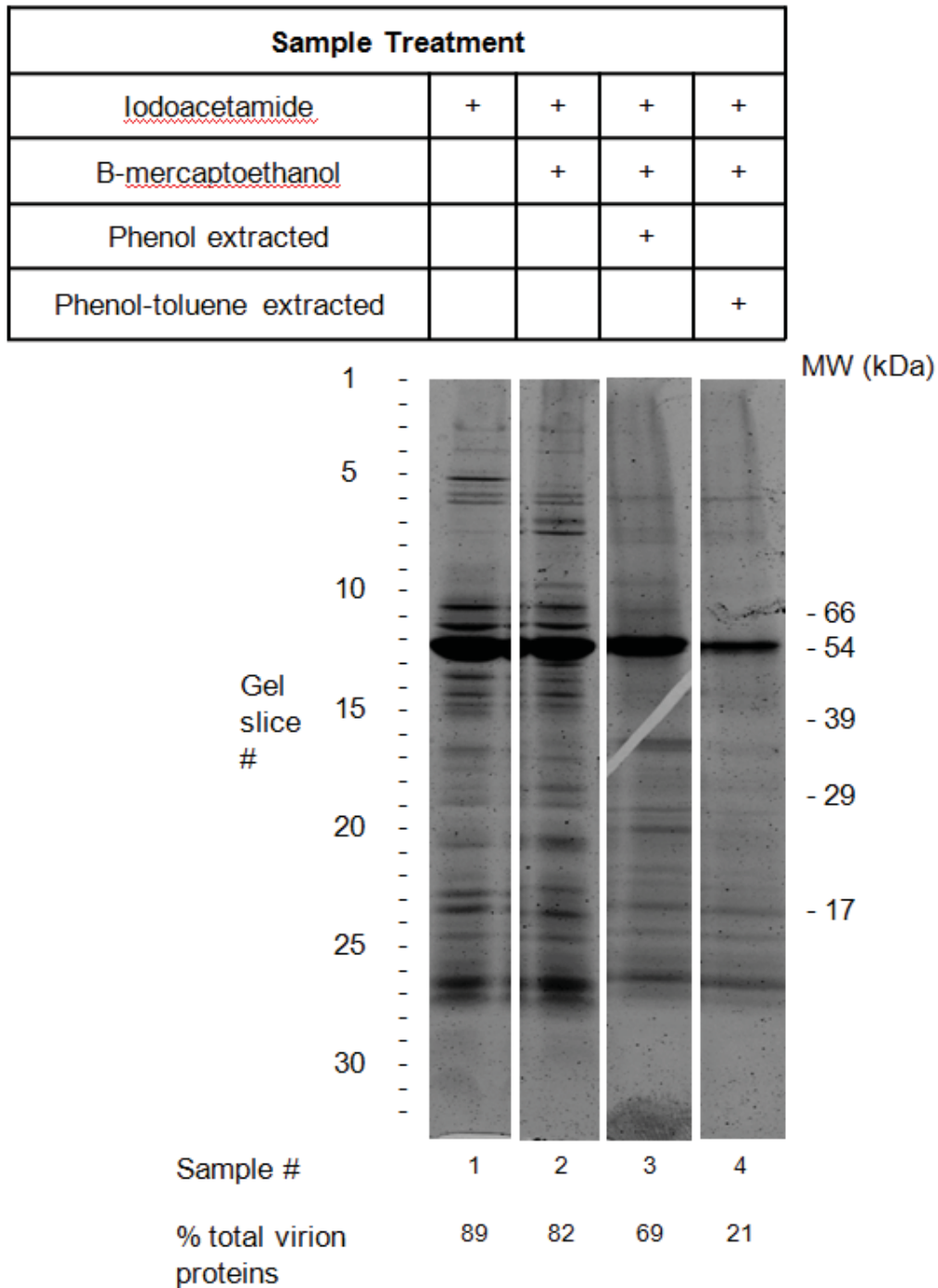
129 *emPAI: exponentially-modified protein abundance index

130

131 Table S3. Chlorovirus accession numbers and capsid protein paralog classes

Chlorovirus	CDS	NCBI Accession Number	Capsid protein paralog class
AR158	C048R	YP_001498130.1	III
AR158	C470R	YP_001498551.1	IV
AR158	C558L	YP_001498639.1	I
AR158	C675L	YP_001498756.1	III
AR158	C741L	YP_001498822.1	II
ATCV-1	Z151L	YP_001426632.1	II
ATCV-1	Z280L	YP_001426761.1	I
ATCV-1	Z506L	YP_001426987.1	III
ATCV-1	Z558L	YP_001427039.1	IV
ATCV-1	Z664R	YP_001427145.1	III
FR483	N074R	YP_001425706.1	III
FR483	N254L	YP_001425886.1	III
FR483	N274L	YP_001425906.1	III
FR483	N395R	YP_001426027.1	IV
FR483	N470L	YP_001426102.1	I
FR483	N737L	YP_001426369.1	II
MT325	M078R	ABT13632	III
MT325	M269L	ABT13823	III
MT325	M381R	ABT13935	IV
MT325	M463L	ABT14017	I

MT325	M748L	ABT14302	II
NY-2A	B059R	YP_001497255.1	III
NY-2A	B529R	YP_001497725.1	IV
NY-2A	B585L	YP_001497781.1	I
NY-2A	B617L	YP_001497813.1	I
NY-2A	B748L	YP_001497944.1	III
NY-2A	B825L	YP_001498021.1	II
PBCV-1	A010R	NP_048358.2	III
PBCV-1	A011L	NP_048359.1	III
PBCV-1	A383R	NP_048739.2	IV
PBCV-1	A384dL	NP_048747.2	V
PBCV-1	A430L	NP_048787.1	I
PBCV-1	A558L	NP_048914.1	III
PBCV-1	A622L	NP_048978.1	II



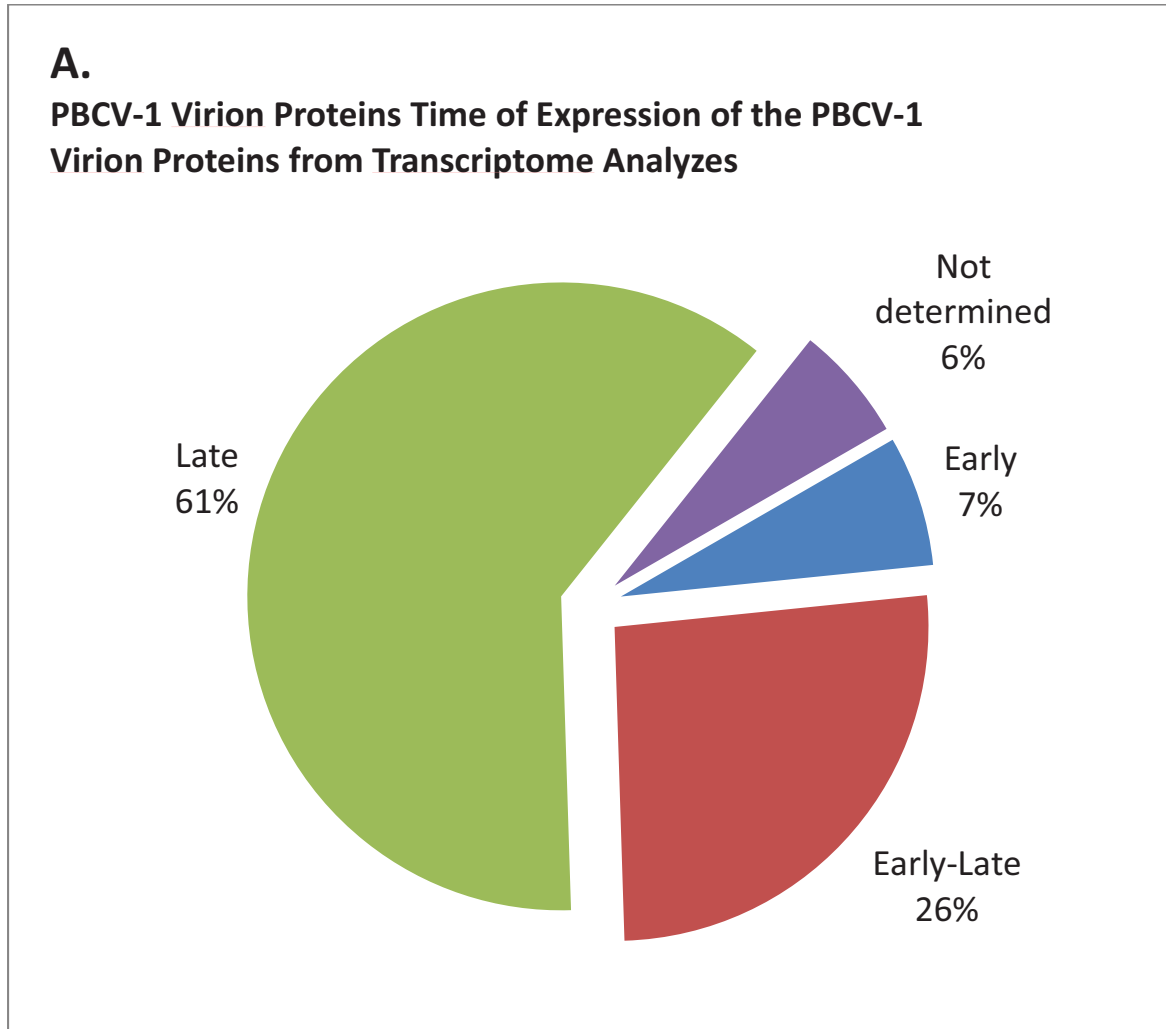
133

134 **Figure S1.** Protein extraction and separation of PBCV-1 virions using SDS-PAGE. Gradient

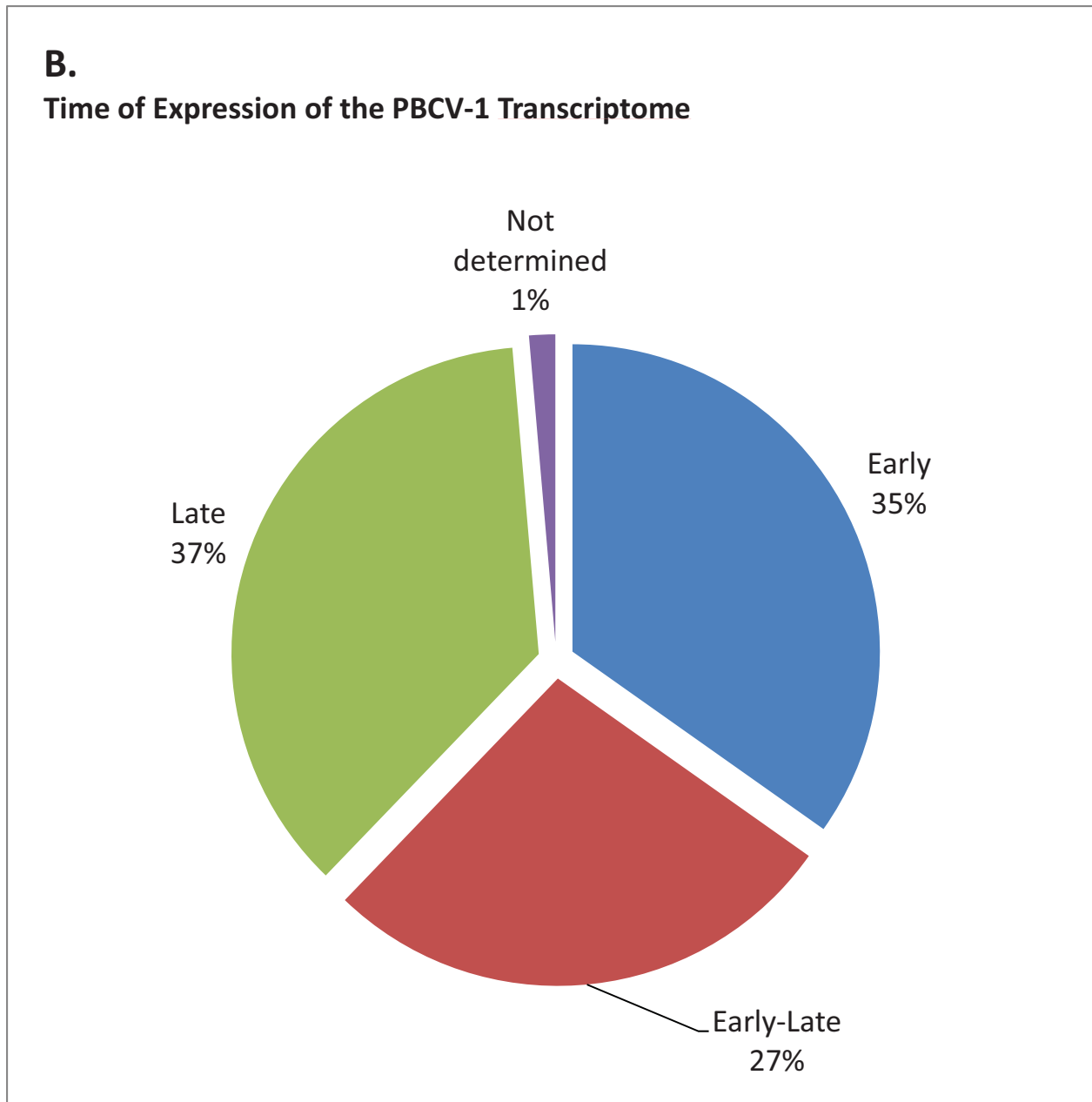
135 purified virions were solubilized and treated with various reagents, then separated on SDS-

136 PAGE. The gels were cut into 32 pieces and each gel piece was analyzed for tryptic peptide

137 fragments by Method 1. All the samples were treated with iodoacetamide; sample 2 was treated
138 with beta-mercaptoethanol or dithiothreitol; sample 3 was treated as # 2 with phenol extraction;
139 sample 4 was treated as #2 with phenol-toluene extraction. The percent of the total virion
140 proteins detected with each method is at the bottom of the figure.



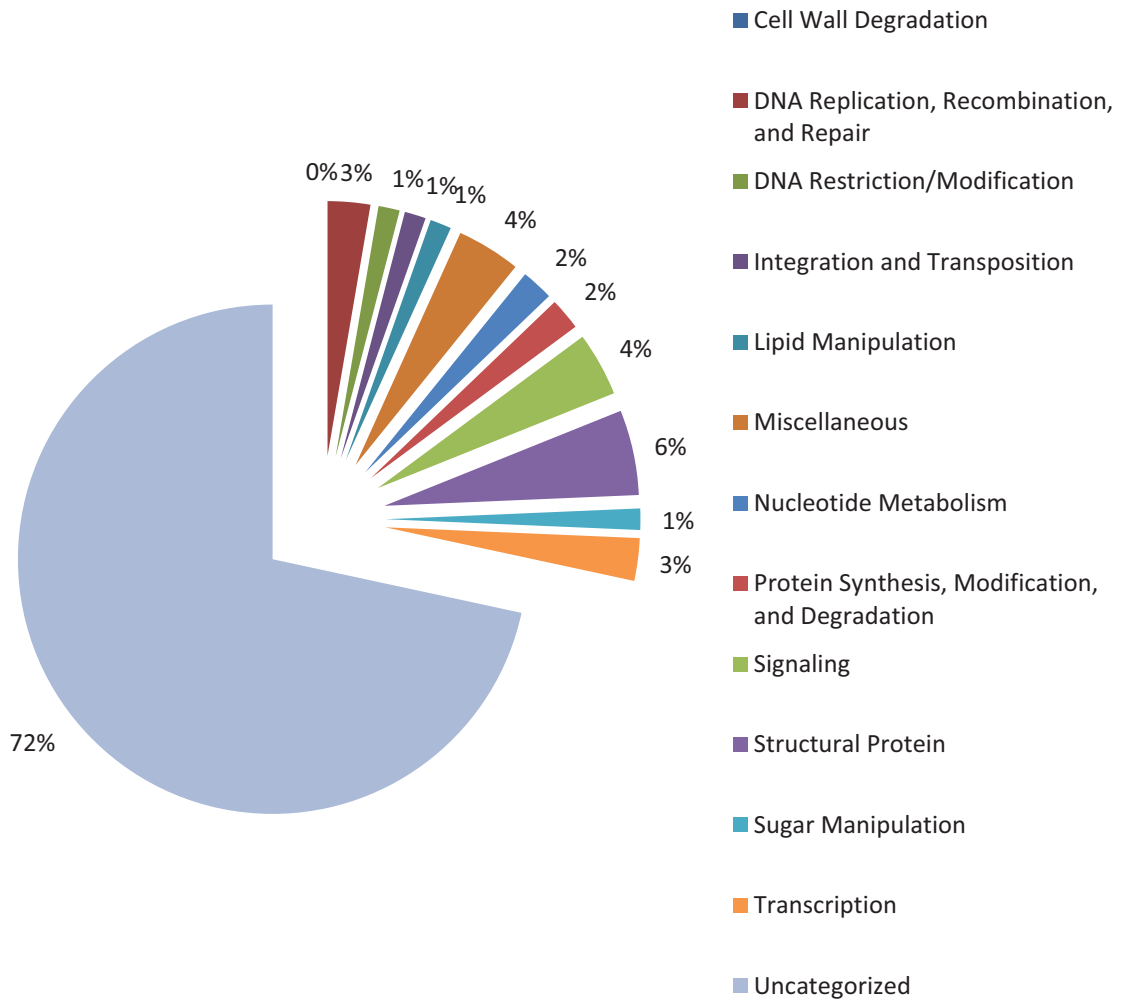
141



142 **Figure S2. (A)** Distribution of PBCV-1 genes as measured during the time course of replication.
143 (A) The expression stages and relative abundances of the total CDSs as measured by microarray
144 sequence hybridizations (10). **(B)** The expression stages of the genes associated with the virion
145 proteome. Note, the fractions labeled “not determined” are genes that were not included in
146 the microarray analyses.

A

PBCV-1 Virion Proteome Functional Distribution



B

PBCV-1 Total CDSs Functional Distribution

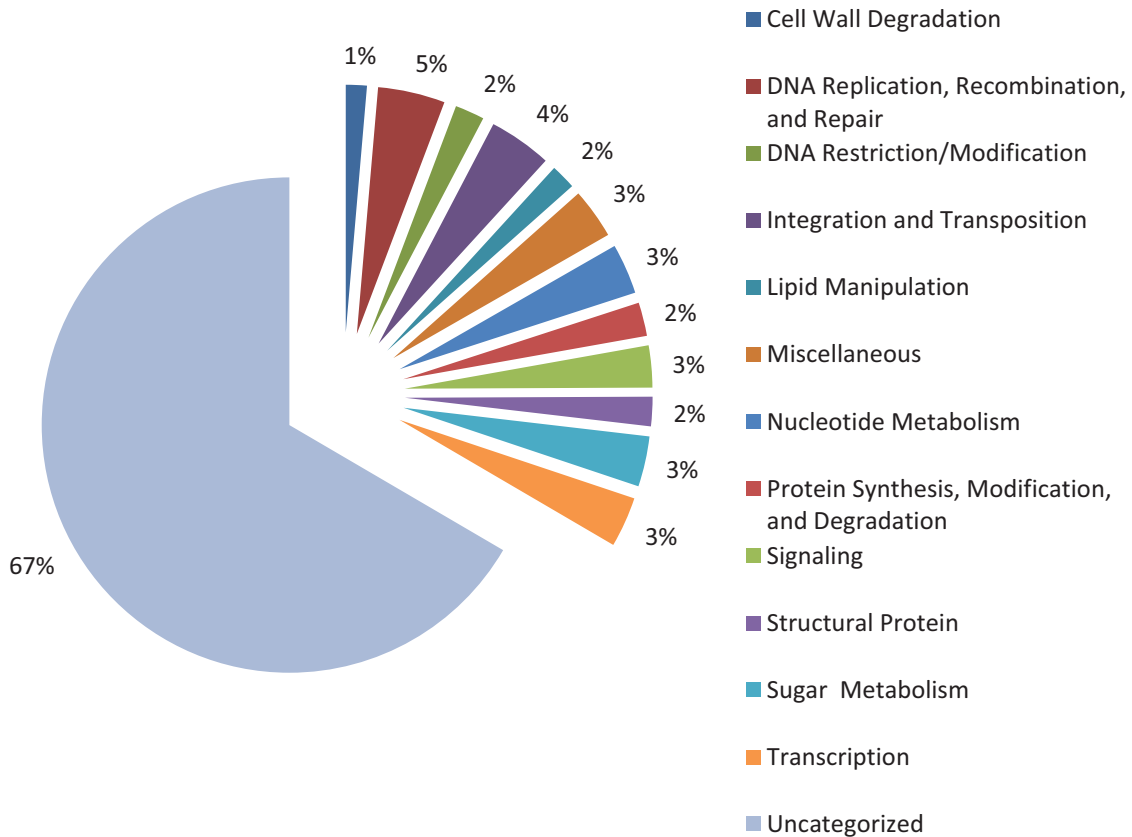


Figure S3. Functional distribution of PBCV-1 transcriptome and virion-associated proteome. (A) The functional distribution of the virion-associated proteins into 13 functional categories. The functional categories were determined from either sequence similarity analyses against the GenBank non-redundant database, or the proteins had activities determined biochemically. Note, there were no proteins detected with cell wall degradation associated with the virion proteins. (B) The 416 PBCV-1 coding CDSs are distributed into 13 functional categories.

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