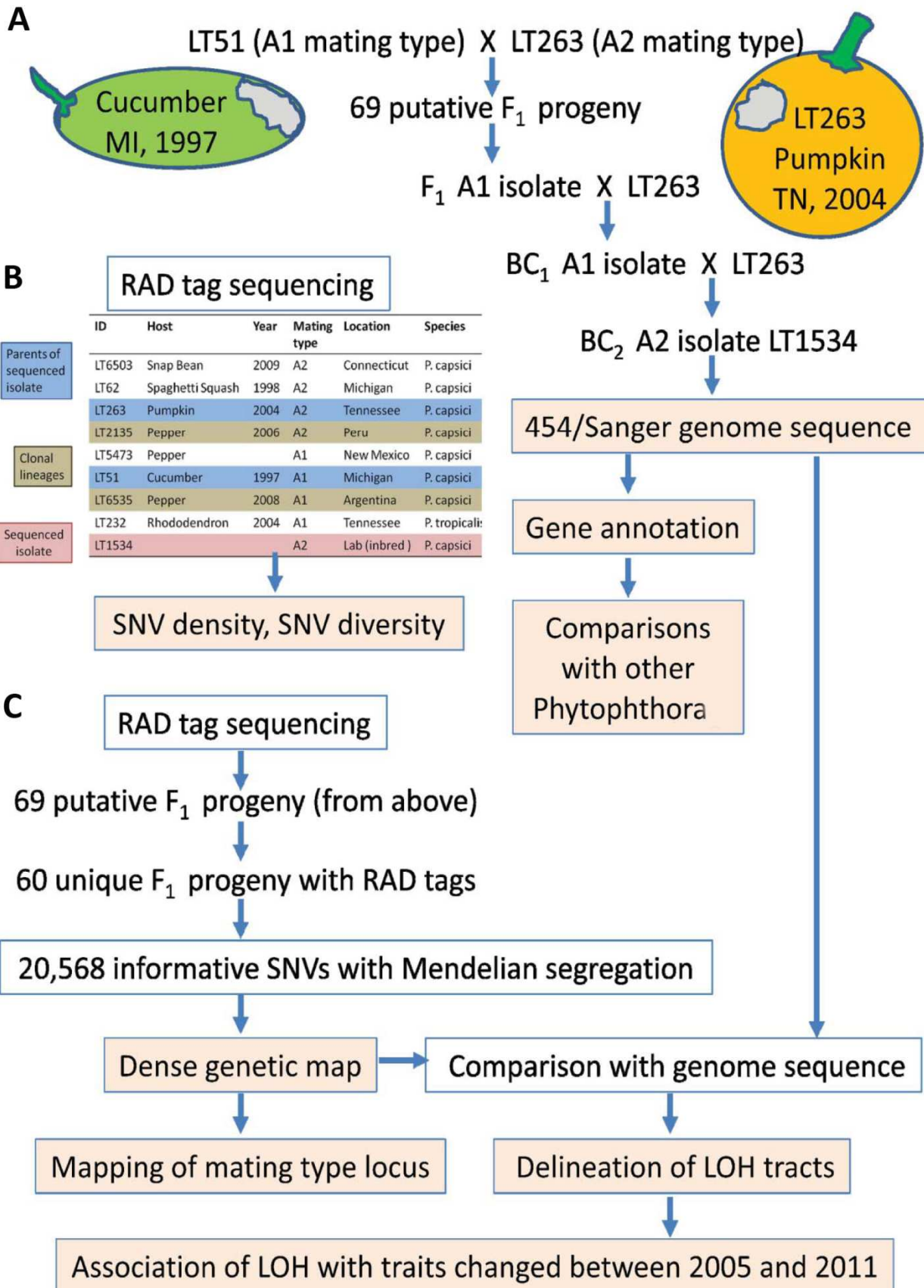
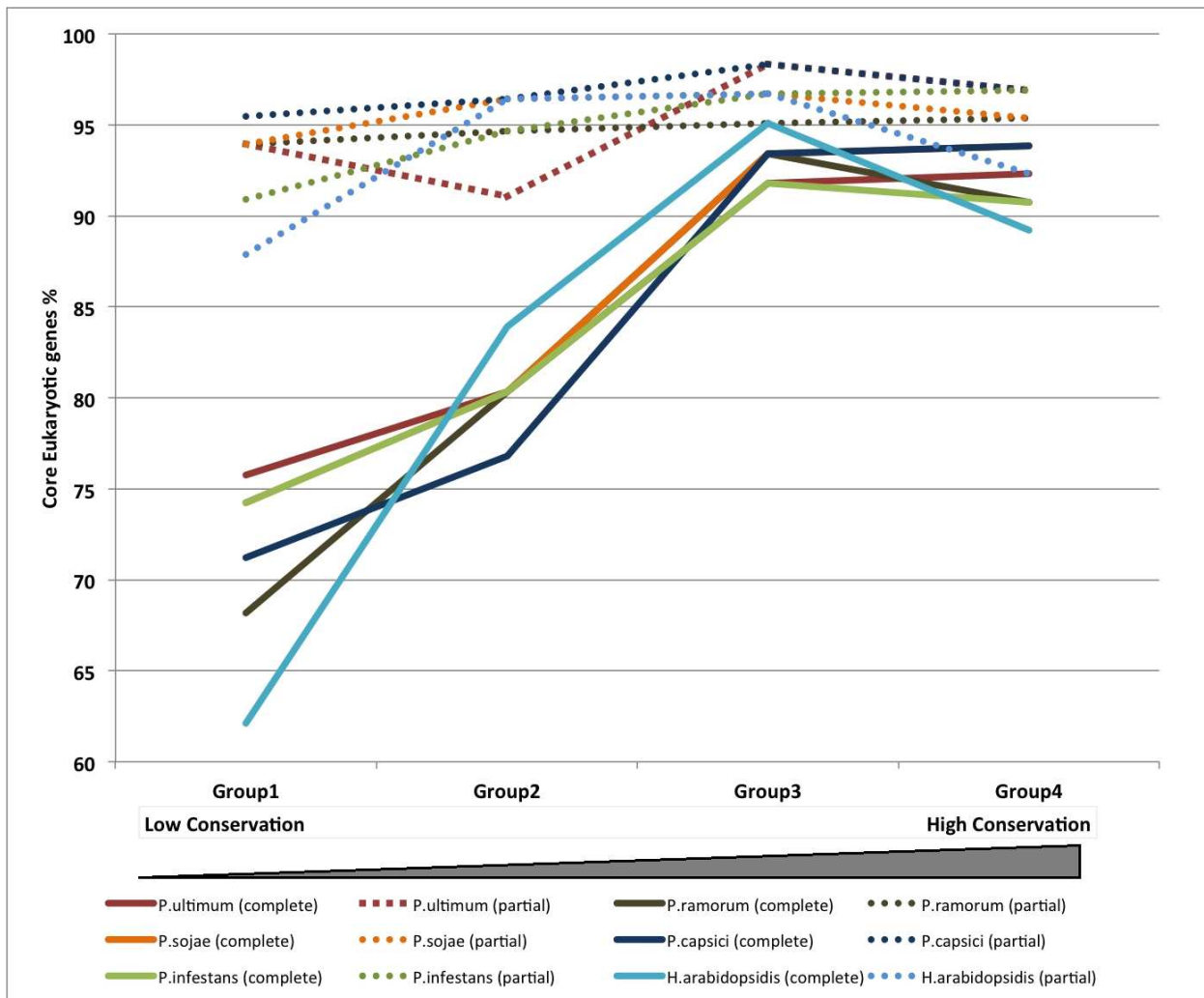


e-Extra supplementary figures and tables

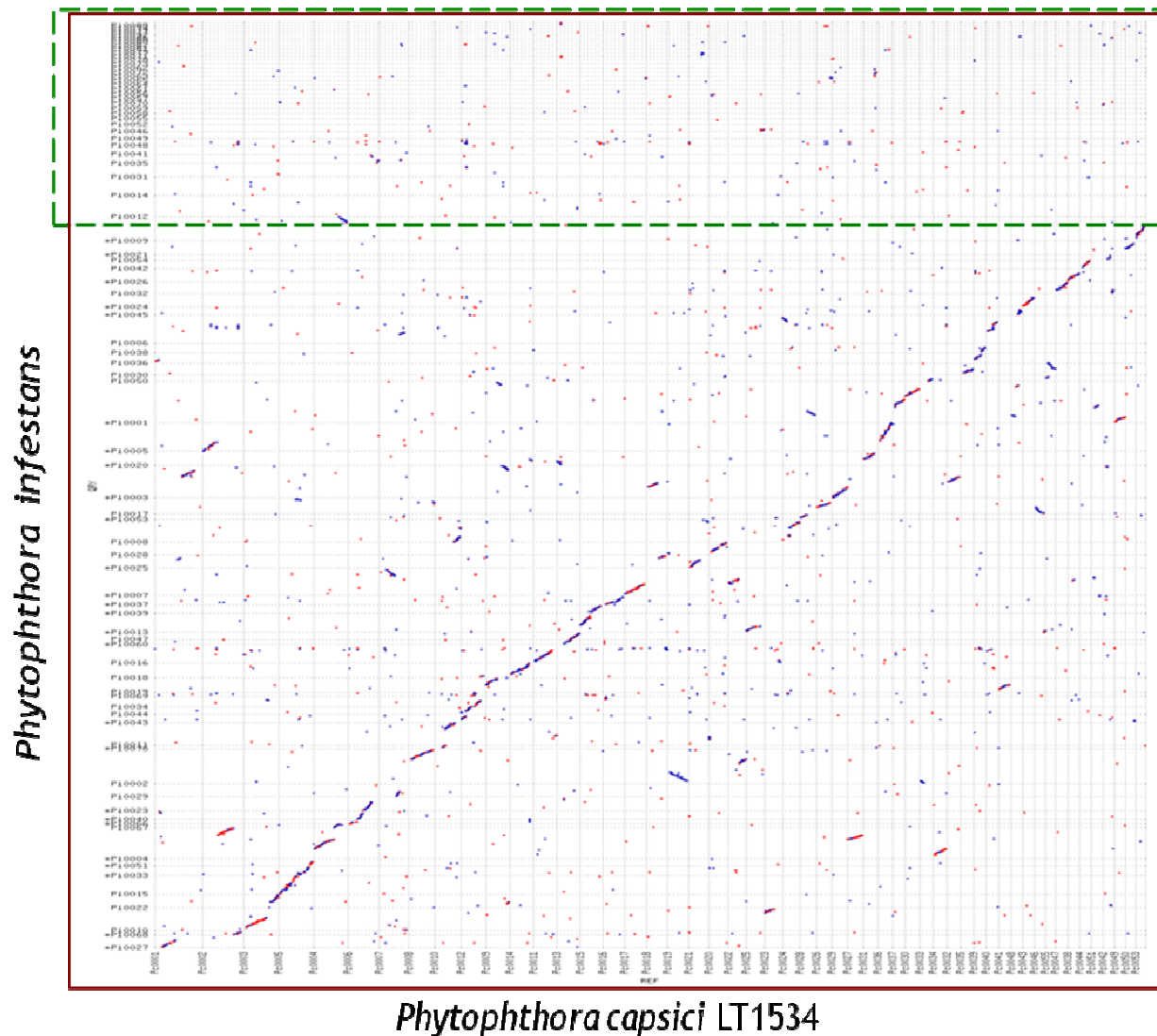
Supplementary Figure 1: Experimental design for molecular and genetic analysis of the vegetable pathogen *P. capsici*. (A) Two parental isolates of opposite mating types were crossed. Of 69 putative F₁ progeny, one was backcrossed (BC₁) to parent LT263. A second backcross (BC₂) yielded a moderately inbred isolate (LT1534) for genome sequencing. A hybrid Sanger/454 sequence assembly was performed and gene models were annotated both *ab initio* and using substantial EST and cDNA sequences generated for this project. The *P. capsici* genome and gene content was compared with other *Phytophthora* species. (B) LT1534, the parents, five other *P. capsici* isolates and a *P. tropicalis* isolate underwent reduced representation re-sequencing (RAD, Restriction site-Asociated DN_A sequencing) by SBS of 72nt tags flanking *SgrAI* sites. Single nucleotide variants (SNVs) were identified by alignment to the draft reference. SNV density and diversity were calculated. (C) *PstI* RAD was performed on 60 unique F₁ progeny from the original cross. 20,568 SNVs had simple Mendelian inheritance and were present in ≥55 of the progeny. The program JoinMap 4.1 was used to test the inheritance of markers from the largest ~100 scaffolds and to generate a genetic map. 18 linkage groups included 90% of the 19,805 predicted genes and covered 84% of the draft genome. Graphical genotypes revealed minimally spanning tracts of LOH in 23 progeny that ranged in length from 299bp to >1Mbp and spanned more than 30% of the genome. LOH was associated with a mating type switch in 4 isolates and with a change in pathogenicity in two isolates over the course of the study.



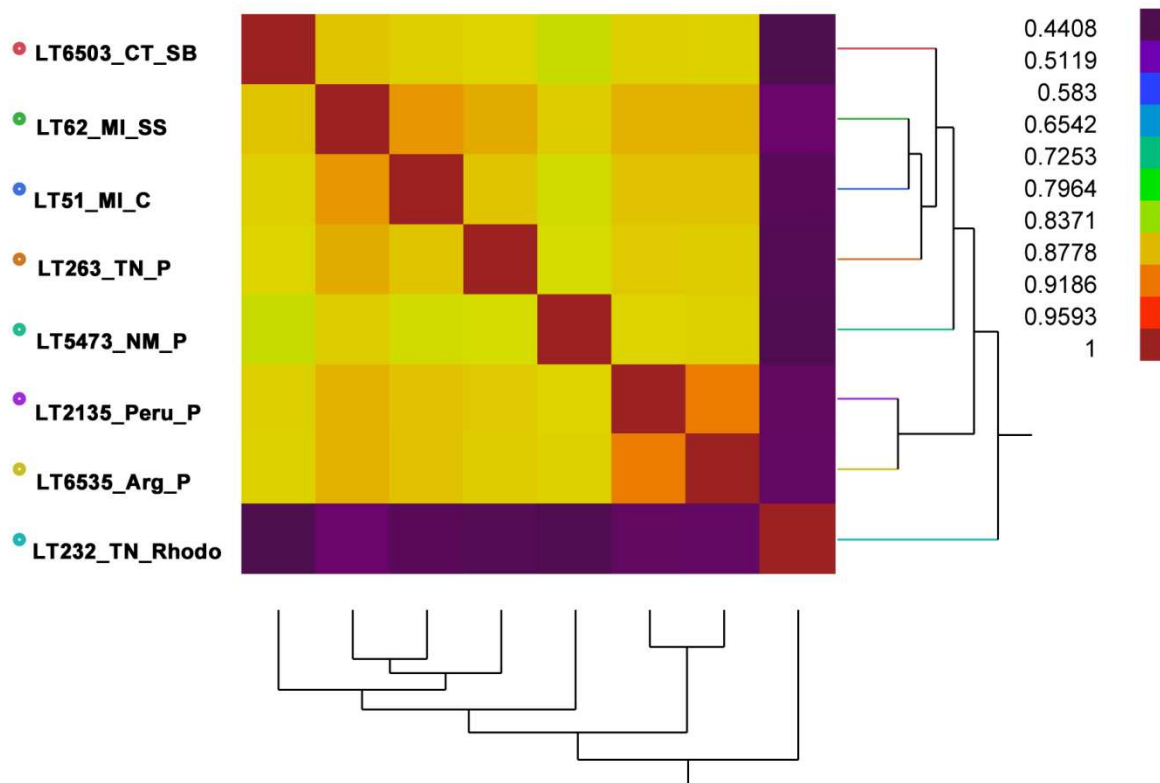
Supplementary Figure 2. Graphical display of CEGMA analyses using standard parameters. *P. capsici* has the highest number of identified core orthologous genes among all oomycete genomes of the *Peronosporaceae* and a comparable amount of fully covered genes. The increase in fully covered genes towards the most conserved ones is a result of the algorithm used for assessing gene completeness in the CEGMA analysis. The high percentage of fully covered genes for CEGMA groups 3 and 4 (most conserved), supports the high continuity of the assembly.



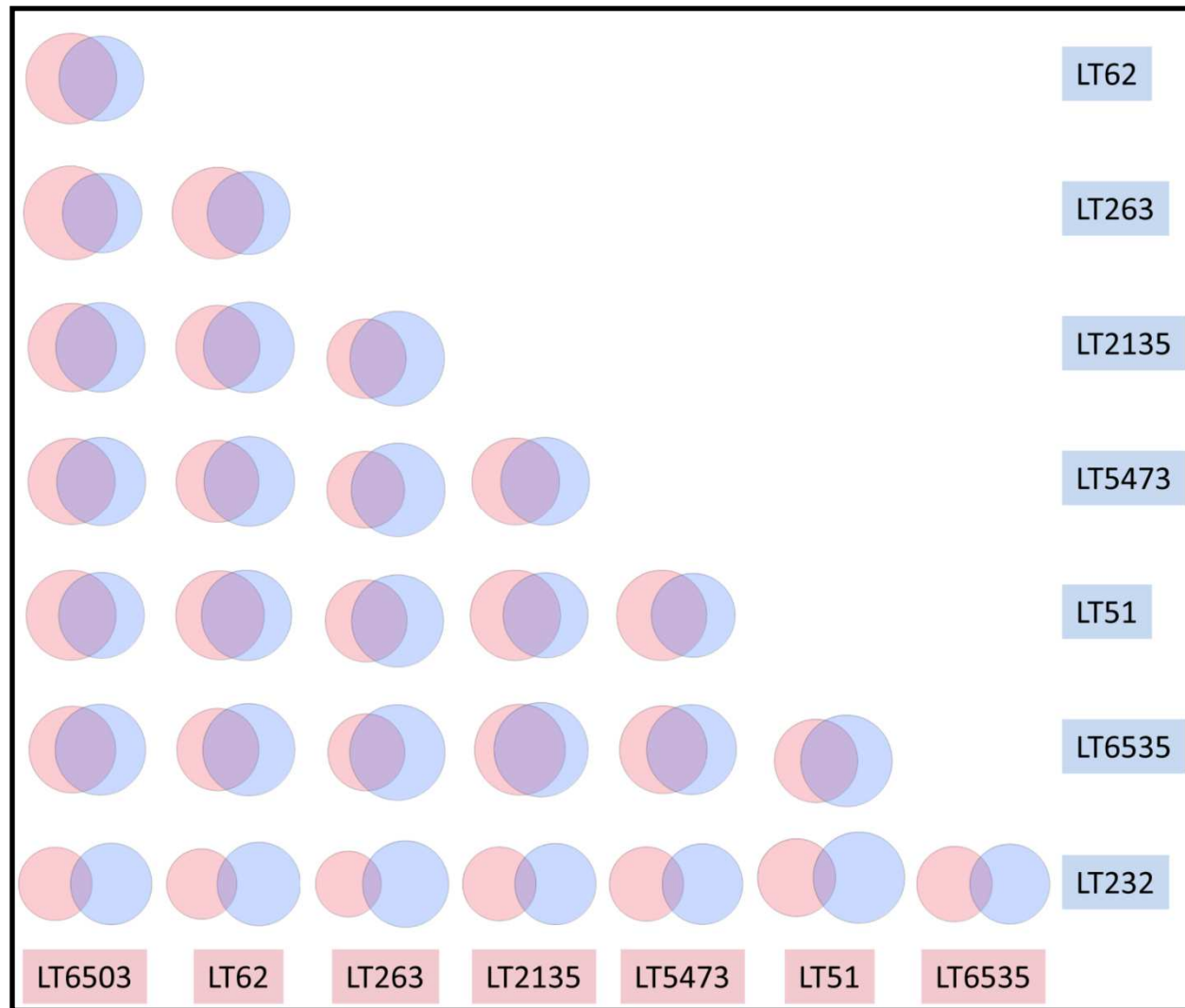
Supplementary Figure 3: Pairwise comparisons of gene models from *P. capsici* with *P. infestans* showed good synteny with some duplications and translocations. Sequences other than gene models were replaced with Ns. Nucleotide sequences were compared using PROMer package of MUMmer, which identifies Maximal Unique Matches in all 6 frames as anchors for amino-acid-based alignment. Large scaffolds (>450 kbp) are shown (155 Mbp for *P. infestans* and 46 Mbp for *P. capsici*). Scaffolds were ordered to maximize the center diagonal. There were some translocations, which possibly were assembly errors. A region (green box) of *P. infestans* had no syntentic genes in *P. capsici*.



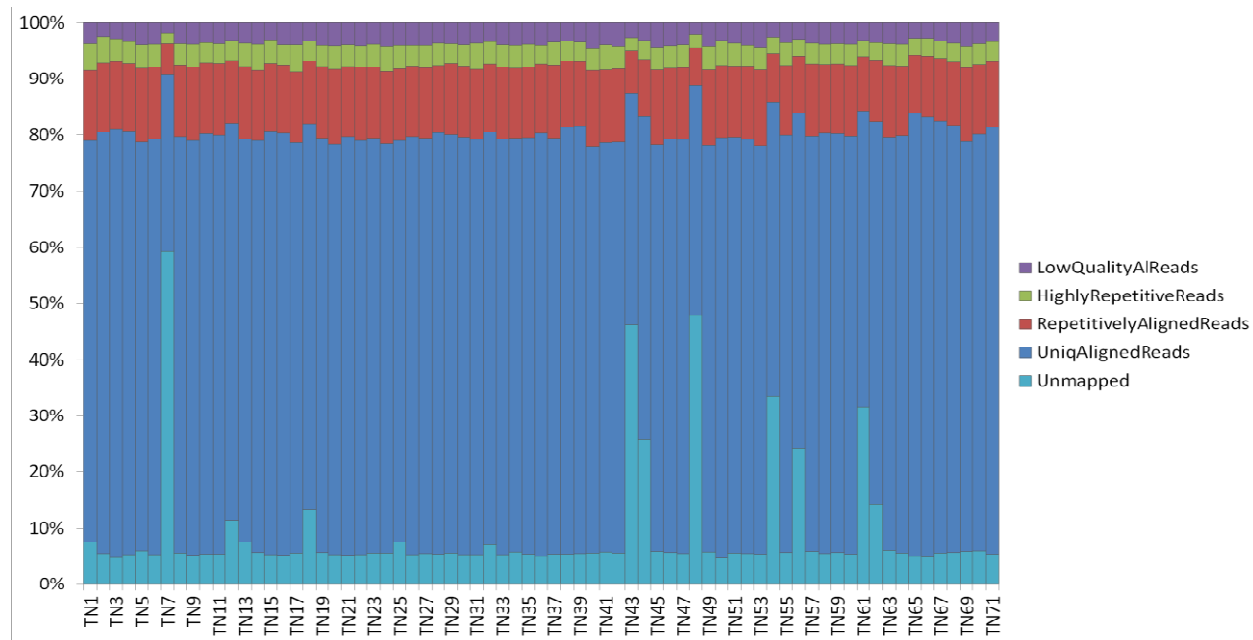
Supplementary Figure 4: Dendrogram and cross correlations for SNVs from 7 *P. capsici* and a *P. tropicalis* (LT232) isolate at regions with 30X coverage in all samples. Left side of the panel indicates isolate origin as abbreviated US state, Peru, or Argentina (Arg) followed by host designated as SB = snap bean, SS = summer squash, C = cucumber, P = pepper, and Rhodo = rhododendron. Branches on the similarity tree on the right side of the panel correspond to isolate directly across. Tree on bottom of panel is identical (isolate LT6503 on the far left) and correlations are designated by color legend on right.



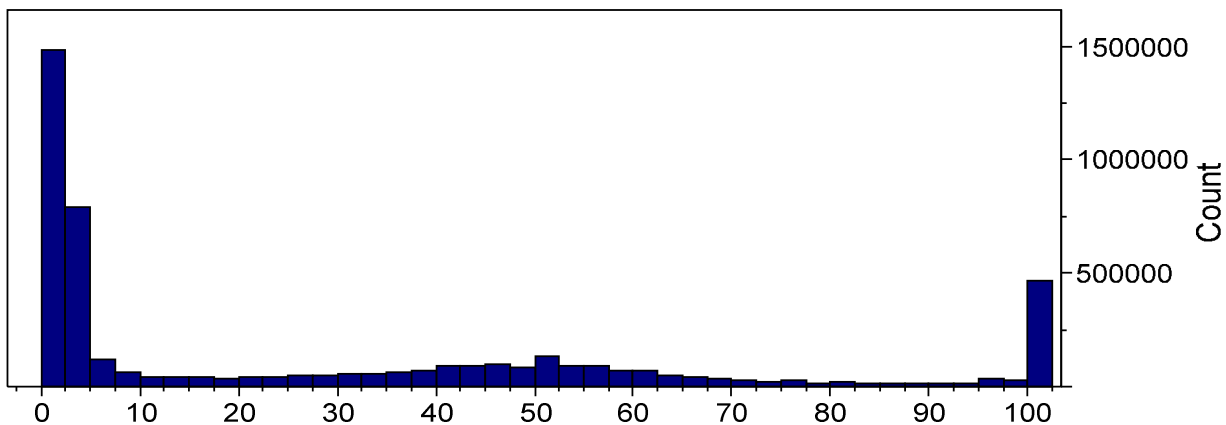
Supplementary Figure 5: Overlap of SNV content of 7 *P. capsici* isolates and one *P. tropicalis* isolate (LT232) sequenced by RAD. SNVs were called at sites with 30X coverage in all samples if supported by ≥ 4 uniquely aligning reads of Q score ≥ 20 . Of note, LT263 (recurrent parent of the LT1534) had the fewest SNVs of all *P. capsici* isolates; the two clonal lineages (host = pepper) showed the most overlap; *P. tropicalis* (LT232) showed the least overlap. A total of 64, 537 SNV positions are included. Positions with at least one alternate allele per isolate are LT6503 (20,445), LT62 (17,914), LT263 (14,637), LT2135 (20,869), LT5473 (21,386), LT51 (18,527), LT6535 (22,431), and LT232 (25,308).

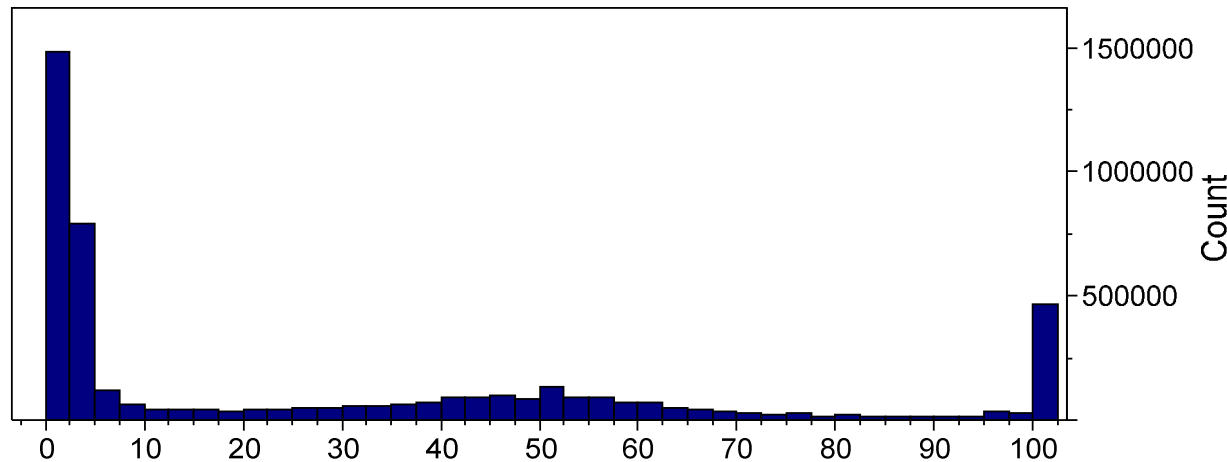


Supplementary Figure 6: Relative abundance of RAD sequences from 69 progeny and two parents based on alignment to the reference *P. capsici* sequence. The X-axis shows isolate ID's and Y-axis shows relative abundance.

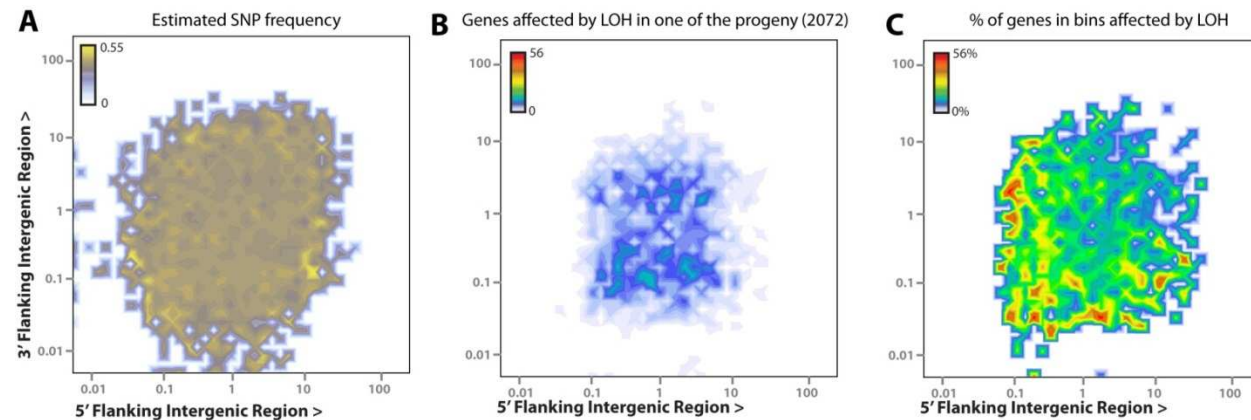


Supplementary Figure 7. Frequencies of alleles based on the requirement for 10X and 20X unique coverage and an average quality of >20 for the alternate alleles. For 10X unique coverage (top) this includes 5,849,663 SNVs and for 20X (bottom) this includes 4,449,193 SNVs across 71 samples.



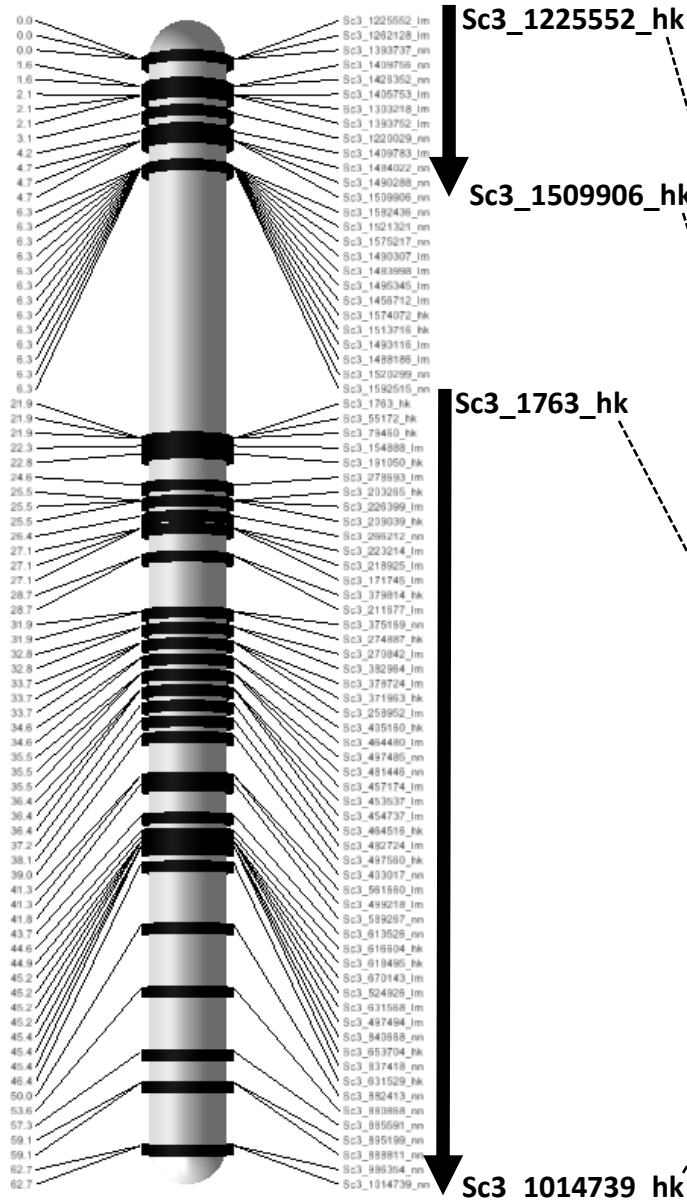


Supplementary Figure 8: Gene-poor environments in *P. capsici* genome show reduced LOH frequency. *P. capsici* genome architecture illustrated by the distribution of all predicted genes according to the length of their 5' (X-axis) and 3' (Y-axis) intergenic regions, counted by two-dimensional data binning. (A) Estimated SNP frequency is homogenous across the *P. capsici* genome. The SNV frequency was calculated across each individual scaffold (in SNPs per Kbp) and used as an estimate for local SNV frequency of each gene. The average of frequencies for genes in each bin is shown. (B) Genome architecture diagram showing only the 2072 genes affected by LOH in at least one re-sequenced strain from the progeny. (C) LOH frequency is reduced for genes residing in a gene-poor environment. The percentage of genes in bins affected by LOH in at least one re-sequenced strain from the progeny.

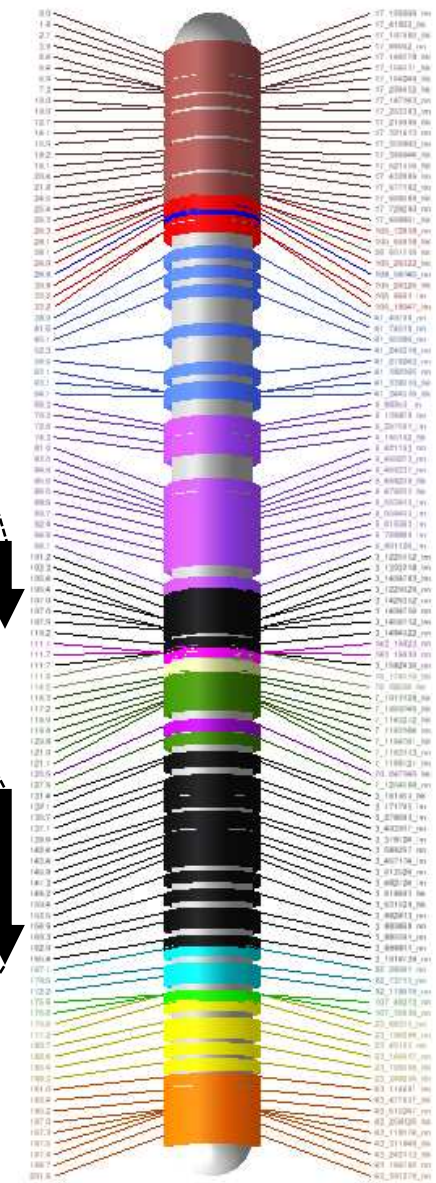


Supplementary Figure 9: Comparison of the genetic map of *P. capsici* Linkage Group 1 (202cM) with the physical map of Scaffold 3. Two blocks of Scaffold 3 comprising 1.6Mbp contained correctly ordered markers on Linkage Group 1. However, an assembly error occurred between these blocks.

Assembly Scaffold 3



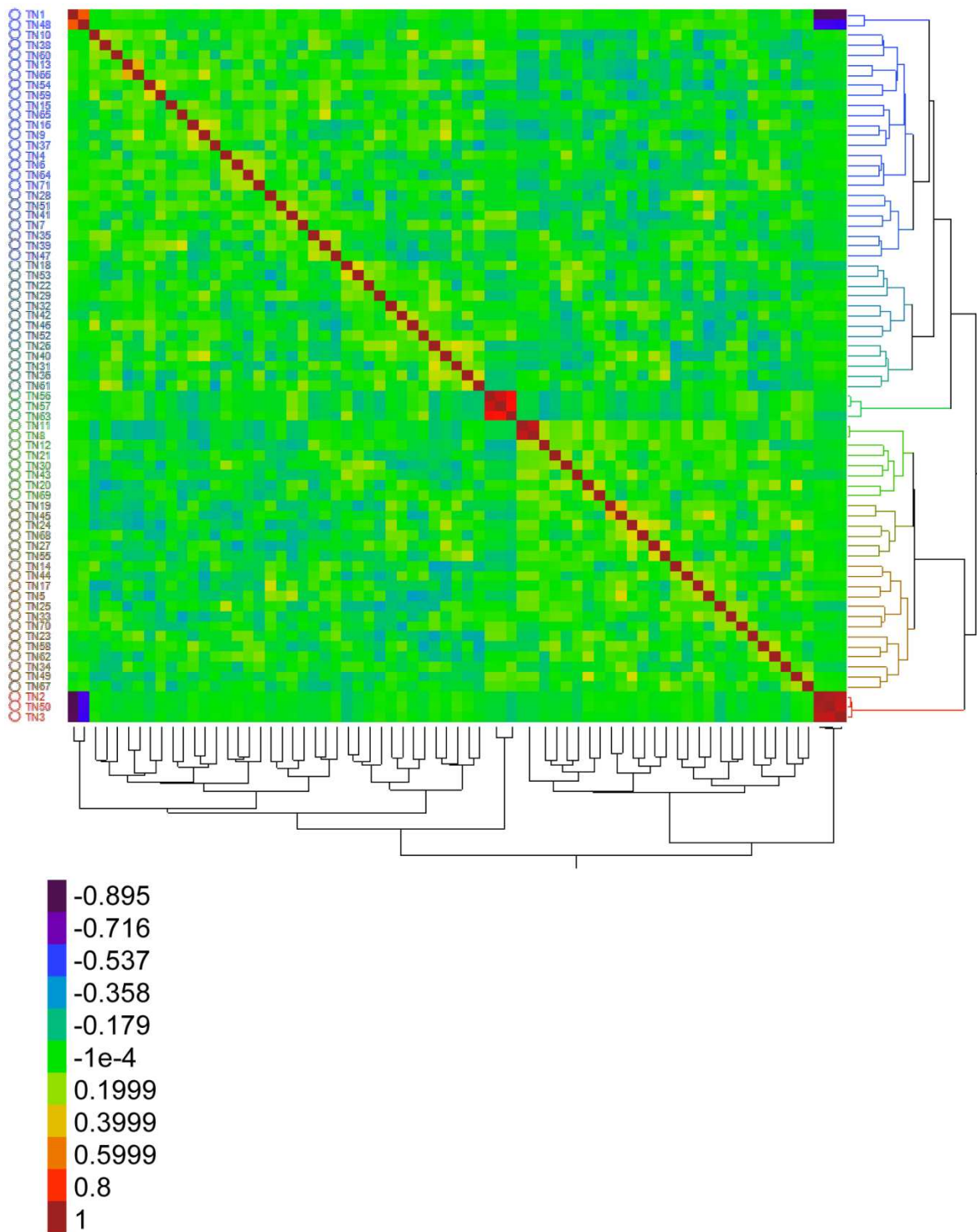
Linkage Group 1



Supplementary Figure 10: Graphical genotypes for four progeny with co-occurring LOH tracts. Genotypes are designated as 1 (yellow) = RR, 2 (gray) = AR, and 3 (blue) = AA (R = reference allele and A = alternate allele). Parent (TN1 and TN2) and progeny (TN63, TN18, TN34, and TN47) genotypes span 38 markers and 53,404 nt on scaffold 2. Columns 4-6 list χ^2 test values (>0.05 in green). Column 7 lists “yes” if all non-LOH progeny are AR for an RR x AA marker. Columns 12 – 15 indicate gene context, the predicted reference amino acid (or “silent” if mutation is synonymous), the alternate amino acid, and the gene model. Note fixation of markers 2_1447270 and 2_1486458 for the reference and alternate allele and the three different composite haplotypes across the entire tract of LOH.

SNP ID	TN1	TN2	RR:AR	RR:AR:AA	AR:AA	Fix	TN63	TN18	TN34	TN47	Gene	Ref	Alt	Gene Model
2_1437332	2	1	0.58	0.00	0.00		3	3	1	1	C	Silent		e_gw1.2.451.1
2_1437339	2	1	0.41	0.00	0.00		1	1	3	1	C	Silent		e_gw1.2.451.1
2_1437367	1	2	0.68	0.00	0.00		1	1	1	1	I			e_gw1.2.451.1
2_1437372	3	2	0.00	0.00	0.49		3	3	3	1	I			e_gw1.2.451.1
2_1437398	2	1	0.41	0.00	0.00		1	1	3	1	I			e_gw1.2.451.1
2_1437400	2	1	0.41	0.00	0.00		1	1	3	1	I			e_gw1.2.451.1
2_1437402	2	1	0.41	0.00	0.00		1	1	3	1	I			e_gw1.2.451.1
2_1437403	2	1	0.41	0.00	0.00		1	1	3	1	I			e_gw1.2.451.1
2_1437404	2	1	0.41	0.00	0.00		1	1	3	1	I			e_gw1.2.451.1
2_1441337	3	2	0.00	0.00	0.59		3	3	3	1	C	Silent		e_gw1.2.451.1
2_1441385	1	2	0.59	0.00	0.00		1	1	1	1	C	Silent		e_gw1.2.451.1
2_1441919	3	1	0.00	0.00	0.00	Yes	3	3	3	1	C	Silent		e_gw1.2.451.1
2_1441939	1	2	0.89	0.00	0.00		1	1	1	1	C	P	R	e_gw1.2.451.1
2_1441973	1	2	0.89	0.00	0.00		1	1	1	1	C	Silent		e_gw1.2.451.1
2_1443263	3	2	0.00	0.00	0.49		3	3	3	1	C	Silent		e_gw1.2.451.1
2_1447270	1	2	0.89	0.00	0.00		1	1	1	3	C	N	H	estExt2_fgenesh1_pg.C_PHYCAscaffold_20222
2_1452313	3	2	0.00	0.00	0.49		3	3	3	1	C	Silent		fgenesh2_kg.PHYCAscaffold_2_#_197_#_4100895:1
2_1452392	3	2	0.00	0.00	0.59		3	3	3	1	C	Silent		fgenesh2_kg.PHYCAscaffold_2_#_197_#_4100895:1
2_1452394	3	2	0.00	0.00	0.59		3	3	3	1	C	Silent		fgenesh2_kg.PHYCAscaffold_2_#_197_#_4100895:1
2_1456984	1	2	0.58	0.00	0.00		1	1	1	3				
2_1457008	1	2	0.68	0.00	0.00		1	1	1	3				
2_1482757	3	2	0.00	0.00	0.68		3	3	3	1				
2_1482805	2	2	0.00	0.69	0.00		1	1	3	1				
2_1482817	2	1	0.41	0.00	0.00		3	3	1	1				
2_1482847	1	2	0.59	0.00	0.00		1	1	1	1				
2_1482853	1	2	0.59	0.00	0.00		1	1	1	1				
2_1482862	1	2	0.59	0.00	0.00		1	1	1	1				
2_1484820	1	2	0.58	0.00	0.00		1	1	1	3				
2_1486427	3	2	0.00	0.00	0.58		3	3	3	1	C	Silent		estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1486428	3	2	0.00	0.00	0.58		3	3	3	1	C	Silent		estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1486435	2	2	0.00	0.62	0.00		1	1	3	1	C	M	K	estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1486458	3	2	0.00	0.00	0.58		3	3	3	1	C	A	T	estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1487541	2	1	0.33	0.00	0.00		3	3	1	1	I			estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1487551	2	1	0.33	0.00	0.00		3	3	1	1	I			estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1487569	2	1	0.33	0.00	0.00		3	3	1	1	I			estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1487581	2	1	0.33	0.00	0.00		3	3	1	1	I			estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1487584	2	1	0.33	0.00	0.00		3	3	1	1	I			estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1490736	2	2	0.01	0.71	0.00		1	1	3	1	C	Silent		estExt2_Genewise1.C_PHYCAscaffold_20980

Supplementary Figure 11: Dendrogram of SNV cross correlations derived 23,809 SNVs from 2 parents and 69 putative progeny *P. capsici* isolates. Isolate names on the left side of the panel correspond to the tree branches directly across on the right side. The same tree is reproduced on the bottom of the panel with isolate TN1 on the far left to create the cross diagonal. Correlations are designated by color legend at bottom of panel.



Supplementary Table 1: Arachne hybrid assembly version11 (20 Aug 2010) of *P. capsici* LT1534 454-Titanium and Sanger gDNA reads.

Scaffold count	917
Contig count	10,760
Scaffold bases total (Mbp)	64
Contig sequence bases total (Mbp)	56
Estimated % sequence bases in gaps	12.5%
Scaffold N50 / L50 (# / kbp)	29 / 706
Contig N50 / L50 (# / kbp)	397 / 35
Number of scaffolds > 50.0 Kb	140
% in scaffolds > 50.0 Kb	91.9%
% assembly masked by repeats	14.4%
# finished cDNAs	1260
% finished cDNAs that align with assembly	97.1%

Supplementary Table 2. *Phytophthora capsici* filtered gene models classified by prediction method. Annotation v11 (20 August 2010) was produced by the JGI Annotation Pipeline, using a variety of cDNA-based, protein-based, and *ab initio* gene predictors.

Prediction method	# models
Total	19,805
protein-based	14624 (74%)
cDNA-based	2920 (15%)
<i>ab initio</i>	2261 (11%)

Supplementary Table 3. Properties of the 19,805 *P. capsici* filtered gene models, including genes with homology to transposable elements.

Property	Value
Avg. gene length	1253 nt
Avg. transcript length	1028 nt
Avg. protein length	314 aa
Gene density (per Mbp scaffold)	309
Avg. exon length	467 nt
Avg. intron length	189 nt
Avg. exon frequency	2.2 per gene
Multi-exon genes	9861 (50%)
Genes with similarity to proteins in nr	14925 (75%)
Genes in LT1534 multigene family	15489 (78%)
Genes supported by ESTs	9981 (50%)
Genes with Pfam domain	9234 (47%)
Genes with signal peptide	3162 (16%)
Genes with transmembrane domain	2890 (15%)
Genes with EC number	2821 (14%)
Genes with GO term	8544 (43%)
Complete models (with start and stop codons)	69%

Supplementary Table 4: Field isolates chosen for RAD sequencing (first 8 isolates)

	ID	Host	Year	Mating type	Location	Species
Parents of sequenced isolate	LT6503	Snap Bean	2009	A2	Connecticut	<i>P. capsici</i>
	LT62	Spaghetti Squash	1998	A2	Michigan	<i>P. capsici</i>
	LT263	Pumpkin	2004	A2	Tennessee	<i>P. capsici</i>
	LT2135	Pepper	2006	A2	Peru	<i>P. capsici</i>
Clonal lineages	LT5473	Pepper		A1	New Mexico	<i>P. capsici</i>
	LT51	Cucumber	1997	A1	Michigan	<i>P. capsici</i>
	LT6535	Pepper	2008	A1	Argentina	<i>P. capsici</i>
	LT232	Rhododendron	2004	A1	Tennessee	<i>P. tropicalis</i>
Sequenced isolate	LT1534			A2	Lab (inbred)	<i>P. capsici</i>

Supplementary Table 5: RAD sequencing statistics.

Isolate	Reads	Reads Aligned	Reads Uniquely Aligned
LT6503	8.1 M	7.0 M (86%)	6.0 M (73%)
LT62	7.8 M	6.8 M (86%)	5.8 M (74%)
LT263	8.1 M	7.2 M (88%)	6.2 M (76%)
LT2135	7.4 M	6.4 M (86%)	5.4 M (73%)
LT5473	8.4 M	7.2 M (85%)	6.1 M (72%)
LT51	8.5 M	7.4 M (87%)	6.3 M (74%)
LT6535	5.9 M	5.1 M (85%)	4.3 M (72%)
LT232	5.0 M	0.8 M (16%)	0.7 M (14%)

Supplementary Table 6: Single nucleotide variants (SNV) in RAD sequenced isolates.

Sample	Nt with \geq 30X coverage	SNVs	SNVs/kb
LT6503	1,761,096	20,445	24.88
LT62	1,714,370	17,914	22.08
LT263	1,829,566	14,637	16.87
LT2135	1,631,155	20,869	25.52
LT5473	1,760,096	21,386	26.04
LT51	1,903,934	18,527	22.30
LT6535	1,597,281	22,431	25.88
LT232			51.67

SNV detection required at least 30X coverage of that nucleotide in all samples and at least 4 uniquely aligning reads with Q scores of at least 20 to call the variants. The *P. tropicalis* (LT232)/*P. capsici* SNV/kb ratios were identical at both 10X and 30X coverage and in comparisons of all samples. These SNV densities are conservative because the regions had to contain the 6 nt invariant endonuclease site and sufficient reads to align at 90% identity in order to meet the 30X cutoff. Densities were increased to ~26/kb when the coverage requirement was decreased to 10X, and to ~33/kb when the restriction for 30X coverage in the *P. tropicalis* isolate was waived.

Supplementary Table 7: Distribution of SNVs among seven *P. capsici* isolates and one *P. tropicalis* isolate. Bi-allelic SNVs = 59,471; tri-allelic SNVs = 2,413; tetra allelic SNVs = 80; total SNVs = 61,964

# isolates with a given SNP	Number of SNPs
1	34,065
2	8,447
3	5,470
4	4,265
5	3,479
6	3,243
7	4,309
8	1,259

Supplementary Table 8. Summary statistics of RAD sequences generated for the parents and 69 putative progeny.

	Parents and Progeny	Parents	Progeny
Total	158,460,781	9,588,155	148,872,626
Mean	2,231,842	4,794,078	2,157,574
Median	1,655,467	4,794,078	1,646,382
Std. Dev.	1,602,718	138,864	1,563,741
Min.	594,851	4,695,886	594,851
Max.	8,951,606	4,892,269	8,951,606

Supplementary Table 9. Summary of the markers for each of the linkage groups. The parental isolates differed at ~25% of the 20,568 loci. TN1 had 5273 RR, 8632 AR, and 6663 AA genotypes, whereas TN2 had 8141 RR, 11767 AR, and 660 AA genotypes (where R = reference allele and A = alternate allele). The smaller proportion of AA genotypes in TN2 reflected its use in backcrosses that produced LT1534, the reference genome.

Linkage Group	SNPs	RR:AR	AR:AR	AR:AA	AA:RR
1	3003	1576	453	519	455
2	1277	625	275	264	113
3	1324	702	204	253	165
4	532	269	139	98	26
5	1621	807	264	333	217
6	839	448	122	156	113
7	448	231	51	92	74
8	1493	865	210	274	144
9	925	465	161	159	140
10	2940	1330	447	684	479
11	1412	704	138	219	351
12	871	386	108	279	98
13	933	493	94	198	148
14	547	217	50	63	217
15	687	319	242	104	22
16	1231	493	43	293	402
17	371	134	80	63	94
18	114	78	22	0	14
Total	20568	10142	3103	4051	3272

Supplementary Table 10. Summary of genic and non-genic markers showing simple Mendelian inheritance.

Linkage Group	Coding					Non-Gene	Genes Per Linkage Group	Total Genes in Linkage Groups
	Coding Silent	Non-Silent	Intron	3'	5'			
1	858	488	382	215	48	1012	716	2,271
2	384	218	107	82	32	454	309	1,016
3	392	187	160	96	27	462	382	1,146
4	176	89	26	24	14	203	140	648
5	450	286	170	97	25	593	371	1,392
6	274	144	43	37	17	324	152	646
7	111	66	44	21	4	202	91	467
8	427	238	114	77	18	619	365	1,087
9	277	157	96	80	16	299	242	1,076
10	800	492	355	196	47	1050	645	1,941
11	378	244	124	75	30	561	357	1,267
12	249	141	51	64	13	353	197	870
13	285	147	57	47	16	381	190	778
14	164	100	36	26	15	206	203	570
15	207	104	63	42	6	265	178	655
16	322	204	130	100	25	450	288	1,194
17	95	62	23	27	6	158	157	546
18	23	15	12	4	1	59	34	119
Total	5872	3382	1993	1310	360	7651	5017	17,689

Supplementary Table 11. Summary for linkage of scaffolds breaking into blocks with LOD scores ≥ 7 .

Scaffold Block	Lowest Marker	Highest Marker	Total Covered	Linkage Group
Sc1.1	1_53205	1_945508	892303	LG9
Sc1.2	1_1041839	1_1173436	131597	LG2
Sc1.3	1_1283380	1_2077147	793767	LG11
Sc2.1	2_113526	2_546537	433011	LG13
Sc2.2	2_731366	2_1916562	1185196	LG10
Sc4.1	4_87661	4_936084	848423	LG10
Sc4.2	4_953874	4_1526218	572344	LG12
Sc7.1	7_95189	7_945588	850399	LG13
Sc7.2	7_961256	7_1255438	294182	LG1
Sc12.1	12_27274	12_214345	187071	LG18
Sc12.2	12_711003	12_1052066	341063	LG14
Sc13.1	13_112552	13_176739	64187	LG11
Sc13.2	13_290642	13_1028147	737505	LG2
Sc14.1	14_454507	14_856026	401519	LG14
Sc14.2	14_893013	14_991161	98148	LG11
Sc18.1	18_98284	18_475978	377694	LG11
Sc18.2	18_597329	18_965332	368003	LG2
Sc21.1	21_96574	21_533490	436916	LG2
Sc21.2	21_630881	21_910293	279412	LG5
Sc22.1	22_56130	22_491901	435771	LG3
Sc22.2	22_536539	22_839077	302538	LG08
Sc23.1	23_45180	23_357600	312420	LG1
Sc23.2	23_653052	23_764546	111494	LG2
Sc24.1	24_9834	24_190368	180534	LG7
Sc24.2	24_227914	24_769629	541715	LG5
Sc25.1	25_17541	25_427916	410375	LG15
Sc25.2	25_657899	25_803448	145549	LG6
Sc26.1	26_63597	26_631721	568124	LG8
Sc26.2	26_641731	26_706136	64405	LG1
Sc42.1	42_26570	42_320442	293872	LG14
Sc42.2	42_349280	42_479514	130234	LG17
Sc48.1	48_22671	48_192342	169671	LG10
Sc48.2	48_212975	48_350345	137370	LG3

Supplementary Table 12. Summary data for linkage groups and scaffold blocks.

LG/order	Scaffold Block	Scaffold Block Size	LG Total Nts
LG01.01	Sc43	531,961	
LG01.02	Sc23.1	312,420	
LG01.03	Sc82	207,552	
LG01.04	Sc107	135,171	
LG01.05	Sc3	1,611,504	
LG01.06	Sc70	280,435	
LG01.07	Sc7.2	294,182	
LG01.08	Sc9	1,114,135	
LG01.09	Sc41	565,801	
LG01.10	Sc109	125,037	
LG01.11	Sc26.2	64,405	
LG01.12	Sc105	102,149	
LG01.13	Sc17	985,188	6,329,940
LG02.01	Sc13.2	737,505	
LG02.02	Sc23.2	111,494	
LG02.03	Sc172	15,116	
LG02.04	Sc11	1,076,090	
LG02.05	Sc18.2	368003	
LG02.06	Sc1.2	131,597	
LG02.07	Sc21.1	436,916	2,876,721
LG03.01	Sc48.2	137,370	
LG03.02	Sc95	141,949	
LG03.03	Sc30	669,438	
LG03.04	Sc65	307,771	
LG03.05	Sc37	577,888	
LG03.06	Sc51	529,628	
LG03.07	Sc22.1	435,771	
LG03.08	Sc64	301,055	
LG03.09	Sc102	120,866	
LG03.10	Sc560	6,948	
LG03.11	Sc800	3,230	3,231,914
LG04.01	Sc6	1,423,605	
LG04.02	Sc86	162,230	
LG04.03	Sc44	536,020	2,121,855
LG05.01	Sc10	1,190,366	
LG05.02	Sc63	355,987	
LG05.03	Sc68	334,182	
LG05.04	Sc24.2	541,715	
LG05.05	Sc50	475,612	
LG05.06	Sc21.2	279,412	
LG05.07	Sc52	444,141	
LG05.08	Sc80	185,445	

LG05.09	Sc60	366,873	
LG05.10	Sc67	292,061	
LG05.11	Sc112	78,237	4,544,031
LG06.01	Sc87	182,599	
LG06.02	Sc118	67,328	
LG06.03	Sc139	29,350	
LG06.04	Sc191	15,330	
LG06.05	Sc93	150,113	
LG06.06	Sc108	116,431	
LG06.07	Sc25.2	145,549	
LG06.08	Sc89	163,789	
LG06.09	Sc81	202,973	
LG06.10	Sc132	34,092	
LG06.11	Sc39	591,971	
LG06.12	Sc92	170,487	
LG06.13	Sc74	127,009	1,997,021
LG07.01	Sc24.1	180,534	
LG07.02	Sc106	144,267	
LG07.03	Sc77	232,015	
LG07.04	Sc16	1,020,712	1,577,528
LG08.01	Sc26.1	568,124	
LG08.02	Sc8	1,194,346	
LG08.03	Sc161	37,761	
LG08.04	Sc22.2	302,538	
LG08.05	Sc58	327,658	
LG08.06	Sc72	230,452	
LG08.07	Sc66	294,895	
LG08.08	Sc62	322,952	3,278,726
LG09.01	Sc5	1,593,284	
LG09.02	Sc100	129,988	
LG09.03	Sc103	136,111	
LG09.04	Sc1.1	892,303	
LG09.05	Sc90	152,720	2,904,406
LG10.01	Sc34	622,686	
LG10.02	Sc40	570,383	
LG10.03	Sc4.1	848,423	
LG10.04	Sc27	705,730	
LG10.05	Sc2.2	1,185,196	
LG10.06	Sc84	182,275	
LG10.07	Sc49	508,508	
LG10.08	Sc48.1	169,671	
LG10.09	Sc28	757,753	
LG10.10	Sc79	222,129	5,772,754
LG11.01	Sc1.3	793,767	
LG11.02	Sc32	608,388	
LG11.03	Sc97	154,197	

LG11.04	Sc18.1	377,694	
LG11.05	Sc29	707,326	
LG11.06	Sc31	686,815	
LG11.07	Sc14.2	98,148	
LG11.08	Sc13.1	64,187	3,490,522
LG12.01	Sc71	252,329	
LG12.02	Sc38	550,336	
LG12.03	Sc94	190,298	
LG12.04	Sc53	465,517	
LG12.05	Sc91	167,065	
LG12.06	Sc4.2	572,344	
LG12.07	Sc57	397,303	2,595,192
LG13.01	Sc36	679,425	
LG13.02	Sc111	91,577	
LG13.03	Sc2.1	1,925,921	
LG13.04	Sc61	347,729	
LG13.05	Sc7.1	850,399	3,895,051
LG14.01	Sc12.2	341,063	
LG14.02	Sc14.1	401,519	
LG14.03	Sc42.1	293,872	
LG14.04	Sc101	136,107	1,172,561
LG15.01	Sc15	1,032,281	
LG15.02	Sc25.1	410,375	
LG15.03	Sc46	525,601	1,968,257
LG16.01	Sc20	899,172	
LG16.02	Sc19	944,885	
LG16.03	Sc33	640,966	
LG16.04	Sc35	603,175	
LG16.05	Sc55	463,907	3,552,105
LG17.01	Sc104	135,144	
LG17.02	Sc42.2	130234	
LG17.03	Sc47	556,112	
LG17.04	Sc54	427,428	
LG17.05	Sc85	206,069	
LG17.06	Sc73	348,431	1,803,418
LG18.01	Sc12.1	187,071	
LG18.02	Sc78	194,772	381,843
Totals	125		53709626

Supplementary Table 13. Flanking markers and tract lengths for isolates with loss of heterozygosity.

Linkage Group	Isolate	1st LOH	Last LOH	Tract Length
LG01	TN37	3_1774	3_901039	899265
LG01	TN37	3_956368	3_1023899	67531
LG01	TN60	3_1032846	3_1165737	132891
LG01	TN65	3_1032846	3_1165737	132891
LG01	TN37	7_960715	7_1232128	271413
LG01	TN37	9_76547	9_313726	237179
LG01	TN37	17_31860	17_976370	944510
LG01	TN47	17_29488	17_870195	840707
LG01	TN37	23_32662	23_422316	389654
LG01	TN37	26_642307	26_711717	69410
LG01	TN37	41_23637	41_340818	317181
LG01	TN37	43_20676	43_531378	510702
LG01	TN37	70_203860	70_279147	75287
LG01	TN37	82_27984	82_206524	178540
LG01	TN37	105_6822	105_76260	69438
LG01	TN47	105_51382	105_63734	12352
LG01	TN37	107_2162	107_50059	47897
LG01	TN37	109_25479	109_78466	52987
LG03	TN15	30_22501	30_610494	587993
LG03	TN31	30_22501	30_610494	587993
LG03	TN15	48_309043	48_463219	154176
LG03	TN31	48_203358	48_463219	259861
LG03	TN31	65_43166	65_189594	146428
LG03	TN15	95_68974	95_100010	31036
LG03	TN31	95_11127	95_120313	109186
LG04	TN25	6_517600	6_1330434	812834
LG06	TN45	39_45571	39_218172	172601
LG06	TN45	39_269905	39_467526	197621
LG06	TN45	74_57229	74_92332	35103
LG06	TN45	81_83924	81_172214	88290
LG06	TN37	87_72040	87_152869	80829
LG06	TN45	89_41061	89_116125	75064
LG06	TN45	92_10433	92_128655	118222
LG06	TN45	132_10333	132_22914	12581
LG07	TN60	16_155019	16_945332	790313
LG07	TN65	16_134053	16_974846	840793
LG08	TN3	8_159857	8_160156	299
LG08	TN47	8_148811	8_1163217	1014406
LG08	TN13	26_91745	26_340248	248503
LG08	TN3	26_142703	26_609109	466406

LG08	TN47	26_63564	26_620971	557407
LG08	TN37	66_126127	66_190579	64452
LG09	TN50	1_87441	1_742969	655528
LG09	TN50	90_44554	90_110297	65743
LG10	TN18	2_734379	2_1650759	916380
LG10	TN34	2_734379	2_1555552	821173
LG10	TN47	2_932367	2_1752644	820277
LG10	TN53	2_734379	2_1062403	328024
LG10	TN63	2_734379	2_1650759	916380
LG10	TN18	4_86143	4_936679	850536
LG10	TN34	4_86143	4_936679	850536
LG10	TN47	4_42590	4_936704	894114
LG10	TN52	4_86143	4_936679	850536
LG10	TN53	4_86143	4_936679	850536
LG10	TN63	4_86143	4_936679	850536
LG10	TN18	27_8619	27_697509	688890
LG10	TN34	27_8619	27_697509	688890
LG10	TN47	27_1128	27_697509	696381
LG10	TN52	27_8619	27_297350	288731
LG10	TN53	27_8619	27_697509	688890
LG10	TN63	27_8619	27_697509	688890
LG10	TN18	28_38583	28_704354	665771
LG10	TN18	34_47706	34_548650	500944
LG10	TN34	34_47706	34_548650	500944
LG10	TN47	34_27259	34_571709	544450
LG10	TN52	34_47706	34_548650	500944
LG10	TN53	34_47706	34_548650	500944
LG10	TN63	34_47706	34_548650	500944
LG10	TN18	40_53500	40_510721	457221
LG10	TN34	40_53500	40_510721	457221
LG10	TN47	40_53500	40_510721	457221
LG10	TN52	40_53500	40_510721	457221
LG10	TN53	40_53500	40_510721	457221
LG10	TN63	40_53500	40_510721	457221
LG10	TN18	48_66747	48_185568	118821
LG10	TN18	49_163203	49_500583	337380
LG10	TN18	79_90923	79_155337	64414
LG10	TN18	84_57519	84_178288	120769
LG10	TN63	84_57519	84_177335	119816
LG11	TN54	1_1282528	1_2051166	768638
LG11	TN54	18_96129	18_475984	379855
LG11	TN62	31_358469	31_633967	275498
LG11	TN54	32_50547	32_548375	497828
LG11	TN54	97_62889	97_123104	60215
LG12	TN64	4_1416514	4_1526252	109738

LG12	TN64	53_92504	53_418139	325635
LG12	TN64	57_52048	57_123257	71209
LG12	TN64	91_44469	91_134135	89666
LG13	TN63	7_132139	7_207249	75110
LG13	TN63	7_346854	7_427074	80220
LG13	TN66	36_95503	36_626762	531259
LG15	TN17	15_476066	15_890139	414073
LG15	TN44	15_476066	15_838906	362840
LG16	TN25	19_412214	19_765034	352820
LG16	TN37	19_412214	19_765034	352820
LG16	TN70	19_412214	19_765034	352820
LG16	TN25	33_58644	33_570713	512069
LG16	TN37	33_58644	33_570713	512069
LG16	TN58	33_508481	33_570693	62212
LG16	TN70	33_58644	33_570713	512069
LG16	TN23	35_141346	35_556998	415652
LG16	TN25	35_141346	35_556998	415652
LG16	TN37	35_141346	35_556998	415652
LG16	TN58	35_141346	35_556998	415652
LG16	TN70	35_141346	35_556998	415652
LG16	TN23	55_62577	55_381570	318993
LG16	TN25	55_25260	55_381570	356310
LG17	TN47	85_67570	73_215288	147718

Supplementary Table 14. Summary of total nucleotides affected by LOH per isolate.

Isolate	Total LOH Coverage
TN13	248,503
TN62	275,498
TN44	362,840
TN17	414,073
TN03	466,705
TN58	477,864
TN66	531,259
TN64	596,248
TN45	699,482
TN50	721,271
TN23	734,645
TN15	773,205
TN60	923,204
TN65	973,684
TN31	1,103,468
TN70	1,280,541
TN54	1,706,536
TN52	2,097,432
TN25	2,449,685
TN53	2,825,615
TN34	3,318,764
TN63	3,689,117
TN18	4,721,126
TN37	5,556,816
TN47	5,985,033

Supplementary Table 15. Source and version of genomes used for CEGMA analysis.

Organism	Source	Genome version	Reference
<i>Phytophthora ramorum</i>	http://genome.jgi.doe.gov/Phyra1_1/Phyra1_1.download.ftp.html	1.0	Tyler et al. (2006)
<i>Phytophthora sojae</i>	http://genome.jgi-psf.org/Physo3/Physo3.home.html	3.0	Tyler et al. (2006)
<i>Phytophthora capsici</i>	http://genome.jgi-psf.org/PhycaF7/PhycaF7.home.html	11.0	
<i>Phytophthora infestans</i>	http://www.broadinstitute.org/annotation/genome/phytophthora_infestans/MultiDownloads.html	4.1	Haas et al. (2009)
<i>Pythium ultimum</i>	http://pythium.plantbiology.msu.edu/download.html	Release 1	Levesque et al. (2010)
<i>Hyaloperonospora arabidopsis</i>	http://vmd.vbi.vt.edu/download/index.php	8.3.2	Baxter et al. (2010)

Supplementary Table 16. Predicted Crinkler amino acid sequences and gene names.

Gene Name	Amino Acid sequence
>jgil104199le_g w1.9.417.1	MVKLFCAIVGEAGSAFSVEVDETDSVDDLKTAIKAVNEDITCPPRKLQLFLAKKADGT WLDGAGAAAVTVDEADRVPLMLDKGHNHHKFVKMNPLLWIKNDQHFGENFRPVHV VVVVPDVAHAQTGLWLTGFVKNALNTKGIRCKLYWMATLRIGYYDPARRTDKKNV AFWYEDTKLCFHVLFETKDAALLFETDLRIEPQLGSPLTNQVVERAPVNAVTELQ RVFYGDYVPDDSKSPQNSVSSISLTSGSNLDSSTDEFRFQRIEHEKFFLPYGKAESCHLV SRKQSRNHKREFAKYDRDSNSRLALS RDHMHGWF DMSIEVPIVNMLPGSVKENQSIGN RRKVVEVFVKVLDAGCTDRVFSRLKEGSTTNDPLMMKTFVHVEDPETFCLCMRWKH DDNAERWRSFWDMTPAVD*
>jgil106435le_g w1.12.217.1	MLKLFCVVVGAGDAFPVNIEPNETVGDMKKIKHEEMYQFPASELQLFLAKVPKEKH DMAWLSSRSEDVKKLKGEKTPLIDILTEEDQELQAEDPLDDVLRGMPPSLCQIHVLV MAPPQDSLRSHLTTLLSVLLCHVLTKAPPTPTDRNVDFKDDVCNFYGCYSPDESCVRC MLLNDAPSELVVASHLFRCSNEDVSDVMMQITLSIDDERNLLFKPLKYAFDHFQI SFIRDDTDVFRWKVFDPSILATPIVDLKDRKGKVLSTEQTQLLSRISENPCRFTNTQTF GDVDDSALTFTGLERPFYRCLNLQARVARVMALEKKWIDASYDFQDFWSEVSLDDKM EMFHRSILNS*
>jgil106457le_g w1.12.537.1	MKLFCAMVGMAVGIEVDIDNNAYVTALRDAIATKNEEIKKSALRLKLFACKGNAWL TDTEAAGVGGDLES LGFKL MKSV RLLK NPEY FGED F QTGE GHV H VV PEE M TVG EPVVDV DG VNI YV TS NMT LNP DLV AF WRAF QAI DT KIE AD S VIAL PEGT FIL GNP KV GS RIYIRPCYPQLWEVCWHIIHHETPNL VILGNPGIGKTYFGYLLLFLARLGKTVVYESRR TKRRFLFSRNVVVIKGSSQDFDDILEQD TYYVVDAMEPREFQARTILVTSPDRDVWYTF NKISCQTRYMPVWTEQEIFSCREQVYSTIPKSVVQKCFYRGGIPRYVLQYAQFDNHQ ALIEKALEVVDFDWLMNAYGK LDDNNSQAHRLHYRVNERFTCDYFGFASSFVQHEV YQHLHKKEKRKLLEFIGRSVGSGDLSVLRDRLAEEHDHHCAR THKKL KRF*
>jgil106668le_g w1.12.123.1	MVKLFCAIVGVEGSVFSV VIGEGQTVEELKKAIKERNDDKINVSWLGLQLFLAKKAKG DGDWLTEKDQVQEGVYDMSDLQMLRAARAKLRLVKSNDVNEEQKVEGGKSVNVL VVLPGTTTIKVNERENNALTDELAYYQRIGQEIQSNCQQHC GPILD KIDS IYEKKPYPM PFICVQGSSGMGKSQLA FGEGREHPRPW FYWTHGT VSDYDQ RIYRNFA SIGSAF DS VVKDEV RKEE DDI LNCTS VLYMTK KLWTYGF IELL RYCS RS NVGAQM VRV ENQ TF YVTKSNLEDVIEVRSRMEKNGEVLPFFI DEMPPSRTK KLSAFQLNVFRACGLV VIG MG TDANISNLV GKPEHSRTDPHWWMTV VSCFPW QSIPFGDPAKEEVWQKVIELHPV VKH IAEH SRG LFSRC FVDAV VKF AMEEVSEN ETF ALAD LLDA AFK A VY A ELR ITKG FMF TEE GRDAQML ALSYTIGS NKPPGT KRP VT DSD STSEPTPK RRK LDV DVGV ASM RAHF AN SG YE GIAD VD VLQGDLR FRD TSD AW SPIC QF PAIE KDV LLYLA V LG KK FSDY NDT SSTD VFE FLER NGN REES NAL RRD FTN FIK FEN VVA HAL FCAS RR NG ARGIAL YE FLS GLV SE FQ DE YY QR DEF DAS ALL KE FD GL KEK FA EKK PFL APPN ARW PDY ILE AGG DCN FGH FE RMDGYVMVPGV DGPLFACDCQYWKDDLDSDAMKKIIAGLNGGATGCGDDQQPKW SNWSLALVFCRKLEEF ESEQREDWEFPSTGIATVDCKAWEVN WISKPEAGEKL VIVV QT GRVLCLP*
>jgil106736le_g w1.12.192.1	MMKLFCAMVGMAVGIEVDIDNNAYVTALRDAIATKNEEIKKSALRLKLFACKGNNG RVEWL TQL DVVKGVMDANGFTHLQFADAKLRAVGLKSSELVEVNEEDVAVGKGHVH VLVKIDQTAPNVELMPFTSTCWLVTS GVNALNTRGVRGQLYR LAHTELGYYDPANV RTDTGNIPRAFWHENNDIQI HVLF KEEHALYFQSHLVDDFSIPIKSFVSKACP TDLQRI FRDHYVADET VSPQLSMSF SSEISVFDPSNPVFKYQRIAERLFGSHGKAES AHLISASHCR NVT SYDEYDKDDNNRLALSREM HGAYDG INCDFPLVNIEVVSASDHPELDHR FKVELQ VSVY SHEYV FLLGRLKDG STKT EDPL VMKTFVYVQDKNIFCTCIQWKYNKNRQLREEF FTTNPRAT*
>jgil108643le_g w1.15.666.1	MAKLSLSCVIVGLMGSAFSVDIDADQFVGDLKKIIQKKNDLHNVDA DKLQLFLAKKD KGWLPDKSEA AALELK KGEV HEDIQV LINGE EMEAT KTL NYWL FEKNQ MEEQ LSSEQ IH VL VV VPEIR DGKK RY RSSL WFAE SEPL KRRV KT DESE DED DDV SEERT FFQ LCG CPM AHPARKDSKLMERKAYTVIFAQLVEHV KDC FEFNRTSNNPGLSSNVV TGNPGIGK SW

	FYLYCIFQLIRNREREDIKQLPPYELVMNYDDNFVKYDAARAECVRLNKEDVDDLMKD PFVLRVDARSTKLMGWRGVSVLFAPDAEDLHDFEKVPGPKFIMPWAWSLEELQDCNQ VLDDLKLAEDELVSFRDAFGGIPRYVFSKNKTAIENKLKRAMASFSVKEILSYCKRGA AVKESDQSDCVLQMVSEANFRLFYLDLSSDICEKIVFQAEGEDLTMLAKFAMGK*
>jgil110274le_g w1.18.174.1	MVKFFCAIIGVAGSAFPVIDASLSVGDLKNAIKGKNDIKCTARELQLFLAKKEKGAG AWLTEDEAAAVSLDEGGHLQGFKMMKSSLKKNPKHFGSNFQSDEDQVHVLVVVPD QAQPQTGLWLVTSVENALNTKGIRCRLYRLMASYLGYDPVRRTGDKDTALWYED KTLCIHSIFKSEENALLFDNVLQDECITQTSPLDGHDVSTNVAPVSRQLSELRIYSRHYA PQDTESPQVMSLISINTSIVDVTMDEFKYZQRIESEWFGDVGKAQSCHLMSRDHCRKC PSDRKYDNDPNNRLALSSAMHDWYDGRMYNPVMNISVESVSEPVIGNRYKVNLIV RALNARYAKWISLILKEGFVASEDSLEMHTCVYVQNPKVFCVCMEWKRKEIDKQWK YYDMEPAVD*
>jgil112192le_g w1.21.161.1	MVKLFCAIVGAMESAISVEVGEGLTVGDLKKAIATDQKFDAASKLQLFLTEEGGGW LSSEDDAAIAMRTGAPEQVKLLLKDEIDPAEEIGDKFGSAPTKTIHVLVVVPEGVTNV TAATVFENNKRREDDQPDFWMKAIEDDKVLTPLTCEELKEHLQRELPVKIPLGGRL SQIVESQNVAGKMNSKLFDEARHSVTDTVAILNGVIAPVWIGGESPTEATYHFLWNE VIAKVLMYVSDGTCSRNSSFVFTSTGLFRPNLCFYNSANDKVCVFRGEEQARGEMTVPL TDLQRLTWRYDDAPYVFGYAAAQDVCLVTIRESETNAPGEKRRAKVEIERYNLRDL RGRLSIFLALLNLSTLFRPVVGRIRPLGIAEYKTIIRPNGVKIAFGENCVKTYPLTMP KIISDLRDLHWQMKENAVPNVVELKSTNMRKRSVELAPVGRQLPLENVHQLLMAMRD ILRALVALHTIGLMHRDLRWENVLRYPDEDKWFLIDFDEGASSPAVKVDHLKAETHAP EILSSSTHTTMVDIWSVGYLLETSHVHDLVALEDIKTQCLQENPSVRPTAQSLLEAVEA LIAN*
>jgil113302le_g w1.24.271.1	MVFASQDNDFNRCSPFFSQLPTAEETDEWLEFPSLLPLTRRSLYIRPSFKSIAAQALLK VDSNRRKYAVVTGTPGIGKSVFLYYVMWKLIKDKRMLFVTGRPPIYFDGESVLDYCQ LPYAGNRNFWSPLWLVDATNPCKIAGLPIHHCSVLLASEPRDDYVRHFRKLVPTPQV FYMPIWTEEEIMEKIPLYPSAASVWRDRFETLGGIPRLVLQAVQTDPQEFMRCVCFYSL CMRLVLFHSKTKTGFIDSLYTLEMPRHVHILSQEPYHEYTLAYASETAMRAVIDAKWIV NRAEMLHFLVMNLKSTDLSLTQTTCHCIFELYAMQLELGGSFSYRSLQAGVEQSSETLD EDENDIDIPMSWREIVDRVEADQNEDQLYVPKSAKDVAIDAWMPEVGGFQIALGKEQK IKSQADELALLGQGGNRLFFLFVPRDFDSFTKQEPLSIEQYALLIPYPEV*
>jgil114043le_g w1.25.260.1	MVMVKIFCKIVGEAGSIFSVKIDDSESVEDFMRKIKDRCDGKIVAPWMDLQLFLAKKKD GVWLTKRDVLEVTSELDPLDDTLAPLNSVGLSEEDVRYRMTKEDIKAKKVPVHLLVV PTEDYLRSPATILLETILPHVLTHATTTLTEDNRDFRHNLNCFYGCYTREQSLVRCMLLD VPLPKSLVLA SHLFRRSNEYLSFRMMQISDIDDVKNGLLLFKPLKYAFDHFQISFIRDDT DVFRKLKF DSTIKETPLIDLVDHHGKKVLSEEQTGELLSDVNNDTCLFDVGKTFGDVG CALAFTGIERPYYHCLNLQARVALMVALKKWIDESYDFKDFWSEVSLDDKMEMFHR SILNSVADI*
>jgil117630le_g w1.33.284.1	MIKLFCAIVGEERNVIEVDISDVESISALKEAIKTKMPITLKVDANTLQLPAKMPDDK WLRDDSAADDLMSGRIHDDIKAMIADSKILRPACRIRDELSAKKTGWKWSLPSPITP KIKTKQIHVLVKVPDMMTGTLPLCGVAPEAQWRWLWQSIVRTSGTYSGTALVLRSSR NLYLLTNLHFWLAEYEEEIFFEHMSAGFKMNVELYLRKNPRKRGKQKSDGGELQQ TFNGSPVVIVDQLLPGHTTLTEVHSFVFKSDACWCSSVDYDYAVFKVLAPPRIQLGV EPSVSHFPTNVYVFGFHDGHEEKKFGHSYAIVPARIKYHRYKCLILSSTVSLPESGVICT SRGLSIGYLAGSTIDESGNEVNQWLELGFFRRIL*
>jgil128631le_g w1.77.29.1	MVKLFCAIVGDPWSFEVDIDENVSVSELNKAIKTKNKKIKLKVDAKTDGP SGGKVWLDRAGAAA VETDDLHFRTRMDSTLYVRNPKHFGSGFKPDEGQVHVLVVVP K WAVEAETIGQNVERVGLPRTTALNDPKKYAEIIALDDWGVTNHQIPSIWEFMSL GCTKTGEFLWRLEEKQVASLLDGFWRESSPGSINQFEDMKSI LMGSPGIGKSTLLCVM AFHLVLYKKNLVYRQLKGENCLLYLG YEDDEVYFTVKRCKADRAVSIYEELGYR QGFPNVWLLDGFRYKEIPEGLETFRMLATSQQVSLKSQESTDAYCCLLPCWSKDLLS MGSLIYNFTPDEMEERFFYSGGSVREFTYATWEDIQRAMDVAVSGVEDYSKLLTASC MFTDTSQVVRTFVENTNDRSHYFSSRYWEPMDSEYAVLALSVRLKADALHRYTWA KMA GHGSLAGCAFEIYLHRLAIDNRLE
>jgil129711le_g	MMKLFCAVVGEAESVFVDIEPGETVSDLKDAIKEINKHDPVLKNVTAMNLQLFLAKK

w1.86.23.1	GGAWLSGDDPAVLELEESEIHPDILEMMDAKPMLDDKTLQFLLFEKNKLPQPSTNQIHVLVEIPAGARAIPYSNRPPKPFVSSEGATWDFQNPLDNEQLSNAIRKHYDAWKRGYYDKILHPLFTCWSPGTGKSRLLDEFPKLLKDWLFLAGKSENPMDMIRLLQNAFTFNIAFDKETPHEAGSFSSAAELIGTRMLYQLQDTLKWDVFVQEKSRSVPSDVMDKLSKILGTRHKDMCVILCVDGMEKLSHENGKDCEFYKVLTVLSYLIGTSKWCWVIAICSAIYSPVKNFLLSSPQWAYEVPTAILSRPTVEGEDIFATFNGDQLIELLIDDMGGFRALEVHVMMRKARRKGSLEFMSVLTAVLAELRVLYPRIKKMASMQEAFALVVARRPVDKYSRGKLSLDDVI STGLVRREGRFLTCPYVLYLLLDPDSPWSKYCYSSQETRENAKPWTWEAFNYKLRALKSVACQGKVDWRDIHRGARFGRCYRVVIEEPRYSLDVNRKTAKLDFGEGNIFRCKYDPDQQGYFFEDAFTGVKDAESRAFHIEHQCKKIKDNLSLEDLLEEKKAAGPHDLFLYCTSEVEGDIESLENCAVVDRTCWEKYYGPFAARALYVSTVSPPDINTSAIQSQLQVNIGIGPAISKRIVEKRPySSLEEAHEKTGVSMNILSQTSCKRSKVDK*
>jgil131973le_g w1.125.9.1	MKLFYVIVGVAGSPFPVDIEPSETVGDLTKAIKKENKCRFKHVDAYDLQLYLAKKDKNGA WLTEDDVATVRDDAVFQTYSKLMKPTLFLNNTEICGESINQCDVHVLVKAPMRLPTIHQLHYRHFTVGSVDIRTKKSMTFDPPLVRFWRALQGDRTEFKADAVLTPEGTFLLGNPMLGSRIYIRHCYPRWLQVCLKMINDEAMNTPHLVILGNPGIGKTYFGYVILWLLARS GNTVVYESRVCHRRFLFSQDMVVQGSKKDFIEILEQTYYVVDGVEPRYYSAKTILLTSPQREVWYEFNKDDCRSCYMPVWSRDEVLTCRELMSDIPESVQDCFRRWGGIPRYVLHYATAGGRQQWLEKAMENITLDSLMDACGDLYENPSEESLHLLHYRTKEFNTDYDFDFASQYVLEEVYRRLYNHNKKLLEFIDKSGWVGAAAVLRDHLFEVYTVAAASLTDEDGDGVAVEQFKANKLL*
>jgil20879lfgene sh1_pg.PHYCA scaffold_75_#_8	MVKLFCAVVGVGQGSAPVDidASQSVGDLKDAIKTKNKKLKNIDASDLQLFLAKPKDGPWLRSDDSDVIRMRSGAIPEQVKLLNEQIDPAAGIGALFGDAKPTMEIHLVLRVPDYDSDSEVNQQRKLTFSQKLRKESGATGELPVQGDFMFKLFDLTD DIGVQLVDIKAIGDIVGFTGSEFYIRKEILTSQWSGSARLLVAMIAQDNPWNKNIMGELGINCTQLSLNNNDTKVSTTVDAQFIAARNNQLQVLPDSD*
>jgil39322lgw1. 83.18.1	MVELSLQCAIVGQIGRSFDVEIDDGKKVSKLKEMIQVKNRETIKCDAKDLRLFLAKMGPPTTQLLKIWKKESTDGNSPFSFIPEFGEMVKLFYLNMMFIQEEQMELPSTDQIHVLVVVPPPVGSKRSADIEVAKVLKRLKMIEPKVLTEIPLPVEKEFQLDNLSMVQQSDDPIVMTPTLHEFWKKFGEFPLYYFVRMEEVFWKVIKKLLFGEDRVVIVGSPGVGKSCFLMLLAFLYACIKRKVVLVIRRLK
>jgil506739lfgen esh2_kg.PHYC Ascaffold_21_# _111_#_Contig6 65.1	MVLVALTCALVGKTGVLGVKIDDSAQVWELKKAITVEKSNDLKDVADKLEFLAKMMEGGTWLDGAGAAAVALDERGHPQGCVQMDPTLWIKNSKHGDNFKPGEQVHVLVVVVPKDENDRSAAMALGVPSLPPPTFHRHPERLKRWAAINEMIRQKNQDGNEKTGTRDTNKKRKNRDIDKSMPYSSLWNLDLEPILSVEDFDLKASA VPPNVVEALRDRLMQVRKLYGDVYSGKEAKRQVFIVAIIEAVCLMLGDATILVEEEVKGNVLVHGRFEVLRKGKRNVSIVEAKRDDIPQGIAQNVAGLEALSDVEGLERTLGIVTNYLEWWFISDDDEKIRRMNTTLKVGAVPSIQLTEIVGMIYGFLANSS*
>jgil511381lfgen esh2_kg.PHYC Ascaffold_83_# _9_#_Contig727 .1	MVKLSLQCAVVDQAGSSFDVEIDDSAKVSKLKKVIKEENPATITCDAKDLQLFLAKKDDAWLDGAGAAAVALDEHGHGPQGCVQMDPTLWVKNPKHGDNFKPGEQVHVLVVVPEGVVGASSETS KMDFVVDKVS KLYEH S VLSKRTRYVHSEMSSSKGNKLV KELKIRVT PVD AVPFTGGSPTPVEEFEWIKRTEEQQSGRYRDYVEANIGDVL RNNKLCVFSVEKGANILSVEVPGCDVDLAGRTDMIVLSAIVQKFPHLPHLPGVKMLIEVKREVKSASEFQALS ELIAMDFIVDESVMALLTNLNTNHWEFLWVSNKSNNRPIATTLLTPGEAFEVIRTLAQ SSTADADIMLPCLAEPVKRRKLNQMLPFIGEASGDGIRESIERYDIASCLGPDFDMARA VARQVTRSIPTLSYFS*
>jgil527369lestE xt2_fgenesh1_p m.C_PHYCasca ffold_180095	MELRCGVYGEGSVFSVKIARDVKVSALQKKIASVLSTEQHTVFPRLLA LY LARKKEGEEVKWLKDDRHA KDFL RG GTITEYEEMRPSWTL DDEELFGPDFQPGQEEIHVLVELPKAA VESASLVKMEKQLDEMERYERIAENKRKRYVHSEMSLIKGR ALLQDLKMR LT VV DTV PFT TTDASPVQVPAFEWESICDGRGQNL TEEQQRVRYREYVENNIGDVLTTKLCVLGVEKGMDILSVAVPGHDIDL AGRSDILMVIEVKRVLKSGCTFQALSELIALDFLVDDPVMALLTNLTDHWQFFWVSDKKKKKHTLLAQSSADA EINLPCIEEPVKRRKLAEVLPVSEGGESNGVREAIERYYDIASV LGPDIEMARAVANQVTRSIPIYSSYPS*
>jgil540976lestE xt2_Genewise1P	MTIYCAVVGGENESA FGIDIDE GKSVDQLKQAI KERNEDIHVPSHRLKFLAKKSGVWLT EIDVMEDVSDTTDLELLEAGRATLRSVGLSDEDVGEVDEADAAGEGPVNVLVVIPEL

lus.C_PHYCAs affold_50927	YNVPSPVMALFNVMLPHVLTQVPTTESDVVIEFKMELCKFYECYSRHTWVRCLLD VAFPKSLVSASHLFRCNNAFMAPLTQVLWDIDDMRNGLLFKPLKHAFDHQLSFI TNVFRKLKDPSIYNTRLLDKCDNENVLSMEEMGVLFYNTSLTRNPCEFDQTTFGD VDGSALVFTGLKRPFYRCLNQQARLARVFALKHWIDESYNFTDFWSEVSLDDKMDM FHRISILEN*
>jgil556686lestE xt2_Genewise1P lus.C_PHYCAs affold_940014	MVLVALTCALVGKTGVLGVKIDDSAQVWELKKAIKEKSLNKLKNADADELELSLAKK GAGWLSIEDLAIIQKGEDVPGFERVSLVDEAAYSASIRDVLKTNGMPPPQTRQIHVL VVVPKDENDRSAMALDVPSPPTIHRHPERLKRWAAINEMIRQKNQEGRNEKTSTRDT NKKRKNRDIDSSMPYLSLSWTDLPEILTMDFNLEASA AVPQNVVEELRDRLMQVRKLY GDVYSGKEAKRQVFIMPIFEAVCLMLGDATILVEEDVKGKVN VHGRFEVLIKHGKKR VSIVIAKRDDIPQGIAQNMAGLEALSDVEGLERTLGIVTNYLEWVFISDDDEKIRRMNTT LKVGAVPSTKELREIVGMICGLLANST*
>jgil558527lestE xt2_Genewise1. C_PHYCAscaff old_11463	MMKLYCAIVGVAGSVFAVEIGEDKTVYDLKDAIKTQNKKVDA GKGMWLTEKVQKGVNNTSDFNLLGTVGAPLKFGVGLKDDVEFEPTLKDV ESMNTPV HVLVAIPQQWTISKKTDAKRLEKEENIPL EMLWQYSEMEITTFPQPDELSSLLQRPLPFQ LNLKFLTPKTIFDPSPGFLVCNELSALIDGFSYSCDYRPDPMASENTW QRMYDQLLDIS YRLCRAHGFDVSNRN
>jgil559084lestE xt2_Genewise1. C_PHYCAscaff old_21063	MIWLYCAIVGKAGGVFGVKINKGDQVWELKKKIKDENQATITCD ANEQLFLAKKDG MWLPDEDLVAVDLENGTIHPDIDKMMNAEQQM QDNKTLQFWLFEENEMP KPSTDQIH VLVVVPKQDGTLNETSVA QTPLQDEEASAYSFSEL NSAMRDQIVRK MRLVEN VPDVKE PEDTSIGGYSWIPKIEENEESQRAGYMAYLQQHLKTL IDRGDFLLDDIADDK SVLDIVDP RLPFAMSGTADVLLINRTSKNPLIKLAGVSL VIELKKV EPDV HVPQAIGQLV CSMKAPL NCYPLSLLTDLNDRWHFSWFSDNHTLT QVTLK YKPKNA FRLIEAA VLRRT EVSLPPSFIP GPFKI K VDDFLLQPDDGYAEEMMERYELMADV VEPEFL MAR RAEYA QHLV QSMPMY AHTFK*
>jgil563460lestE xt2_Genewise1. C_PHYCAscaff old_120265	MMKLFCAIVGARS AFSVEVG EDQTV EDLKL AIKNQN RNK LMSV DANDL QLFL AKKDG GNGME WL TQ LD VV KG MD ANG FA HLL FV DA KL RA VG LDS SEL GK V ND KD V AV KG HV H V V V V P K T G V SK D GG R SI Q K F S A PI E F SE ME HE EG GP ST F R T E R K D T V A A I A S L L D R I C R D V L L I D E A Q I T Y N D E Q L W R G F V K D T L E S Q I P G L R L V F S S Y G S F D V Y R K Q E R P G T P I L V P T D N T F G L N V T P K G L Q L S R V E E M V L N S I G A V S D L I W V L C S H G I G I A R V E L L Q T K F G F L Y E D Q A K Q L Q F A S N M H L K I W L L S S R T D P I G Y M V K D I S H G D F I V A C V Q R M S A S R L Q K F A T E N T T R V A R E R Q I Q M E L Y G A T T S C L P K G V L V T P E W R T D D G K G F I D L V I R G S G I L W F W E L L V N G D D A V C H S K R F E T G G T N Y G S L T R N C R Y M L I D F R Q N M G V R K R K D G F L Y V S F A D S F T K A H V F G L D K P T V S V E L L S *
>jgil567378lestE xt2_Genewise1. C_PHYCAscaff old_240430	MVNLF CIIVGVAG NAFEV KIDD GAS VA ALK KEIK QENN NIK LG V DAG DL QL FL AK KDG AW LMS KD LL RM W NE ET PE E D D P T V R I K E F P S E Y P D R S I H V L V Q V P R E LL Y N Q P K P R D S S N E W L A E F F N H K V E P R S L P F V G K L S F V T Q P L A K R V K Q E W L N E W A L S P G L Q E K M F V L D D A P C M E F T S L I F N K R T L N P F R R G K T E N A F I S M W D S I F R N V L D V L F T Q H I D R S C N G S S T R Q K R P D F L V L D Q V C F R G E E K P P D V N I S V P T E E L C S K L V W A Y G S V P Y V F G Y A A S G Y D I Q L H A C P L D P L N V G G V V T K N I G T F N L E V K E H L F Q I V L V M L N L S L F Q A I D C P A S G R D E F R D I T R S S G W P S K P L Q T M L I S G K N F S T N V G D K R T T L K R A S P V N D L H F Q K R L A V F K P R G T M V R P S N L L F G A L K D V L Q A L V H R L G W I H R D I R W S N V I R Q T G N S W F L I D F V D A A T N P Q Q Y P G Q H L S V E E H A P E I F V E N G V H T T A V D I W A V G F L I E T S G V E W L D F A G R T S L Y R R L I A K D P A A R P N A E E V L A E E A K S E K E A C E N S R R S E T Q C R K R K L P D S*
>jgil570403lestE xt2_Genewise1. C_PHYCAscaff old_370147	MEVV KL CV IV GE AG SA FV K G Q D Q E V V D D L K E A I K D R S D G K I D V P R P D L Q L F L A K N G N A W L S S D N D D V K A L K E G V K T L I D E L T Q K E E L Q G E S G L K K V L A G M L T P S T D Q I H V L V I P D Q S K A P S V F E G V L D R C R D S F F L Q L P T A G E D D S W L M F P Q P L L T E R Q K L Y I R S S Y K S I A A Q A L S K M D P K R R K Y A V V T G P G V G K S V F L Y V M W K L I K E K K R V L L M A E E P A I Y F D G E F N K L E P T D V F Y M P L W T K E E L S T I A P L Y P N A Q D Q W E N R F E G L G G V P R L V L K D L K V T P Q E L L Q T A C S N C C L D G D Q F E D Q N G S D S D H P Q P *

>jgil573037estExt2_Genewise1.C_PHYCAscaffold_510047	MVTLFCAIVGIESDAFAVEVNENDSVYALKQAIHARKMYEFSADKLLPAKTEGPAWLSNSSEDVVKLEEKEKTPLEALTTKCQELQAELPVSMDLKEIDPPSLSQIHVLVVLVPDLKKRKRGGTTRLSLLETCSKEGSLPTEGDFLQMFEWDDQDCGVKDITAIGDIVGFTGFRFFVRKEILCVLENLKHKFKANFDRGEVGQFIFLGSPGTGKSCVLALLCFYVAATSDHPVLWYRSVQYGREMSFTCLFYQKYYRWNGAEVKIYDRLYDEV
>jgil573801estExt2_Genewise1.C_PHYCAscaffold_550284	MIKLFCALVGAQGS AFPVDIDASQSIGDLKDAIKDQKQNDLKNVDAKLQLFLAKKGDWGLASKDLPsiQRDMASPTIFEKLPLVDPTCSIQEVLTENELPPDQPTRQIHVLVNPQVHVLGKRTRADEFINQIGMKSRVIVDGNETDRITRYFEMAGFPPLAHPKAERYKILERNAYIVIFTELMIKKAKLSFEKGADC SLVVTGNPGIGKSRFYLICIFTLFFALTWKLRSSPSI
>jgil96357le_gw 1.1.1010.1	MIKLFCVFVGAQGS AFPVDIDASQFVGDLKYAIVEKKKEDPNLKSVTAKNLQLFQTKE NGDGWLSSDDGAVIAMRTGAIPEQVKKLLKDEMDPAEQIGDKFRNAPTKTIHVLVVPPAPENERKRKRMEDEVAPDAWIKA KDEPVTTLPTCEGLKHLLRALHVKIPINRLF QIVSAQNSTGELFTVLEKL FEPQPRNVDITGAVLRIIDPLPSGPTTKSSYHHFWDCVIATLLKVVTDGNYHRSTNASASTGAYRPDMCFYSRKSNCVFRGEEKANGELDVPMAELHEKLTWRYDDAPYIFGYAAVGLLVCLVTIQKDEKTSSRAKAEKIETYDLGNLKDRLLFL LALLNLSTLFDPVVDLIRPLGIPEYITRERTNGVRIEFAEDCVIKTYPKNIMPSDGIIRNLKS LHRLMKEHSVPNVVELKNANKKKHVKLAPIGIDRRPVNVDQLMALCDILKALVALHAINVMHRDLRWENVLKYSTEGDKWFLIDFDEGRSPAATVTHLKAESHAPEILSSSH TVKVDIWSVGYLLKTCLQDLPPELKRIQSQCLQTDPSSRPTAKSLLAGIESLIES*
>jgil100678le_g w1.5.1392.1	MVKDIKLRCGAYGE GS VFSV KIKQNADVEALQLAIVNARKGVNNRFNVDPSTLTLYLAGKQE GEEIKWLKDED LGDELRGVYPKQYM KMRSSRILDEDYFGENFQPG RHDVHVL VELPEKVSSVQSVRAF FWLVTGLVENALQTRGVHRLIYRIADAQLGY YDPANMLPDNKPRAFWYTNN DLQFHVLFKEGEVCVHCTCWCC*
>jgil102004le_g w1.6.669.1	CVIGVAGNAFSVNIDENLSVGHLKKA IKGENVNDPTLNVA AKNLQLFLAKAEGGARLSSLTDEASEDATKVKKGEWLSNLTDDVKKLKKGEKTPLVESLTHE NKG LQGESGFKRV LGPFLAKRKEWIWL*
>jgil106201le_g w1.12.351.1	IKL SLC AIVGQTGSSFDV EIDDGEK VSKL KEMITEKNKQDPNLKNA AKNLQLFLAKK GDAWLPDDDPAAQDLIEEGKIHT EIKALIDGNKMKEAW TIEDV LVDNNMTGEGRAPKS RQI HVLA VVPGI LTIERER VDENQD*
>jgil108281le_g w1.15.479.1	MKKVSLQCAIVGIGSSFDV EIDDGEK VSKLMRAIKDRKPLTITCEADLLQLFLAKKGNA WLS SSTD DV KALK GEKTGF IDELMHEKE KMEEYPLSDY LANMNDPEV KQI HVLV VP
>jgil108895le_g w1.16.149.1	MAKLSLQCAIVGQIGSSFDV EIDDGEK VSKL KDAIKTKN KDDPILKTVAA KNLQLFLAK QGNAWLPDDDPAAQDLNEGKVHTEIQALIPDEHRAA LKL VNGESDDYINALTAGEQIL ASKTIETWLYEKIKMEEPSTAQTHVLVVVP
>jgil109378le_g w1.16.566.1	MKL TLQCAIVGQTGSSFDV KIEEGQTVGDLKEAIAV DQKFGFAAS KQLFLAK QPV EDD DGKEVVPVYHPCAEGMKKESFKWLPDKHRAA LKL VKGESDDYINSLTAGEPIASKTL TIWLYEKNNMEDPSTQQFHV LVVVP
>jgil113678le_g w1.24.344.1	MNR LERYA ALNEIVQEKNQAENGKTSNQDTNHKRKKKAL DNSVPFSSLSWDEIEPVLQ LNMFHLTA KPVDEF VRKILAQLADLHQ LYGDVSTGKEE KRKMFTMTVLEAVC LHLG DVMIFVDEELTGT KIHMHG SIEF VLQRGA KRPV IARRDNVEQGM AQCVAC VEV LAD AEGLERTFGIVTNYLHWIFI RDEDESIELIDQPLN ASMPSF ESLK VILGMICGM LESE*
>jgil114206le_g w1.25.261.1	PLDDTLAPLNSVGLSEEDV RYRMTKEDIKAKKVPVHVLVVPPADSPATILLETILPHVL THATTFTEDN RDFT HNL CNF YGCY TREQSLVRCM LL DVPLPKSLV LASHLFR RSNEYLSFRMMQISDIDEVKNGL LFKPLKYAFDHFQISFIRDDT DVFR LKLF DSTIKDTPLID LSD RY GKKV LSEEQTGELV SVADNGSCL FDVGKTF GD CALAFTGIER PYYRCLNLQAR VALMVALKKGWIDESYDFKDFWSEVSL DDKMEMF HRSILNSVAEF*
>jgil117855le_g w1.34.341.1	MVTIFCVVAGPGSVFAVDIGITQTIDHLKKQIKENKPNMIRFDADLLKLYLARDGGAWL NSNDDDFKALKRREVPARIKNLQM EQQLLDETA KLNDDDYFGKHF PGDRDIHV LVEL PEDPTEV LHYKSELVCGDCFLAELLVVTNSFRL*
>jgil120859le_g w1.42.251.1	MVTVFCAIVGVPGSVFSVKIDENESV AELKKAIKKETPNIFQCNA MDLQLYLT KGNV WLTEAHVKEGLRDTSGLKL LNSMKT KLKFLGLRDENDEE GEEE EGMLGV DVL V
>jgil122221le_g w1.47.236.1	MVTVFCAIVGVPGSVFSVKIDENESV AELKKAIKKENPNIFQCNA MDCLSDEGRRVAD GGSCEGRLERH QWVEAVGFNEGETKSSGVEERKRRRR*
>jgil124124le_g	MGEELYDPTDKISAKFPSQIPGGTIHVLLV VPEGGKR HLSNEWFTESFHPLKKRRVG ED

w1.52.167.1	VNKEKRFFDMRDFPSLLEHPKVEFKTIVEREVYVVIFSQLVQYAKTCFVPTSAEPDG KSKPGKDSNIVVTGNPGIGKSRRFLYCIFQLREREDVALLPPYELVNVHKTNVKYDA VSKEFVELNKDVRALQRKPYVIRLVEATSSELTGWRGVSVLFASPGVDGIDNFSKVDG LTFIMPTWTFeeLEDYNSLLSDELKLAEDELLSRYDRFGGIPRFVFSQIMDQTEAKIQSAI ASFSAVDVISYCRKNDAVREKDYSHCVLEMVPTKADFRANFYLDFVSMHIAEAVIDKV HGDSLAKVSEFAV
>jgil125925le_g w1.60.118.1	MTELKLFCALVDTQAVFYVTINGEKTVYDLKKAIKNEKPNDLEKVDTAMLQLYLAKK CDGLWLTENDVKNGVSSTAGLTLNAAQAPLQDAILGEGLHEPSKEDKVAGNGPVHV LVSVPESVG
>jgil127207le_g w1.67.186.1	MMKLFCVVGVAGSAFPVDIASDETVGDLKEAIKAKKMYCFPADELRLFLVNTSVKNP DEEEKKAH*
>jgil127255le_g w1.67.176.1	EDEDSSIDPYWPESDLAEDHEGQRKGYMQYLKDNLHGVLEGGSTTIPPGTSTSGQPKY HLKDTSRMTSLLTKASSLPFGLKGTADLMIIGEVAHSRNDIFADLQFVIKIKKNQCGPK ERKELLLELVAANWKSYSPECAPIGLLSNLNDYWYFMWFTTDRKIARMKLSCPANGLKA M*
>jgil127347le_g w1.68.101.1	MLNLLCAIVGAQGSVFPVLIGESESVDLKKAIAMKPLTVTCKADRLQLFLTKTEGGG WLSKDEVVLATRKGGIPEEINKMLIDEIDPAKEIADVLGAAPTKMVIHVLVVVPRGLY TIHGDHVTARTDGWQSFSPTKLRTRCRLLEV*
>jgil12764lfene sh1_pg.PHYCA scaffold_1_#_21 6	MLLNCAIIGGGDVISIIIIEEWKTVALLKDAIKEKKPIKLNVDAGDLHLFLAKKDGAWL MSDDLLQMREDGKGERGYMSEELKDPVAKISAKFPSELPGSIHVLVVQRFRIEDVIDIL AELEKIDDVKNMTVILCVDALQQLVNDDAKTRNGQFRRSYHACSDDMAAI*
>jgil128012le_g w1.73.136.1	MKVSLQYVVVGAGSAFPIDIGKNLLVGHLKAEAIKEKHDDIKCPARDLQLFLAKAGGN AWLANSTDHVKKLKKGEKTAYIEALIHENKELPRKDPISKYLERMDEPQMEQIHLVV VP
>jgil128338le_g w1.75.57.1	MVKLSLQCAIVGQIGSSFDVEIDNGEKVSKLKEMIQVKNRERIKA AWLPDDDPAAQDLEEGKVHTAIQALIDGNKMKEAWTIADV*
>jgil128353le_g w1.75.129.1	MVKLFCAVVGVQGS AFPV DIDA SLS VGD LKDAIKTKN KIKL KNID AS DL QL FLA KPK DG PWLR SDD SD VIRM RSG A IPEQ VKK LLNE QID PAAG I GIG ALFG DAK P TMEI HVL V RVP DYD SD SE VN Q QR KLT SF Q KLR K ESG AT GEL PV QG DF M K F DLT D DDIG KV LN I KA IG DIV GF TG SDF YT FAR PN*
>jgil128403le_g w1.75.131.1	GSAFPV DID VSL VGD LKAIN VE KTI KLKNV DA ADL QL FMT K TDG QW LR SD ASD VI NM RSG V IPEQ VKK LM NK WTR QM K LAN CL VKN H Q RR QFT CWW RL QRL I*
>jgil129581le_g w1.85.127.1	MVSITLCCVIVGVVGSEFDVNIAKGKSVHQLKEVIKA KNK LL EN VDT REL QL FLAKTA DGTWLS LTD HAV N SL RNG I IPT KV K ALL KRE
>jgil129660le_g w1.86.101.1	LWN DEV V AARI PREK VLM HER ISRG GF GEV YVGVYNGRK VAI KM LPE IRK RI QSV NE FL VEV KLM AALEH PRIVE FIG VAW DS LT D C VV SEL MER GDL R ALL SQF QAEN HPH GF DH DKV KIAL HVA HAL TY MHS FS SPP IVH RD LK SKN ILL TNE FD A KLT D FG AS RER V D RT M TAG VGT SLW MAPE IMAGE KY DEK A DM F SF AV VL SEL D SHV LPY TKL R QET RAS DV AIL QLV LQG K TQID FSD AC PSS IA AL GMAC A K DP TARP TAA QAL YEL QQ TL NS NE FY*
>jgil129685le_g w1.86.155.1	MVKLFCAIAGISVFLVEIEALEVDDLKNAIQKEKPLTITCDAHQLQLFLAKKDDGK GAWLTEVEVKNGVNDTTGLKPLDAVRAKLKNVQLSDSDVGGVDEADEVAGKGPVNV LVML
>jgil131615le_g w1.108.42.1	MKVSLQCAIVGQTGSSFDVEIDSEKVSKLKKAIKQEKISFKDVDAVDLHLFLAKVQK DMTWLYSRSEDVKKLKKGEKTP LIEALT MER HEL QGEDPLENV LNG IDPPS VRQI HVL V VV PKG
>jgil131976le_g w1.125.17.1	MDV RYES VAHYFKPINGTRVLK SIEN CGED FEP D EGRV HVL VV APV T VML QDP PYCH FVVDGVNIPITDNMAFNLPGLTGFWKAFQEVDTEIEANTAIKLP EGTFLLGDSNRGSCIYI RSCYQLQWEITQKV VQDEVKKATNLVIDGNSIGKTYFGYVMLLYLARLGETV VYESY G T K K R V L L S H N V V V E G S Q Q D F S D I L N L P T T F Y I V D G V E P M H Y Q A K T I F L A S L D H T L W Y T FNEKRDQIRYIPVWSWDELSTCREVLYSDVPESVVEGCFHRWGGIPRYVLQYAQSEEK QILLEKTMEIADFFWLLNGYEKLKANNPEAHRLLHYRVNDHFAKEYFDFASPYVQQEV YDRAYKKDKRVL LGFIGGGDGW KW*
>jgil133238le_g w1.380.5.1	MQLFLAKTEGKWL PDN ED LDT LL TLH RIC TCSH P TS WKLSNP DLF GPG VSL GED VV H VL V VL KDAEAGV AS VEL SAL PT VK QRH PERL KRWA A IN EM VR QRN QER NE KT STR DT

	NKKRKNRDIDCSIPYSNLSWVDLEPILPMEEDLKLEECRPSKCSSRST*
>jgil13996lfgene sh1_pg.PHYCA scaffold_5_#_21 0	MVKLSLQCAIVGQIGSSFDVEIDDGEVKSKLKEAIRTKKHLITCEADQLQLFLAKQPVE GDDGKEVVPVYHSSAEEVKEESFKWLPDEHRAALNLVNGEPDDYINSLTVGKQILGSK AIATWLYTKNNMELPSNEQIHVLVVHHTQFK*
>jgil15135lfgene sh1_pg.PHYCA scaffold_11_#_1 21	MMHNLHGEAGNLKLTAELESPTRLVTDVVTQWLASMDEMREDEAAPGRTPKTQME KSKDEERDEEVQKKRPKRKFRKSTINVRKEEKAHLLLEELSGIAYQDGGDQSSSICHIAL MSEYYLFNSPARGKMKTLTCAIVGVAGSAFPVDIDANKLVGHLKDAIKEKKMYQFPAD ELQLFLAKKADGAWLSSKDPEVISMRSQDIPEQVKLMNVEVDPTDDIEDVFEGAPTK TVHVLVVERR*
>jgil15224lfgene sh1_pg.PHYCA scaffold_12_#_4 9	MSDMVFLMLLFEHIVICVSLSSSIFFDAITEKNKKDPILKNVTAKNLKFLAKTESGWLS YDEDLVNKLNNRVDTNTVTATIVDEEREKF DLLSEKKGCLNLFICATKYGDNLQE AFGPAKSVIYTEAMQSTEKKATTQYIHEVILLDLTPQNRAAFFGLAWDDTLQTRLENV IHKAGGGNTAE*
>jgil20945lfgene sh1_pg.PHYCA scaffold_77_#_1 3	MVQISLQCAIVGHAGSSFVVGIDDGAKVNKLKKAIQRENPLTITCEADQLQLFLAKDGK HFGSGFKPDEGQVHVLVVVPCAVAGSKAETIGQNEMNGDGEAGVDIFSL*
>jgil34764lgw1. 83.9.1	MMKLFCVIVGEAGSAFPVGIKPEDTVGDLKEKIKGKNTMTITCDAKDLQLFLAKTHAG WLDSDNEDLDTLLQSEIDSSSYLRMRASWKLSKPNLFGPGVSLGEDVVHVLVVPEG
>jgil49590lgw1. 51.244.1	MLELLCVIVGVTEWAFSVEVDETKSVDNLKGVIKNAKRNALTGINASDLQLFTAKTTD GKWLESNDPDVIKMTSGDIPKQVKLLKDVIDPVKDIGSVFQDAPTTMTIHLVLVAP
>jgil502542lgen esh2_kg.PHYC Ascaffold_1_#_ 2_#_Contig261. 1	MVSSPLPSSTLDALAPPFYPSTMWYPSITDDYYLERPHSPGKSGFVIADPVEPMMAEIPDEE LFDPAFYPLSAMEMQELEQVDEINEILAELDLMESHQUELHYKSEKTRELRSSSDVDAEI YSMMAKASKAKNFSKQHVHLHKNTSFHSKRNMRSAHQPRSVK*
>jgil505958lgen esh2_kg.PHYC Ascaffold_17_#_ 43_#_4096633: 2	MAASPNNKIREASQLDQLKQFTTVVADTGFQINKYKPQDATTNPSLLKAAQMEQY SALVDDAVAYGKGLSPDLSESERLGYVIDKLSVNFGLEILKVVPGYVSTEVDARLSFDT EGTIARAHRIIELYEKAGIKKDRILIKIASTWEGIQACKHLQKEGISCNMTLLFGFAQAVG CAEAGATLISPFGVGRILDWHKAKTGKSSYESHEDPGVSVTKIYQYYKKYDYKTIVMG ASFRNTGEITELAGCDRLTISPNIIEELTKSTAALKLDAETAGKAYTGEKLSYDEKDF RLSMNEDAMATEKLAEGIRGFSADIVKLEQILKAKLSA*
>jgil508616lgen esh2_kg.PHYC Ascaffold_36_#_ 69_#_Contig46 95.1	MVKLFCAIVGVTGAFEVKIGYTKSVDHLKDAIKTVNKITLKDVDAPDLQLFLAKPKD GPWLRADNSDVISMRSGAPEQVKLMNEEMNPAAIRGDLFGDAKPTMEIHVLKVKS SS
>jgil50894lgw1. 79.118.1	MLKLFCAIVGVAGSVFAVEIGEDKTVYDLKNAIKTQNKIKLKEVDAGDLQLFLTKKK KGEGMWLTEKDVQKGVNDSDFNLLGTAGAPLKVFVGLKDDVEFKPTLEDVESMNTP VHVLVAIP
>jgil509061lgen esh2_kg.PHYC Ascaffold_41_#_ 57_#_Contig18 45.1	MGQQQSSIPTTEVAALVELYDALSGDRWRRRDGWKQPTRDPEQWFGVEVAMGHVVA LELPANELSGCLPVASLARLPNLRVLDLSKNQLRGEIPAELGQLTALKRVDLSCNDLSG AIPRQIGACNQLQELNLYQNSLSGTMPKELGKLQLSRTLQLQHNNLCGALPETLCELTQ LTKFSVRGNCLTGRIPTDIGRLQLSQLVFLSLRNNELTGVISSLGCCKALEFLNLSNQLSG PIPETLGELEDLEYLYLFDNALEGRVPGSIARLKFLKESDFRDNRRLRGELPNFLDGCSSLE AVMTKWKNRKASYRHAILGDPMPSDTPPTSSHQLQTLEDPPSSATFLSQSFHDKG SEGLGDGAPEDDPANNSHLVAEFSRKRVFQLPDSVVQAK*
>jgil509444lgen esh2_kg.PHYC Ascaffold_46_#_ 29_#_Contig27 50.1	MFWSAALDLKTQRAPNRLSYGSFNRSASNSSFHIPQPSQFTSPLPDKRREEYV ERPDTAFARTRASLQLSEHTQTALKMLLQGEREFKKPSLGRTIENPKPKFAQDMYSMS KISSYSAHRAPMKEPLERTPLKPIARAHGQEEPNPQENSNEHNYWMQELEERRAGLRR SRRRTCTPMSERATTSPLKVGVMCSVPLVLFIFLFGGGSDASSNVMDVMNN LLGIGRGDDAETIAARHAQDRDRIASTFARPPEKIQLEESIVMQLDVSSDHQKLRGSQKE KAGE*
>jgil509452lgen	MESVVPCLLDELMHGVVILHHEAALVLGTTQFSVETFFPLLAYVLVHCRLPIIHAQLH

esh2_kg.PHYC Ascaffold_46_# _37_#_4103400: 2	LLENFAITADNANGEESYYVYCVHAAVEYVCNAAGLSGSTMPLGPASSASSAVTPGG CGLTPVAPMSSTSPPQAAFSLDLELEMENELDVVLGLSRDSGEQPEESGTSALQ*
>jgil511091lfg esh2_kg.PHYC Ascaffold_75_# _1_#_Contig233 .1	MKLFCAIVGVAGSAFSVEVGEDQTVDDLKDAIKVKNDDIKCPARELQLFLAKDKG
>jgil52465lgw1. 36.563.1	MVELFCVIVGVTGSASFVKIGNTKSVDHLKDVTKKNITLKDVDAPDLQLFLAKPKDG PWLRADNSDVRMRSG
>jgil52481lgw1. 67.119.1	MAKLFCSVVGVTGSVFLVDIDQAESVRLKKAIKAKKMYQFPSNELQLFLA
>jgil530721lestE xt2_fgenesh1_p m.C_PHYCAsca ffold_750003	MVKLFCAIVGVAGSAFPVTIDEQGSVGDKEAIQVKNRETIKCDAKDLQLFLAKQPVEG DDGKEVVPVYHRSAEEMKEESFKWLPDEHRAALKLVGESDNYINSLTAGEQILASKT LTTCKST*
>jgil532971lestE xt2_fgenesh1_pg .C_PHYCAscaff old_90155	MSIEACTCSLQARLHQVENARSREMKAHGTLVQSLEIELESRRICEGESRLALAKLRGV NCKLEAELQAAHGKARQLLEQLEAQRVETSKCEERLKVFCEESDCQLRAAREKSDNLM EEKRQLNVEVSEVKACLEKAKAENKVLHAKYEEKISEANAKTADIVQLTSQITDKQQEI EVATERSQLLRELTTYRVQQELAQERIKLLEKDVNRTMVTRNRALNWRRKRQELWE NQFISHRIFLNWKLRSAQAKFQSAAAMARCNDDTRSRLQLIEMNVLAQCSSAKEELNR VRQACINECVQMFLRREICQTELPKLLTIVRNHQRSFDNQQRHLEMQLVNQRESFERE RERLNMSRQDFEHCVQKAIEEKRLFQKRQHHAMFQVFTSKKKWELHRRVFGAWKEFY LRSIVGHATHHTAMVFQSQHLRASTSEVSPIRRGESHQWTMPAPIRGESHQWTMPAPI RPISLASQHWRRIDRASS*
>jgil533583lestE xt2_fgenesh1_pg .C_PHYCAscaff old_150061	MMSEEQAATLSEDDGQLQRLETEKRELIDLKKA VVAGKAMKKQLDDTRAENVLMD EEKQDIVVQLQEVVTRYQAIQQELESKEGELMLAQSKEAFQVELQAAQEEADAKAIKE LTTKFSQEEARVMVQEEENNRLTSQVAAMLEDKQQQEKTQTLMQDNQLQNKVDDL MQQVAGARQLQEDTEQNYLEVASKLNQVLRENDELKNKPGTDASVELQLSQLRAVETE LAASADKWRQEQLQLEQQIETLTAVNKKNTTELADVADIREKLLVHGIDLTYASIESL LDTKNNEIQMLTEQLAAAEEPQEQQIIREADKTVEDLRAELERAEEHSSLVTMENEYR LACEELKTEAEKLTGQLDAQTQTQQLNDDFSVERIHKSEENRLLVEKLEECEAALKA RASEIEKLVARARTASSDTKSGSSEMAACKAENLRMIFEVAKTADSVSKLKDHEELL EAHRKKSSVEDTQLVQLASLESANQTLESSDLKKSEELHNHLETCSMQKEDFESVAN LNNAVIKAEEKEKSIIKRDLEDMIKTTENDVLEEQISELHAVAETKTEPDESQEKDREVEEL RSSLVQAKVDFLEQKQQLTALEKKLAVSKSSGPTCTVNSTSLLDAERREFEAALIEMIE MEKKLQVAYEAKQGLESTLQERMEAFTDLETRLSIAEDKITELEQQLEQKVVALIATIEE QLSRGKLEEKAIEFETFKTTTNMLKDERSLRFNEIALLDKIAKSEVQKAAMADSQELA NEELEEQLNELADRIAIEAEKQDLLARLDETVRSEEDIHQLRERLYMLEEEKGSLDDE NFKLERTIEHLESKLDLSEEQKAGLEAAANEASSQQLSLLERVEKATSEIATLSAEKDSL VELQSQLEENVSVLQEDKVKLEQTLDETSSKLRDELDRTVEQMESFQAQLAQSTAKE EVTIALAELRERSEADKLASGEVTAKLEAQAENQTLLENKISVLQMAEKALQLSQSSR DELSEAEARASVLVEERDAVKVLLQEKTAYEQLKTQREELQLRVEQLSSELESSQKKT TEAAIAEETIQTQLKLTEAKLTESVKHDLAEAESAQMVLVEERDAARKEITARDLKI EMMTSQQGELDLAAQKLESELSSILRSKSSADSEATKEMLRDLEESKAQAEETVRSQGS SAQAKESLQSIQEQLAESELRVATLEKERDATRTSLNENMSTQETLSALQEDLQKKVDA LEAELKELRETSSAELAAANETIENLKNAEAAAETLAALRQELAEAEAGCVMVLVEERD ATKKTLSKRDLTLEVLSSEHGEQLKSQSTTTELENLRSKSADAQAAAEEALQLREEL KRATESLESTQNELAESESHVAALTAEYDMVTKAMADVQSERDSLSGLNEELKESVED LKNEMENRQKQFSSELESAAEAIIRLKTSEVKMTASLDSVEQELCEAKSSASLTAEYNS ILATQKEKEAGISKLTSEIDSLQERIQALESQELRNQRETEAQTAEETIRSLKESESQTV ETLEAIRVKLSETEARVVSADEEREAASKALSEMESHRDAQSTEVESLQERIQALESQ ELRNQRETEAQTAEETIRSLKESESQTVETLEAIRVKLSETEARVVSADEEREAASKALS EMESHRDAQSTEVESLQERIQALESQELRNQRETEAQTAEETIRSLKESESQTVETLE AIRVKLSETEARVVSADEEPLEIGSLQERIOPLESELQELRNQRETEAQAAEETIRSLKES

	SQTVETLEAIRVKLSETEARVVSADEEREAASKALSEMESHRDAQSTEVESLQERIQALE SELQELRNQRETEAQTAEETIRSLKESEAQKVVELDSVKQVVLSESSSSALTDLASLR EQWVAKDESHAADIASKDEVISTLKSKEEVMAAYKRLKGHLHELQDRLTQQTSTNDT LKASYEELSGQKIAAVEELEALKIELASSHEHSNTMAEELRQAEEKVAVAEAQRESQME NFKKRVLAYDDELAQMOKQHAVALQEQQETLLKTSAVREEAAATKLLEQKHDQAGT SHEEASQKHEAERMELESQQLSANETIEKRLLASVTVGLEKEIAELKLQVETETEGANA ARAALETYKKRAHTALKASSENKLNLKKTSEATAKLEKELISAKSRVNTLETEDETR KRMAEVESAGDLAQSTRAVESEKRSLEVTLRLEIDSLKAEVNRLEEALENDRQPLEA QINQLSERNAALNQDVITLKEEIRSQTESMEQEVTKEEEIRDLSKQLQALAAAASLAT NEAGRYSPTYSPTEKERRSTASSSRFDSDGNNSFLHRSSIEQSEHMAAAVADSCPIP LASKMAPANGVNKSQTDEDEVCRKLQLNELETASHLFQKKYEDTSALLEANQQKQ RLQELCDGSTQAINIEYLKNVIMKYIESQVPSEKEQLVPVISTLLSFTPQEQQKVMAVHR PNDEGAGLFGGVFSFNGGAAAAPPKPLAAPLNFKPSPTAMSNTTGAALGSKDKNG VLSFGSDPSDDEEFATPLNPFAA*
>jgil537103estE xt2_fgenesh1_pg .C_PHYCAscaff old_740004	MTVKLFCAVVGFGNIFPVVIDMTDTVEDLKYKIKEKKPNAIYFDASLLKLYLAREGDK WLNSTDREAMRAVKARQCPDRIKNLMQEHLLLDAMNFQTPDGVHVLCYEPREDELGSFLENWFPCWSSLQSLFSVEKEKRA*
>jgil537210estE xt2_fgenesh1_pg .C_PHYCAscaff old_790005	MRPDFLLHYLGMVLLRGEEKSATTEIDEPVKELTAKMSWWNPMDYGDLPYILGYATSG ARLRVVMIDRHLHSDAILEFQSIFQRAEVIKLFYNLAFFFHKMSVLAKRTCPSSLMPFT PDVNRKRKIELMDDVIVRTIKRNQCRDRVFRRRLADIYATLQELNNVRERTHLQIVRK LRLKKRLLRVQLSPLGTVRPPMVDEVRVWLQGMLTALKYWHSCDYCHGDLRWSNI VYIPVSSDSGFWVLIDMDESRSKNTTTIDWNHEFQGYTLEHDLFQLGQLMNSFTFSLP SDLEDVQKALLTAVHTPAGDLPTILLNQLHD*
>jgil537308estE xt2_fgenesh1_pg .C_PHYCAscaff old_830012	MKLFCAIVGVAGSAFPVTIDEQSVGDLKEAIQVKNRETIKCDAKDLQLFLAKQPVEGD DGKEVVPVYHSSAEEVKEDSFKWLPEDEHRAALKLVGEYDDYINALTAREQILASKTI ATWLYTKNNMELPSNEQIHVLVVQGKRIEDVIDILAKLEKVVDDVKKMTVILCVDGFQK LVNDEKTFGSTNADQETVDAVDRVCLNKKSKIARIIRGMYSDLSDEENLFSEP*
>jgil537717estE xt2_fgenesh1_pg .C_PHYCAscaff old_1080013	MFEKTMKGFTDEDALSATLVRVHGVLNDIRPVYKKYGKELRDRPVYKKYGKELR DRIHGETSGKLLLALPTACAIVGQTGSSFDVEIDDSEKVSLLKKAIEKQEKFKDVA DLQLFLAKVPKEKQGEVERTEEVKDEVQEDMTWLYSRSEDVKLKKGEKTSLIEALTM ERHELQGEDPLEDVLSGIDPPSVLQIHLVVVPKGEDVVLA*
>jgil544930estE xt2_Genewise1P lus.C_PHYCAsc affold_160416	MPEPPALQTLDIVDEDQHQVVLVGGIEGGKQTVEIANEYGGFTTEEVTAVKTSEGEK LVIEDAQGQATVVEVHELPAEVAATVTEDSQVVNAVTADNQQVQIVVEGEPEHGEQT VQLIDEHGDTVVEQAKAIETSEGVKLEIKTDQGPVSVVVAEVPEEIKEVIKEQEEHQPG EFGGNPFTEEEPAVLQTVDDVADNGEHQQVVLVGGIEGGKQTVEIANEYGGFTTEEV AVKTSEGEKLVIEDAQGQATVVEVHELPAEVAATVTEDSQVVNAVTADNQQVQIVVE GEPEHGEQTVQLIDEHGDTVVEQAKAIETSEGVKLEIKTDQGPVSVVVAEVPEEIKEVI KEQEEHQPGEFGGNPFTEEEPAVLQTVDDVADNGEHQQVVLVGGIEGGKQTVEIANEY GGFTTEEVTAVKTSEGEKLVIEDAQGQATVVEVHELPAEVAATVTEDSQVVNAVTADN QQVQIVVEGEPEHGEQTVQLIDEHGDTVVEQAKAIETSEGVKLEIKTDQGPVSVVVAEV PEEIKEEGPVSVVVAEVPEEIKEAAEDSMQGSKFMTKPEFREWIRKHYEDRLESKVEE QKLEDEERANAGRVKGLEDCIEQASNKFYYGVYEQAPYYQNAVWDVVEEECWTAQQ WEPESPYEHPNLRR*
>jgil547387estE xt2_Genewise1P lus.C_PHYCAsc affold_240409	MSERLDDPTVRIKEKFPSEIPDRSIHLVQVRREFLYNQPKPRDSSNEWLAEFFNHKVEP RSLPFVGNLSSFTQQLPAKIRVKQEWLNEWALSPGLQEKMVFVLDLDPACMEFTSLIFN KRTLNPFRRGKTENAFISMWDSIFRNVLDFLTQAHIDRDSCNGSSTRQKRPDFLVLD QVCVFRGEEKPPDVNISVPTEELCSKLVWAGSVPYVFGYAASGYDIQLHALYPLDPLK VLTKNIGTFNLEVKEHLFQIVLVMLNLSLFQAIADECPASGRDEFRDITRSSGVVVRLLSP TFVEKIFPDISTFGHLELVYGHLKRAASVPNVDRLTNLHFQKRALVFKPRGTMRPSNLL DLFGALKDVLQALVALHRLGWVHRLDIRWSNVRQRTGNSWFLDFVDAATDPQQYPSG QHLSVEEHAPEIFVENGVHTTAVDIWAVGFLIETSGVWLDFAGRTSLYRRLIAKDPAA RPNAEEVLAELMALESAKSEKEACENSRRSETQCRKRKLPS*
>jgil563418estE xt2_Genewise1.	MVKLFCAIVGVAGSVFEDIDDGASVAALKDAIKNQNPNKLMNVANDLQLFLAQKD KGNGVEWLTEKDVGQGGVSDSDLKLLARARARLRRVGLSGEDIVEVDEQIEAEGGGPV

C_PHYCAscaff old_120143	NVLVELPPGTNRAPLSGTDLWLSRFHHARISVLPTLGLGEYIEHSL*
>jgil576078lestE xt2_Genewise1. C_PHYCAscaff old_830005	MVTIYCFAVDVEGSSFDVEIDDVAKVS KLKEAIATNQKF DFAASKLQLYLA KKGKGKG VWLTEKD VQEGVSDTNDLKLLGAAGAPLN VGLSEKDV KFEPTLEDVESMNTPVHVL VVIPSPDVGPKR
>jgil576081lestE xt2_Genewise1. C_PHYCAscaff old_830013	MKLFCAIVGVAGSAFSVEVGEDQT VDDSKLAIKNQKPNDLKDVDREKLQLFLAKQPVE GDEGKEVVPVYHRSAAEEMKEESFKWLPDEHRAALKLVKGESDDYINSLTAGEQILASK TFTTWLFKKNMELPSNEQIHVLVVVPDPSSNAALEYKDPMSNLRKRGAEVMTDVVM EDDAFVAEENRVRVNRVRCIIVGIAGQSFCINIESNARVENLQGAKKSRCRQLKDVEA DDLLIFLAEKADGA WLSADGAF
>jgil577802lestE xt2_Genewise1. C_PHYCAscaff old_4740001	MVELSLQCAIVGQIGSSFDVEIDDGKKVNKLKEMIQVKNRETIK CDAEDLRLFLAKKG AWLPDDDPAAQDLEEGKVHTAIQALIDGEKMKEARTIADVLDDNNMTGEERAPKSRQI HVLVVVPQQWTVKRVTKPDDDSRAKRARI VRLEVAYQGPVPTDFYSVPAETIDTYQQ LKEAFLSHERLVRPLCLLYGPRRGKTTIGHRLVSLDAEPSILVIYCSLTPLSVESEEAF WVALGEFIGEHTRSFQEF
>jgil59630 gw1. 67.177.1	DEMDSVAPYAWMETNEAVHDQRNAYMQYLKDNLHGVLHQDGSTTPHAIRQPRYH IKDTSHMKTLNCTPEALNHGLKGTADLMIVGETAHRMNDVFTDLQLVVEVKKGQCK AYEQQLTLELVAANLKCEKRCAPIKLLTNLDYWCFMWFTPDSKKARLTLC PANGF KAICDFLAGAVDA
>jgil60673 gw1. 22.466.1	MVKLSCAIVGVPRNVFTVTIEDAASVS ALKEAIKTEKKNVMANFAAEDLQLFLAKK
>jgil70314 gw1. 16.564.1	MVKLSLQCAIVGQTGSSFDVEVGESQ TVGDLKKA IKTKEPLTITCEADQLQLFLAKQPV GDEGGKEVVPVYHPSVEEMKEESMKWLPDEHRAALKLVKGESDDYINALTAGEQILAS KPIATWLYTKNNMELPSNEQIHVLVVVP
>jgil70333 gw1. 32.357.1	MMLSLQCAIVGQTGSSFDVEIDDAEKVTKLKEAIQPKNRQT IKCDAKDLQLYLA KKG AWLPDDDPAAQDLNEGKVHTEIEALINGNK MKEAWTIADVLVDNKMTGEGLAPKSSQ IHVLVVVP
>jgil71308 gw1. 16.631.1	MKLFCAVVGAGQSAFEVKIDNTESVSAFKEAIAAELRYKGRPMLELFLAKTESGWLS NDDDLITKLLQN RIDS TSKL KALWPTWKL NRAKNPQLFGRDVSLGENVHVHLVRFP
>jgil71310 gw1. 13.804.1	MVKLFCAIVGVAGSAFSVEVDQDQTVDDL KDAIKTKN KIKL KKVDASALQLFLAKKG KAWLSNDKTGDTVLQSEDVASFEQMRGSWRLNDPDKLFGTSVLLTEKVIHVLVVIP
>jgil72559 gw1. 33.407.1	LVMVSLQCVIVGKEEC SFNVKIDDGANVSKL DVKEQSGGLITVPPPTMQLFLAKTGR EGWLTGDDATAVVSDDLNHTLMDPTL FVKDPKHFGE GFKPREGQVHVVLVV
>jgil73233 gw1. 1.815.1	MVKLFCAIIGEEGSTFPVIDKGQTVGGLKKKI KEENE DDTLKTVAAKNLQLFLKTES DRWLSSNDSVVIAMRKGDVPEQVKKL
>jgil73236 gw1. 1.816.1	MVKLFCTII GDGSVFVVTIDEQQT VGDLKVAIAAALS YTGRPDLLQLFLAKTEKNAWLT YDKDLVNKL
>jgil73237 gw1. 12.699.1	IMKLFCAIVGVAGSAFSVEVNEDQTVEDLKTAIADNQKFNFANSTLQLFLAKKHEGAW LTQL
>jgil80099 gw1. 10.758.1	MVKLSLQCAIVGQTGSSFDVEVGEGQ TVGDLKNAIKT KTLKTVAAKNLQLFLAKQPVG DESGKEVVPVYHPSVEEKKEESIKWLPDEHRAALKLVKGESDDYINALTAGEQILAS KT IATWLYTKNNMELPSNEQIHVLVVVP
>jgil81603 gw1. 5.1157.1	MVKLFCAIVGVAGSAFSVEVGEGQSVDDLKEAIK NLKRNDLKGVDRDKLQLFLAKKK NGKGVWLTEKD VQNGVSDTSDLNLLGTMGPLK FVGLLEKDV KFEPTLK DIESMNTP VHVLVVVPKKRTSTAIVSEEKD
>jgil81615 gw1. 2.21184.1	MVKLF CMIVGEVGTAFSVDIADGDSVDDLKKAI AENQKFGFAASKLQLFLSKKKND TD EWLTEKD VQGGVIRDTSDLKPLVVARQILSTATDVEVKIETKA FEATSPVNVLVVVP RDSTA
>jgil83070 gw1. 12.944.1	MMKLFCAIVGAQGTAFSVDIDVS QSVGDLKKA IKHKNEDIKCPH RDLQLFLAKNEKDE WLSSKDPDVIFMRNG

Supplementary Table 17. Predicted RXLR effector amino acid sequences and location in reference genome. Location is scaffold followed by the direction of the first nucleotide based on the reference genome (R = reverse, F = forward).

Location	AA Sequence
>scaffold_1_R31 4	MKTATAFATVLALIVATNAAQVPPNTPLRGLRPTADTPSVEDDKEDRKDHVVKKVKIAIPVVPVVEVPQ FIPVVPVSPSTVANSNNAVGPSTNVAGPAAAPGAPGAPGAVTPAPTLNGRPAATPAATSTTRSRPTP APTNFAGARASGAQLPAAPVQAGAPGMTGFPSAGVGDNNTNNNAFGAGIGAAGGPFMAGNGLAGGFG GNPMNRFGANGMGSGGIFGGAGTGPAMGNGFFGNGQSMGFMQDSNGLGGFGQQGMNTNG QAGNNNGFGGGQGGFGGGFNRRERRL*
>scaffold_63_R3 81	MVNAMTMAIVLCTIVLCSSRLRAALKQEVRVLRADAAQAAQTASNISSGLTRLNALIARVSDLEKAQAS QASHHP*
>scaffold_22_F1 619	MGPLTGSLTFVACSRGLSWICSSAFLIISLDSTAASWSRVASRQRNKGNACMRISKLRLRSLKTTKMSY PTS*
>scaffold_52_R1 150	MRLGLFLLVALAALFASCDSAARPVNLDPSGRFLRGESFKESLKILPKTYVKEVDDGDKLKAALQSWSMDK LSRKRLAKSLELSTNRIKKLRKIANRHKVNEVFILDQYVKHLKQLKRSPRPLPRVVE*
>scaffold_8_F31 04	MSLIWRLFVAAILLAGCNAFSDEAPSHLTESALMLRLIDTDHDHIGSRDTGKRFLLRDNDEALTDFTTEEG KSSHPSAKMDKAVISPSIQQLSGQIPTALYNFLLKI*
>scaffold_438_R 9	MRLFPLFVAIATFLIATDAFLMTGDSNQISNVDPGGSRQRLLRAHAEVDFEKEAKEMMRMMKVNKTED FAKKLKIAEEIDDIINKRAPGMHEFMQTVKYKRYANYMNFLNDMAKTSEYGDVKEIKAKSRAQVALKTFRK PTTSQSRWQSILASLKGKRIGK*
>scaffold_87_F3 00	MKSIFYVLLTVFALLCGSNAVNSGGRLRVVDSTDERGGFPSYLKDSFTKWRINSKIKSWVKKQKTDEYVLS KLGLSTLTGKDLVKAPKYSQFQDFKVGWLKEATPTTVFNTLGLNKVEGAVEKADDFTYVKYVMALGEK ADDYPLVTWRKLFGGGSLEQLELKRKILLVKRDEIDISVMLG*
>scaffold_29_F3 65	MRFAYVVLAAAFAVFASTDSAVASEQAKLTSVVAENSPLTHSLAEQEGQRLLRAASDGESSFMERTKFYY WYAMGRTPSYVYEDFFKGMDKSIIAENPNYKVWERYKAYYEKRKAN*
>scaffold_10_F6 08	MFCPLCVYLPGLLCCSDCIIPMYSSVQESAERLPLENRRRPRYRNRTPSYTILEIPQGIGRPQVDRHLRP*
>scaffold_52_R8 22	MKILRSLTFLIGTVQAENSSKLLRGPDNPYDVAGLFPGPADTVVDTPTKFEPEVVDEPVIPSTLLNEDE NSVERLVTEVTTYDDYSEYNRHLSRVNDPAFQGQIQLAEPRHPKEHMIPPSPHDDQVNPRLRGWT*
>scaffold_27_F8 34	MQIRVIALALSAALGSVAAVQTTDRSLGTLVCPAVRSAKSPCLWAGENGEVVDSRALRELFLERHYVAHSD REAYGRNLQEHMTYIEGEDDKAFIGLDTVY*
>scaffold_623_F 5	MHLGHLLFTIVILLASSDGMSTMQNAGKKPIEDTVVGPSNSNGIKKTRALRTDNGSEERGFLSNLVKEAKVR WWLETGKSESQVKKAQLLDELSGKAFETNTKLFQKFATKAYANKLNKWQLQDDATTYSVWKILKLENVSSKQ LKSSPAYRTYDVYNQFDELLQRKWGAYKLPEMVSSKEMMAKSSIWGEAKRYEAYVKMALDMEGVTG KALKEHKNYAYNNFLEALKGN*
>scaffold_25_R1 072	MCVAPPTICLIFQLCQGS LAEILNDQVCRQNHPARQQLAISIGYMLDAARAVAYLHSFSPAFVHRDINPSNFL VDAECNVQLSGFGESRSTVKRNGRALRPMKVPELEAKYFTLNATVASPLGTTQLDSPCTHMEKNSAEYI APELIDGLSEPFFYGEADVYALAITLWDILHPNGDKYPQANDDYVQVHVFCVLQGLRPRINSNLPLKCSII ERAWQQDPGLRPSAQQIVKSLEDFQDDLCARLVLDDISDFHDCSSGSSLRTTEDLHVRTYTGAIVDRMIDRR FVRCPAEAIRLGNALMDSSVLHHANHSTS FENNSASRYYFDVDEAQLALPIDSHIERSVRSSSSLSMESSIA MLNVGHNDAPRSSFSVGRTRSEPPCRCRQLGQRLINRKSTHFHRQVQFLTAEPTVDGNTLTALLGGDE ASPQNSYNGLDMTPGIAATMA*
>scaffold_25_R3 36	MYRVLLTVFALLCGSNAKSGGRLRVVDSAEERGFPSPYLKDSFTKWRINSKINSWVEKQKTDEYVLKKL GLSTLTGKELVKA AKYRQFQDFKVGWVWLKEATPTTS
>scaffold_68_F6 89	MRLNLVVLAAVITLVSRCTAVSAASSTNQINLSNLNQALNTMQNHAGDGNRFLRSAKIVNDDIDSDDDIR DEERGGKTWAEKFAKWHARGESADDVYARFALEPVVRQAYKYGQIGRL
>scaffold_24_F4 44	MRLSFLLTATFVLSSGIVASPTTKDESIPSPNQVLSSEGRRLRVHKSSIDDVEERGFNPEKFNRLMNERGYRST RFSNWVNKNYTDYVYNLLRVDSPNPKRIFNYYQTYLENFAPRLISS*
>scaffold_11_R2	MAVVWIVIAAATALVRVSGGGRILVNCSWIASICVLPRLADVDMAKMAARRRVFNGIAALYDCLTSS*

784	
>scaffold_50_F8 48	MRFYFLLTSAALLSNSNATATVSGEGHVMTSADAPARALETNNGKRSLRYYATDEDVETDKYDQKNGKY DNDDEERNLQRHNSPGLKRRRAGSRRIGRREYESDGGR*
>scaffold_46_R1 328	MRLSQVLVIAASFVFASDTVAVATSNQAKISKMEQSPSQRLRSNHYPVKEEEDESDSVDFEERGFTTPDE EDLEERSPLSSATVKKLENIAKGWGTTSSVAMGTSSVSQTAKALLRDAYISGIKSEKNAAKMAILMANK S*
>scaffold_24_F1 432	MRLLFAVVLILAACLAADVAPALNDAPS K R L L R S T V R V D E E E R G A W D T L S N K F A K V I L K P N Q F A V R N M D D P K V A K A A K T T L L K T F K S V D P Q K F N S V E N F F R G K A F N N L E N Y V L R L N K Q D I N K Q T S V A K V L S T G L G D E K A F H L F L T A T Q S S D R A V K S G K F F R D Q L L T Q W A V E G K T S A E V T K L F P K G L G A N Y Y L H L E N K Y S G I L M D L A R D S Q K R A T R L A K L E R A R A A T *
>scaffold_25_R1 150	MCVAPPTICLIFQLGQGS LAEILNDQVCRQNHPARQQLAISIGYMLDAARAVAYLHSFPSFVHRDINPSNFL VDAECNVQLSGFGESRSMVKRNGRALRPKMVKPVLEAKQFALLTVASPLGTTMTQMDSPCMHMEKNS AEYIAPELIDRQSEHFFYGEAADVYALAITLWDILHPNGDKYPQANDDYVQVHVYECVLQGLRPRINSNLPLK LCSIIERAWQQDPGLRPSAQQIVKSLEDFQDDLCA RL VLDLISDFHDCCSSRSSLRTTEDPHVRTYTGAIVDRM IDRRFVRCPAEAIRLGNALMDSSVLHHANHSTS FENNSASRYYFDVDEAQLALPIDSHIERSVRSSSLSMNES SIAMLNVGSNDAHPPSFSGRMSEPPCRCRQLGQRLINRKSTHFHRQVQFLTALEPTVDGNTLTALLGGE EISPQSAYNDLDVTPGIAATMA*
>scaffold_6_R18 84	MRVLSLVTLA F V S A Q L I A A N A D Q T G I A A V D S N T A L L P R V L G V E S K R T L R R Y D P S E F D S E E A V D S D E E A D P V E V A D S D E E V V S E G E E R V G I P G M E K V A S K A T K A D D M V P N A A K G A M K W K I L V Q K N M G Q L V E T A K L V K K L K V G S F Y S N V E L E K M S L S A R Q L D D I Q Q L Q K A D I K S N V F G T K A T A N G M R R K M T R T E N M K L P P E Q F L V S H V G R G A Q R L G E N G Q R L L S A A V I S K G D D I N S K V L I S S D T K K G D F L L P K G G W D H G E T I E K A V L R E V I E E G G V N G Q L L H K L G E Y P F K K G A T A Y A Y M M K A S T V Y D D W A E S I R Y R I W V R T E M L W K Y *
>scaffold_10_F2 067	MRLSVILLVVAAFVAALDPATAANDANTVVAVPNVHESIATGRFLRAHLEDDYPVKDERDEDDNEKDEER MF SFF Q E K A T A L S A F K N L V S Q S G D D L V E A V S N L T K F E V A L F N Q G H T H M A K M V P G H P G M S L D E F G A V V R A A G L S D D M K N A L M V G Y G K Y L A H L M D *
>scaffold_58_R5 67	MVFKCWRITLFVVLTVTAPEILKGEEYGTAA DW WA FG AVLYELL T GLPPW YSENAREMCKRVLRTPLS VPEY V STEAKNLLQKLTRNPYERLGSLLGGPEIKEHPFFQHIDWEMLSFRDAPAPIQPCATSDTVVRPVIIYPVDF V S N F L P C L *
>scaffold_13_F2 050	MVS V S V Y L F S A V L A F A T L Q G T T A T S L S D P Y T T C T S Y E N F P G R D T D I K D G G S C T V I V P E D P S T S K R K L E W V D G N D I A D L E A H F G V P M E R K L R N L P T S T H S S P P W S G G N W L V N V D S I N Y V W D Q G Q P S A A E K Y A T A F G L D V K T F M D D V S A Q N G I D S V Q N A T E C T E D K E C Y E D L I C A K R A G K S S G R C L P T W G M G N A W A A A T L E K E P K C P V T F N G V T F Q P M D I K A L V T D I Y D S A N V S Y V F T G S R Y N G Y E D S I D D Y G R H T D A S Y R D L N P G F L H I A A T N M L G L L N T T F I V D K D P D Y V V W N Q P V V G F E V R E Q M N M T P A E A A K K L Y G L D T Y P W N V N A S S I V V N S Y L S W V T E S L N G P I V T L N T T E T G N Y T L L E N D D E I I G G E W L Y D S N D Y H P D F L W L E A K P A P D T A T S F G L S Y T N V T M L E K A T E C *
>scaffold_12_F1 755	M Q T I R A I V T A L T L A T I V A G D E G A T R T L R G A M A D N S E L D Q L E G S G S N N S I L W V N A S K P E E R R P I R P N Y H H G H Q Q L V C Q I N F P K N C W Y V D I F D Q *
>scaffold_81_R7 5	M R S H A L V L L S T A L L A C I A G V S T T N P F K V T V T E T R F L R K S D P K T N E E R V I G G S L T S K V G D F I G S T K L Q K Y K L R W K T L Q L D D D L S K V L K S P Q V E K I A K Y N L N K A G S Q V S M F R R I A A K Y S D D I L A R T L V S I E R S A D D N P A L L A M V K Q L R E D Q I A N W L K N K E T V P R V V S K L K L G S D E S I F R S K A L D V L E D F I K K Y N T A R N G D E S L L K T L T I Y G G E S E L V T M I S R A Y P Y A Q L N P Q S V E K A N N I E N Q L I S K W R S E N R P D F S V M S K V K F G D D I N E A L S S G K V R V F F K Y S A S K T M A K R L N A K Y G E G E V A V A M A R A K S T L S G H D V A V V V Q Q K Q M T G W L S N G Y S M E R V F I L R F K E A D D F V Y K L D A E E Y V K L L K S K N P Q D T T N I L Q V L R K G F N G E E D K L A I A L S R P L S E E A K G Y Q A L L F K D W M A R D M D P M S V A V S V F K I P E A D V A A G F S K E I T P I M K Q Y T K F Y N G A A D I Q P L P A V R N G R S *
>scaffold_81_F1 03	M I L F R T L L V A I G T F L V S D S V L A A S D D S T F A R R F L R T D D S E E K T I N L S G L E Q S A K L S K V G L K I D N V D W I K L Q A K S D T A F K L L N L D Q A G E K A F K S P Q F K T W I S Y M S S I S E K Y P E S A I M S T L A A R Y S D E T L V K M I E A T K K F E G M E I A M K M Q E A Q V K N W M Q S G K T A D D L F E V L K L D R S M E N L L T N P K L D I W S G Y L N L F N K Y S K P G K E T T V V N T F V T F Y G D E V V A K A L E A A K K V P N T K E K A T E L Q A L F T Q W L M E A K P H Q I W K M L Q M E K A T W M R N P D A N I W R E Y L A F Y K L H K *
>scaffold_47_R6 72	M R L A F A L L V L G A V L L S I S S V T A S T Q A D K E D D V V P S G E D T R F L R G L E T T D S E E R W G I R D L F G R K K F E Q M L L N N W D E L N K F K V D T V I S K L K G T K Y G A E L L T Y L N S R P K A V R A A A K *
>scaffold_17_F1	M G V T V L V L S F S V G S G G V K S S T M C P H S D C I F D L H E L M Y N L Q I S A I S F G G M Q Q N G S M L T R A V S Q R N S N P T F M

914	CVPSLSRMLRILSGKSYREK*
>scaffold_31_F1 334	MRLLLWALLVTLVAFVSSINAETADSKVTQDSKEEINALTRLLAVDSSDAAKRFLRGDAKDLTTANDDSKELS AEGEERGLIPSSITNLVKVKTGWANWKAKALEKAFAQHMKVNGENPTTLAKRLDIGRTVEGRHHRLYEKFTA WWINYHTVAGT*
>scaffold_143_R 10	MRA SIVL VAVIAV FTRSGLASASNTAQLQQTVPGATNKVENAE GGRFLRE GLKDE SDDL TSE ERGIWGDLFK KAKPYLNGDIMYARLFKKNSSQLWRREITPDELKSMVIHLEKAGWSGEKLQKLKD KSHG YEKYFYTPIKD*
>scaffold_40_R1 065	MAFSSFLKFLAMATLLMVQINAEPSSQCTRNLRIQSDDLPQE QARRLGPWNWKALIINVKNNPGVSDGVVNLL KKADFTGTTSKKIKEVKD TAGKVKD KVGTGK*
>scaffold_51_F6 03	MQLTNILWMATVILFCNDAAASRPELPTISGVDAVQLNPVNQRFLRS DYN SEERAVTLPFAKLDVA VAKL TTKLFNTEKLKA ALGKTKEAYNAREALFKEVNAVNP A VRKAFLAKLAKSGDYNWL PYWL GFKLKGIRKVETT EIKNAV RAGNAVS*
>scaffold_2_F40 6	MVVVGVGVCVG VASSATYAVPTDPDCAI FSSPQGVWNKYTGILDIPYGRVWN GRCWA VRPIRKPA KLRKR RSRNQHQHRS PRRMLRLN WLEPLIVTKT KARAMTQMFKNLLTERRRSRKDVRQRPQYAPGTARSVEEK AALRPVGQVICYACSKPGHFAKECPDAEANARNDAYLASRIQRSGAKYNGERSL*
>scaffold_107_F 190	MRVLSLVTLFTFVSAQLVAVNADRTGIAAGDLNT ALLQGVLA VEKS RSLR YD PSE FDE RESEE EADS DEEV DSEDEERQIGLPGIEKLDDAVSKFTKTDV SVSQA AKADDV V EKA AKAS PKW KALMLK TWM SLLRPEDW*
>scaffold_1_R39 4	MKTATAFATVLALIVATNGAHVSPNTPALRGLRPTADTPSVEDDKEDRKDH HHVKKVKKIAIPVPVPV YIPVPVSPVSTV VASSNN AVV GPSTNVAGPGAAAPGAPGAPGA VTPAPTTLN GRPAATPAATSTTRSRPT APT NFAGARPSGAQ LPAAPA QAGA PG MAGFP NSVG IGDNTNNNAFGAGIGAAGGFPM TNGLAGGFG GNPTNRFGGANGMGSGGMFGGGAGPAMGGRLNGQSMGFGM QGGNGLGGFGGQDMMMANFGQTGN NGFAGGQALTQGGFGGGNSFGGGFSRQRR*
>scaffold_17_R1 815	MKACKFLVLLPSQALANHAVLHVCDRLFRYTAIQREL RKKLVGDYVSMRTEMNQCQKWL SALRHEHTT TLDRLK EITR LDSL VEA TISM NKLQ*
>scaffold_11_F1 315	MRIGYVTLLATTAILASFGNVSGD TD LDTQVAKVASLDAVLPINTNRFLRRRENYDDEEAATEERSAFEALA AKLDKSVLP AFLKIANVKD SERAFH ILQ QYPISLEKR LD LALHGLSKSDRKV LLIQ*
>scaffold_14_F2 398	MQPTLVIFVNVL LLLVGLIGRSDAVHF DETAQAS PSD STIESKTATNERFLRGNDKTIRTTEEDRDITV ASAKAVNMRN KINIMLARGSPTRVLKKLN VVRMSDKNFNNFARFYARYLDKYSRKKP NLPKTAEDV MKDWLAQKLLPFQVEQNLKDLASRN VNRYM QLYFKDADNII LPRLERWENQKL LPSHFKTNLIEIGVK KYM EWYMRNGGEQVVKAKLQKWL RQGIDPREIGAKLRKIGVTD AKYV AWYIDAPV PPQILDELIRAGVT DIQ RYY NTISVIY GQRQV RLYR KNL P*
>scaffold_53_F9 3	MRF TYFVAVALTGL LASNDLV VASTADKTVVHDQVLS DREL IDTGL NDNE KRS LRTN ED LEDD SEDLK RFSIQLSNQPRYYVWFEDEMTPKDVR RKFGLTRHSIKLVKRSIYRGYV KYYDEHC SYFENRK KDFC RAQ EY*
>scaffold_18_R4 67	MRLNLVVLATVITL VS RCTAASA ASSTN QTNLSKLDQ ALG YQV GTM QN HAGDGNR FLRA KTV DSDDEERGGKTWDEKFAKWH ARGET ADDIYGRFFLEPIV RRAYKTANIGSLKQH EYYQ KWL DYKAFL KAKE K*
>scaffold_25_F8 10	MRITYILAVTVAATLHSSVTAIPSVKSSKV ATENGAVPAVIDSTHAGAGRMLRWVKEYEE DDDDDDDDDDLEERGF GDTLKKANPLKLVKKGT KLTAEQAAK V KEALKDAADYQKMIENANKLIRS D*
>scaffold_6_F39 33	MRCFCVVLATV FALLS SETPAIATAEKHSEALSFINGNNYGGTANRSLRSHDKV DDIAAALKNYDDEVVLFTRWWQHEDDVN KLM EWP NRMK NMP ILTKF NDFRTSMHYDQVL DLDEVATALRASKMNLAKEQFNVWHKSGVKPAE ISAAIEKV KVNPLKRKG Y GALN GLYKMYV S GEA KKTARA AAKTEFAAKRAQAKKIAAAKRAQEIKTQ KETAKKIADA AAKRAAELKAKAA VAA*
>scaffold_193_R 7	MSKATPNLLGISWLLGSLVGS HYHLLN RPLCAWNGRLNWPPP RLGVGLRFHQ QR RTL TPCFLPTLHWL RHL RRQ RSTKT*
>scaffold_686_R 5	MRF HAL ALLA AAF LAFGSGND AFEL TADGAA AYSRFLR KSEPKT KLQLGDDLTAVL KSP DVAKIAKF NLRAS GSQVSMF QR ITAKY SDD DILARTL VS IERS ADDN PALLAMV KQLR DDQIANWLKN RETV PGV VSKLKL GTDES IFRS KAL DV LED FIKY NTAR NGDE SLLK TLTTIY GG GESEL VTM VS RARAT P YAQ MNP QSV DKAS NIEN Q
>scaffold_5_F19 44	MRLSYALPATIIAVTFLSSGNAVATT DGGTMDL SAMTSPNA LEEEER KYTNMFSTT KLD EMLDG TKM MSRF KK WKEKG YNTY NLPA ETQGD KYT WIR QK YRD LYHN*
>scaffold_2_F15 74	MGCLHLLL AATT FVACT NAADQ LE MPV KRG L RANE ERA FGA GAGVTEGLSNWVARA AP LLL TDNE LEH LA MKV TSTD KVFKMLK LDDG LDG ILRN PN LKA FAS YIRK VHA TNPD QV LIT LTRY GDD TL AKFL FEAK QVR RTE

	ESAKMLQAAQFIKWFDDGKTPNQVFNLGLKHLTAYEDKFHKLWWVYVAYAHLSKLKKPLPVEL*
>scaffold_24_R7 96	MRFYFLLVATAALLASCNAAAATSNQNKLSTMSTDAVSALESSNDKRFRLRSYRKEDDGDDSDDEERS VMTAEQVAKWTKVERWVKKGHTPSYIKDKLTALDGMNAKNREKYRMFGAAWGRANPHELGRM*
>scaffold_10_R1 115	MRLSVILLVVAASVAAVGPTAAANEANTVVAAPKLPTARFLRAHLEDDYPVKDERGEDSDDENDEERMFS IFGEKATALKAFFKLVSESDLVMAVSALSKEEFVALFNQGQAAMAKMVPGFRPGMSLDDFATAVSSAGL KPDMQNALMVGYGKYLAHQMG*
>scaffold_3_F29 10	MATLVVCASFALASTVADDLHQEAEGNVDRRLRAAAPANKDNVAKIAGGFLTIKEGSTLTAEQMIKNA NGDEAAVKKAILLASTAKESAKMSDESIAKLSAMITTAVVKDPKSWPRLQKFVKVTLGAGVGGALYGYKLL FNKNSSSGPVTTTGSADLVAASTSGGSA*
>scaffold_26_R1 068	MLLCVVILAFLNTVILVLCPRIGRVFVFSTVAAVWFDAGDTVSLSPVLAPSSWFVIRLAAAFGEWILGEDS RDLRPDDMDSRSLVTILTEELAEVGGVEAEGRFCSWLVGFLLPSAIVDCEDRMIFELLPAWGRFDTSVSLATC R*
>scaffold_8_R11 92	MWLLLGVGFASFVCVDARLGSRVLLCPPSSPPYRTPYCPQQFSHPRPRYEDTPQRGLRAFGAPALDFVLR CRIVPRSSRLTLQPVLTV*
>scaffold_93_F9 1	MRVLNLTMGLAIAFIVNMSAVSSADKPQLAVAGALPISRFLRHTHYTDEERAIGLNLLPGSKISSFMTDKLSK PQEQQNRRRRFQQTQARQGRQQVREPEIPRLGSIRGRFQPETPDPELDASHACATVWR*
>scaffold_24_F5 45	MQSAAYIAFAAMLAAVKAESGSILQGRQVLRGQLRVHQCADHALGHGRGRQAVPAVHEGGRRLRHPG SVPVLLHVEDGLPVH*
>scaffold_2_R11 64	MHFHSFIFLHHTSMTSAASCEAARASRTIELLAPVYEDVCQAIKGGSTTVPTLERTADDWIQLVFPQT NVRKDVCVNGASGCRCDFVRLLRALLTQLVVAATYVLAHRSADVATLEMRVQVHEEARRGTSRD FDLVLCQLATVLSVSRSGKKIEARHEALESARSSCHVGCRHPWDASSGQFNPSIN*
>scaffold_214_R 4	MRLAIKTLVALAAVLLATSTEAKAVQTGGDVNVVQSSHILPGENKRLRSEHDEGKLEDDEEEDDEEERKY GANLFSTAKMEKMLGNDWYRYQVARRWKRDGYTWETLPKDPVVDLVRYFKFRERHG*
>scaffold_7_R12 23	MVKVVRVAVVGAFFMASVATSMCGTTQATPLGAPMNDCNPAYGGEIDPNCMPEVAITDAEGCEHAT GPAMDEIIGSITGASAATRSLRRMEDASNSDVADELETYFGESLELSFTTLKEQYSSASVPTTPWPGSYWPTYQ DGINVIWKTGDVSASEKYIAFGLDPTDFMNKISAKTGIDSRSSKTCTADTDCTYRNDGSVCAKRKGVESGY CVPTWYGICHAWAPAALLEAEPQCDVVKNNTFHVLIDALKMTDVTYDGSSISTVFTGARFNGPDIEMDA YGRYISAPRRDLPGFFHIAITNILGKHKKPILDVTAGSQVWNQPVRSYQVQTMELVDANEASQYFGVSP YPFNSEMVFLAYVKTTSWIVEAYADGPLVSSGQVDAYTVSNDYQVYLEDANYAVIGGEWVEGSKTDHPD FLWFPTEKPDASTITSTGLSYAHVKELLELSLAGSSGGTSENASTSTGGSTSASISGSASSASKTNESTSS SNTYSNSASASSDSSSASSAGSSASTSAAASASTGTSETASASTSSASGSQTTSSSGYTSSASTSSGSS NYVAPGSTGSGSTKTEAPSTAYPSSGTNEDGFLMSGDSYPARSASGEESAAGQGSTPASGSAGTTDATST YTSLGSTVEDTGAJLPSGKDALILTPESTSPSTTGEESTPATTAAPIVGYYTPPPSTDGQDLADVLDTTSPID SRAAVTDEPSEGESTGTTALHRSC*
>scaffold_568_R 2	MRLVIKTLVALAAVVLATSTEAKAVQTGGDVNAVQSSHILTGENKRLRSEDEGNLLEDDEENLLEDDEEE RKGGANLFSSAKMEKMLGNDWYRYQVARRWKQYGYTWETLPKDPVVDLVRYFKFRQRHG*
>scaffold_81_R2 17	MTRDIVFLAAVVLLVQAQALSVPTDSSTRILRTSVDTEDLFDEQRAGLAESVIKKATAALRPNVTPEKLANW LNKGKSVDDFARLQLNKAQKLDNPQFATWLTYTDDFVKKNPNTETSAILTAHFGTEALTKMIQAAKK VDGTEAIATKLEAAQIQKWLSSGKSADDIFTILKLDQVDDSSLASPVDWNNYRKVFAQDNPTNLHKPSVR VLEFVPEPLQQGPTS*
>scaffold_81_F2 24	MHLQIALFLVVVSLVNVEAVPAKAQSDFSFRHIGRLRDEREERGVSANAVETITDAVESKINTAQLKSW LESGESADDVFKLKLDQADKVLGHAKLDEWIEYMKLFNGQKGSKKTTLIKTLTAHFEDDGVARMIQKALQ VDSTAKMAKRLQFEQIQRWLHGHEKTPEEVTLKLKDINRYDLFEKPELLTVVKYLDWKMPDRQTTLFA RISPLLEEGILANLLIKAKSVASTEKIALRIQAEQTASWLKAETPDDLFTLLRNRAEDSPLENPIFDAWVKYA DDFREMYPKVSDPPIATISEHYTAQVATMIVEASKSPSTSSIAHRLNTEQFRDWLNTRQSPVRVFKLLKDE AGDKLFQSPVITWLNYATFYNTKREKVSITLLRKRGDEVLAGILTEAQQVPATKEATKLLTSVGRWPKS RVHPDNVYKWLVRVQGREKTDGFRFYERYAAAYKAARNG*
>scaffold_37_R1 044	MRVASLVLAAVTLLASREVVCATTNTLAKADTDTINAIPIDRARPLRNLRKTTSGYDAEERAEEELVDV ELLDKVISDHAYAKQVFLSWLQNGQTSQDIENRLETGQVQDKYGVVQYAHYLLEERSA*
>scaffold_18_R1 184	MRFSFIVLLAAISSLVTTSHALPALPHEQSTVKSSTNPQSDTGAQPSERRFLRSAGVGSKMLKSESFKDAK FAEWLKQNLSSFTVFESKGLKEYRGLYDEYAKLHKASGQYP*
>scaffold_175_F	MWRLSCWAMAETLSVPSLVCSCFAVPAVLELPSEVPAAVIVGSRQVCLRGHRRHRLWNCCSSLR

27	RLRHPFRLRRASLVVLESSLPPS*
>scaffold_13_F2 21	MTALSAWCTAVNSVDAGKCCLLVLMLYCAAQRLVNPSQLPTGVMGCRHWRNHISRRDTVLQASDT LQDAAYATGVSACEVTKLCELLSSGRSGDAGSRLCPV*
>scaffold_81_R2 28	MRVHSLVLLAIAIIISATDAVAAQSLPEEQNVSRFLRTNKAETDSEERVNLAGFELKIGFVDDIIEMHLSPTFK ELLANKGDAKKAFKAHVDDVADDVFQSAQWKEWAEVVKYAVKKNENADEALAAAMSAYNPDGSLKL LAKAAQNPNKTDIATVLENARQSSWINGGFTPGAFKTLNLDKADDIFDTAPFATWTNFLKAYNEKHPKA MTEFEVFSSVYQQQKFAKLLASADDGKYAQAMKELVQSIDDGVHPTNMFAILKLDKVDDLLTPVNL WVRYMKEFNELYPKQATTMIKTFTQSYGDEKVALMIQEAKSSDEKVVEFAKNLQTAQINQWMVDKTPK EMLTVLGINSQTLTENPLGNVWRAYNKEYTKKMANGDFAFQP*
>scaffold_50_F1 228	MRLTCILLMTAAALVGVLDSAATTGNSVVANAAMVISLAPESQGRRLRVYDDEDSADEKDEEEESA DEVDEERGWFSKVALTSASKFVGKSTDDEMGEVIKGLTPAQINTMFESGEDSIQQILPGFKTGMDFKFDE LVQKLPQEQQALVMSAYTKLHNNGKIL*
>scaffold_38_R2 61	MRHNFAKVLVIAIFVVTCSAEAEYHTTSDPLALTSLRGGVPRTRSLRSKYAGNEERAAGTSAVESLVKAT TNVDDVETWLKRGDNTNKVFALKLQKAGDNLLDNPQLPTLMRYLRLFNEANPTKKTSLATLTTHYGNHG LTKIIEAGLASTSKKTVATAKHLQTEQIQCYWAGRSPESVFGVLKLDKLLPFTWISSDLFEKPGLKTWIRYLD EFNAENPKQKTTLISLSKRYNDKTLANMLIAANKAESTSSIAKRIQSEQTLQWLSKGKKPEDIFTLLQLEKAGD SLFQNPLFPAWIKYADDFRLRYSETDLATMSTLINHYSDEAVAKMIMAASEDPNTKMLAKRLQSELQKDWV FSGQTPNDVFIMMNLDKTLKLLENPLFDIWYQYGLFVNKMSAKANWNPITALRTYGSEKELSGILMAALK QPSTKAMASKLLQT*
>scaffold_13_R7 93	MRLSQVLVIAAASLLFASDTIAVATSNOAKISKMAQSSPNQRLLRSNKYSVKEEDESESVDFGKRGFATNDE EDLEERSPLSNAIVAKLDDIAAGWGTTYYKIAMGQSSVSEAKINALLMRDAFISNNKEKELARGMLMLAN KS*
>scaffold_29_F7 71	MRLALLFLLVIATLVATGNTLAVEHTEGKITSSDTLLQPTAISNPDTTQTRRLRTPEPDGEERRLVNVDLAIK DAVHEVRKLTKWLQFAVWKGLKSPKLIQEWMKYPYTSRPAKILAYKAFWGKGPLKYP*
>scaffold_25_F1 705	MQRMMSTPFFLMITTIGDAQGEVEWMIPADNSPLIKSDNFNSRNCATRTRWATGVALPTSMSQTAPRA AGGRAFPFPIRRCLRFQMCLNRSCSGVLPASSIWISILTSSVVALTSGRMAI*
>scaffold_102_R 91	MQQHLVFCKVANFILSTVLFVALPATHADLNFPQSHEQQLRPLTSSVLDQTRAavarGRHSQRLRRRLRG WR*
>scaffold_83_R2 91	MSFTIGVCLVLLQSEVHIICRKTHKKFSKNSVNFLVIAKPEGYGQPYGLHHPTQVLRLGRVRLRRGRERHF YILDNS*
>scaffold_3_F32 92	MRLTCILLVAVATLVGVLDSAATTGNTVVANAAMVNSPLVPESQGRRLRVYDDEDSADEKDEEEES ADEVDEERGWFTDKMALSSLASKFVGKSADEMGEVIKGLSPQQINTIFDSESSIQKLLPGFKTGMFDFKD DLVRKLPQEQQGVVMSAYSYKLHNNGRFS*
>scaffold_81_F7 0	MRLHCLLLAAVALASVDASPLQLNQQSISKANGFLSKRMLRTAESSEDSEDEERMNMPGLDTVNAISSTV TKGKMKLWLAKGEDADDALKFLNLDNGIEKLRDATNFNTYASYVRQYNAQNPDHEVMSLAAIAARYGDE AAKLLHTMTRHASTQAVATKLSQEVDLWMAAKKTPKEVFAALLKLNQNVDLLSSPALYSMDDFLNAYNR QHKNQHTSLMKVVLGSYDNAVAKALQDGLESASTKSRAWELKERQFDDWIKAGIKPKEN*
>scaffold_46_F8 62	MVMILIFIILETGEFGVPDGDNRHADNTSAQLKQTDIGEMREQGASGKEIQLVENSSTWETKTEFSKQ KYLKKKQQKYMMPVRFLRCTAESLCRTYRLKNPAKICNLREDSLGQMLVYGNIFAGGQVLVDTCMGLVTG AIAERQGGSGRIICPYEGQQPAADILRRFNFTTISSFCEQNGWIVANFLFKNR*
>scaffold_6_R31 68	MRLAIKTLVALAAAIALSTEEAKAVQTGGDVHVVQSSHILPGENKRLRSEHDEGKLEDDEEEDDEERKY GANLFSKAKMEKMLGNDWYRYQVARRWKREGYTWEGLPSNPVAALIRYFKGFRERHG*
>scaffold_93_R2 83	MRSSFLVLLGATLLQSIINSTLAETHITSNVQLHSEDIVRRLRFAPEQVEERGLTVPKFQDIVNNQQLAWWLK RGKTTDDVFAKLKLNLAGSSIFENPKFVAWIKYVDDFSKHNKGISAITSLSKQYQGDDVLAKMLQKARQDET TKATATRLQTEQMGWRSQGISSDTVFATFKLDEGIANLLANPGFNIWARYLTEFNPQKTTIFKTLLEDHFSQ NILSQLLIAAQKNPRTEKFATSLQNVQLQGFLERGESPTVVFKLLQLDKGADNLFANPQYKTWLNYATSFQKT KSDAAPAPVIDLTAHYDISLVKMIATAKATEGTKNMATYVEKSLIGKATDGKEPTYVSKLLWASATDKKT LEAVYLEQLIIHSTVKVSSNLQLEQISLWMGKKEPTDAVFKLLRNKGSENVKRPQFDTWLQFANKFTQNP EQSKSVFATLSAHYDDLPLATMIKTAKEVNTKKIAEYVERGLLPKWAADGKAPAYVLNKLATDKQDKERVL TLMFKEIRIVEDKITRAES*
>scaffold_118_R 15	MQTRYRLLLLVLLAVFLASTRASGASLQETTATSENPRPDGPVPTSLRESGAVERLLSPSEEERMGVKEAIG WLWHVVKMKLKIWFLYRGTTPEQVLEKLKVVKSTDENYKYYARYYFRYYVKYPNKPKNAPTKTADAIMK

	ARLQDWLDKNLSPPQVFKEGLFGSGTFASAQGHGPDYKYFEQYAKMWSDLQVRISSGHA*
>scaffold_31_R7 46	MAIFVACSTVATAESVALTLDNEVRRRLNQQNIAKAAGDLISKESATLTKVINIAktANGDEAAARRAV MLAAGAK*
>scaffold_8_F29 12	MIAPLFLIHICILLTVCRGRHPPFGLVVVLQLEQRRLDVRAQVRNLRIHIVTAATNNNQRQYAVLDNIISIN ELTA*
>scaffold_18_R6 89	MSSSTGSQVPSSFLLGSLAGVPFALIAGTHARPLRAVVTVAASNTPVSVGMNSPRSPSFAATAISMGVTA FAGICALPAIRAGAVVRPASASLVRNGITIVRPFPSTLSRSKSIFAFLSVRKSTPSRNNSVQPEMTHASTQIVS PNTFWNWKRQTPRTSSRNPIPSTYSTGASRGSLFPYCIHAVPATEYLAQLSSLMFCSPSIITHAVGNPSSAS TYCTASPSPGSPCRCTLTRANSVSPGLSPHFRCSRNCSSASVACFSLTASRVSSKVRFSTSLRSSLERA RRTRRLRFLILFLRRLRRRACCCNCFSGPSSLPVSGLSATTNSASSPILSGSAAPSSEIACTSSFAGVVRLPPS APTPVASCCAPSPDRASSRTRCSSLAGRCR*
>scaffold_3_F37 06	MPRATSPSLVLLSYNSIGLGELEPNRSWISVGIAHAAQKLWWISARRDERVILARKLRPGSVRGLFAGAT RLLGWVVRVFGRRHGGGGFFFLELPISIRIVERILFRVGSEVGLLRCMLLVESMVLGDREVTLCRETGSSA NGHCRGCACCRGSRRGLRVRWERGLERWRVHSEMDSSVLPET*
>scaffold_2_F13 82	MRFWVWLLTVIVLAGVDTCTARGQSKLSSDFVRPFVKNAPNTQLESKATRLLRTDKYDNDEERAITLPS GLSNMISNVKKWFSNFISKIKLSFGEKRLNAWLKQGKTPDEVFEILNLHKGTKGLANKNLRTWSVFMTMY NRKNPTKVMVNMLGTLTKEYGNEAVAVLLEVSQRQGKTRPLANRLQTQQLHGWARNGLNTDVFNLRLVGE GSVKTLAQNRALNVWVYYFRQMNFYNKGWEALTCKLLTVYDDIPMAKAFAVAKTDPGTKYISSQLQRFQ FKKWLADKVEPATLLEKLKMDKTKLSFEPTEVYVAYSSFYKAYSKTVR*
>scaffold_60_F7 08	MSWMICWICLVTKTISLMQCIRDQCRLHELERTRNLRQLSFKLVALKSSTDFFRHYYLRCSDPPLLFGAPK Y*
>scaffold_24_F1 319	MRLSFIFLLAVTFSSLVATSHALPHEQSTVLKVAATNPDQSVAEGQTSQKRLRSAGLGSQMLKSESFKKSFK DELLDDNVSSFTFFDMLKSKKKYRDLDEYAAIRKARGAYP*
>scaffold_18_F2 148	MRPNLVVLAALVSRCTAVSAASSTNQINLSESQNQALDTLQNHAGDGNRFLRSANIVDDSVDDIDSDDDI DNDDEERGGKWTAKQFAKWHARGETADDVYKRFKLEPFVRYAYKHRQIGMLKENEYRKWSITWPS*
>scaffold_5_R12 81	MQFTYVLLALCIAITASAABVQDNEYQAVVSPFAKRSRSLRSDTGTESDEERGIIPSSLKDIVKKGTSKVSDW SLLRKYKSLQKAGRTDDDIYKLWVQRQKNTDQIYTRWIRLGKSEEVSQFLKHGLNAEVLYNLSRQGKSM DIYSLWKKLKLPEGQIYNIWLKSKTDDQIFSAYWAKMDPEDIGHVRLGKDPTKRNVPFWEKYSEFYRAK KYRTS*
>scaffold_81_R2 13	MHVRLFLPVLLLGVSDTSADKSVPTTRLLRSQKLLQITTSECSESTSDPSTSSKTGSLNRGYPRRSRDP HTDNESNILFYLKM*
>scaffold_7_R22 62	MRLHRTLLAPIAVFLAWNCFDPTTNAQVVAQDLSDKAFAAANTNALRSRSQKSSTSSEDDPIINEETRNTMYT QLFPAWYAAGKTPEDAFEELKNPSTGDENWPIYKNYKMYYDTYKTMG*
>scaffold_39_R4 58	MRTPSFLVAAALLVNADAASSRHLANPNVQNLAHVNDNWLDTRFLRVHVEGERGISAPSVKMLQGW VERGLISDEAVKLISLGDKADDLNASLNAWFSYVKVFNKENPSDKMNMKLTARFGDEALSTMIAKRS SKTAMAELQAKQLANWAALKNPDDIFALLHLSNAKSLLFDQPSVNSWLKYMDFTLKSSVEFAISTLR KYYNDET LAEMIIVALKSSKTEA AKR VETELLRTWLNSMKSSTDV RLLN LSTAPQSEL SLI WR NYV ALL NV DPKF KTE M LQ AW VK GLI DDT F RL LT G NA ADD LL NG S LLS A W AT Y I KV FN Q Q N P T E Q L N L I A V L T A R F G D EAV STM LE AAK KV PT Y HTI AN RV Q T E Q M K L W L D A G K L P D H I F V A L K L N T V K T K L F D Q P Q L N T V V Y L D E F N KAN PNS K TTL F ST L Q T R Y S E A T L A K M L V V A K Q N P N L E S L A V R I Q G E Q L Q F W I K A N R R P G D I F K M L R L N I L G S D L M R N Q L F T A W V Q Y T D E R K L N P G T K L T T L A T L R K Y Y D E T L V T F L K A S Q S P N T A K M G K R M E S E M L R E W F S A G T K P T H P V G V F A L N L G R T G T K V F E S P L Y N V W T N Y V G F M K K A G A F R G D T I T L L R G I Y G D E S L A K V L I A A G K V Q S T K N V A N T L E K E L F A L W K A A R V H P T Q I H K L L R V E N V S R N S P I Y K F Y G D Y V L A Y T R A *
>scaffold_129_R 17	MRLSFVVLAAA VSSLVATGHALPHEQPTVSKVTATNPDQS VTEQGTSEKRF LRSAGIGRQMLKSDSFKKTK FDEWLADNLSSFTVFI DKLGGKEKYRSLYNEAALHKASGQYP*
>scaffold_73_R2 12	MRLLYLAGVAILSFAADATGAKVLVSDDSNHNGAQAASASVSTTRLLRARSVIDERAGGISASASDKLAKL FKSSKVTAVAQQGENCGKCLLSHESREHSLYEGI*
>scaffold_46_F1 215	MRLSQLVIAAATLLASDTVAVATSQNQAKISKMVQSTPSQRLLRGNKYPIYEEDEEDESDVDLEERGFATPD EEELEERSPLSISQVKLEGYARRWHTTWEKVALGRSSISEDKVQALLALRNAYISGVKSEKNAAKMLILRANN S*
>scaffold_15_R1 282	MRLSQLVITVASLLFASDTVAVATSQNQANLAISKMAPSQRLLRSNKYPIKEEDEEEIDLSED SIDVQERDFTAH DEEDLEERSPLSDAVVKKLNTIANGWGTSSVAMGRSSISQAKASALLRDAYVSGDRGAKAAKMAILR

	ANKK*
>scaffold_77_F1 29	MRLSYIFAVVLVAILQSSGTAHPVYEDSKQIVSAGAGDATKVDSGRLLRGVEEKMEAEDDEERLFKNLGTYLKKI PQKIKDSWEVKAKEQLERSRNRKWIREQNAVPTS*
>scaffold_244_R 1	MRLTFVLLAVAASVLTRTDATPATYGEVTSTIESPNRSALKMAETLKERRPLRRPCPSTIYRKSAQTFSETR CSGPACPQSLSRGPSR*
>scaffold_5_R34 48	MMRWRLLFILHALMATLTLPSSCWTVEQILTRGPRTRKLRCSTLPRRDNTEVFDCLYEGNATLQRRIATVTW RRKKLVMKL*
>scaffold_107_F 161	MRVLSLVTLFTFVSAQLAAVNADRTGIAAADLNTALLQRVLGVESKRRTLRRYDPSERDSEGDADSDDEDEERI ITIPGIEKLDAAVSKVTKTDDMVSKAAKASTWKALIQKNINELAETGKLVTKLAKNVLYKDVELEKMSLQGLR QLDDIEQLRKVDIENKVKGKTPDGMRKMEHTANMKLPPAQFLESHIGRDAQLVGESGQRLLSAAVVSK GDDVHGKVLLISSSNPKKGDFLLPKGGWDDGEAIEKAVLREVIEEGGVKSQI
>scaffold_10_F2 271	MRLSVILLVVAAFVAALDPATAANDANTVVAATNVHESIAIARFLRARDEDDEDPVKGERDEDDSDNDKEKDE ERMFSFFQEKAALTEKATALSAFKKLVSQSGDDLTVAVGLSKSEFVALFNQGQAAMAKMVPGFRPGMSL DEFETVVRSAGLSDDLENALMVGYGKYLAHLMPK*
>scaffold_73_F3 95	MRFGGFLLAIALFGLCESTVLNTQQKRNLRLSTTNDEERAIGNISKVDDVADDASKNFAFMINMFKEWDE LSQAQIIARMFKQTTPDEFAAMWIMYKVYQKLGADDIQLKAKAK*
>scaffold_6_F17 3	MRLFIKTLVALAVAFLATSTEAKAVQTGGNVDVVQSSHIVPGENKRLLSERDEGNLLEDDEENLLEDDEEE RKGGNLNFSTAKMEKMLGNDWYKYQVARRWKRDGHTWEHLPQHVPADLVRYFKGFRERHG*
>scaffold_52_R9 85	MRAFAFLFVAAVSILIASGDAALSTQADATSRLRSHHQNTYDAEEEERGLDKSIVKSLPEQFKNMKYKPSNM ENVLESWRTGLQSVDAMVYMKSLGMDFDAISHFVDAYRKHINKKGLPY*
>scaffold_78_F7 7	MHFISSFIIVALPFNVLAELAWELDCSVPLGFPGPLRLGVDFDRAVTLEFAIGFSFELDFLEWPRCWRRVLL GHLSSNLRCWRVSSLRCVLPGRQQRLRSFTSAEIACTIGEPGYT*
>scaffold_26_F3 00	MEAQGLFACLLWIRSVPAGRATVFGDSTIVNQALMLNRCQAPSLRPWVGAIRALGTGRPAFYLQHVRA YNTASDALCNWMMDTLPASDVILRVPSGLVRHRTRRLRALPR*
>scaffold_431_R 1	MKIQLLHLHVLFVQSLDVLHVLRGYYHAILREVHPVPLGLLPDANNHSLQRPLREILVTRISINRTGRPENHYL AQRRRLLLQHL*
>scaffold_6_F13 9	MPLPVAVLAELWLDTVSLYHVVCTWSTAIERQQVRPNGFVRVSVGLRSLRWCRRHLCKLRFHIAPVRRQT SVAIPYPLSCQLLSCSCQCFCAPGIYCCLLYSREFVIRPVHEDHFSLSIVLYHRHTAEIAFNHASSYCWWLDDH KRGVLMHPCKTKEILVTMREFEAVKEIRKVECVEG*
>scaffold_30_R7 9	MRLKSLKVLALSACANVRTLQFSPAGTCASAVSALKHYGRAPARVPFVGHPLRHCCKFESRQRRPQQRKLR KTRLKWRMSSDINLE*
>scaffold_11_F1 229	MKIYTGAMLFVAATVSYALESRQLRSQQQQTGARIVAGDGIETAGDDWNVLLSDRKRGAKQAGTLVRD SGKDEERLVADSRTQTYPVQ*
>scaffold_10_R2 236	MALGTVVVAHTTLFTHASSACLHFSGCPAVKRTAVEDFDSRYSGASTSPPRVRALRRKRAKRKEISYGVWRL HQMTAHTRGMRWSTGFCGPSPPLLVPFHLVFVLATYVIRCSSAPKPSKR*
>scaffold_29_F6 61	MRLNCFLVVAALVASTNAASSIQQNHSQTPDIQAATQTDVVKRSRLRYVKDEEDSIDTKDEERVQGI NVKELNKILSAKRIEIPANIEALSKKAQRDLVHTFKGQDLTKMFATLGMRDVDDVANRHYPFFKWEHLF RPGKEKKVPEMLIGNHEYYY*
>scaffold_46_R9 61	MRLAHLFVVLAATFLVSSGALAATNTDSVIKTD SYVKGSKPATHRALRTYKADFEDDDDDDDDEERGIKI PLERLQSLGRKVGVSADDVLSDAAGVMRTMSESQIKQWTKGKLNKLQKLYKAKSPRISDY*
>scaffold_46_R7 83	MRSSFVLLVAVVALLFRIDAASGSSMKLRSQNLIPSTETRETEQIGRVLRGAEASDPSEERGITDLAKLLGT SKTMFDNNALATIVRLKSQFATADDAVQAFMTRNINPDavykwlklykannqvVKtgdaapeyLLWIRY AKEYKDAHPKVVSKLPKG A*
>scaffold_29_F6 55	MRLAYVVVAVVVALFASQDAVSAEADVQLGNTLVNSESTTRFLRVESTIEAEDEERALPSSLTKISEKLKPG DKVPLTNKVALTEKLTDMIAVRLKPIADTFTGKLKPIADKLMPILKPFAAKLKSAPVVQKIIEKFKTFAEK IKNYKVGGHTLERYTMKFENWFQNKPDDVKTLLKGEGATVNTKNYDLSVQYNAFFQWATRDKEM KAKAAAAATA*
>scaffold_710_F 2	MRLTYITIVATAAILACCDGKSAVGDSKPLAVPQSHVQTGNNANRFLRVHDDEEERIGTQFITSLSDFKASAA DKLRSKSLNSISKVDDIAPGRAGKTLGEVLKIDDVAYLKAAGKWSKNLDDVEDAKLVSLSKLDHKEALRVLD EKNAAAFKSIEDMGYSPDGMFKKMELGKDYFDPFLKKHYTRYWMAKHPTWTSSL*
>scaffold_2_F16 79	MRI STLSLLATVAVVLSSCLASKGEDPSQSIANVDNIRGAGRFLRAQAQVDEERVNKLKLSPESEKWLAGLAG KSAAELEKAAQAPKNVNPEKLLAMMKANAEEHFMLNLAKGVTPKIMRKEYNIRMRKVLSKDALMRNAN

	YQEYERFKEFWNKLPKKLREFLAKAASLH*
>scaffold_225_R6	MVFSVFFSAILLSVSASFNYNVNLETPRVQYRGWPTTWFRSPIAKILWERYLRNLADSIDIPLNPDIRGWHMSNPPLVCNRYLIIYPFVEQRIVHSLGQFCSAGDLLRFB*
>scaffold_237_R14	MQRMRMSTPFLMITTIGDAQGEVEWMIPADNSPLIKSDNFSRNCATRTRWAIGVALPTSMSQTAPRPAGGRAFPFPIRRCLRFQMICKRSICSGVLPSASSIWISSILTSSVVALTSGSMAI*
>scaffold_614_F1	MAFSSLKFLAMATLLMVQVNAPSQQTRNLRIQSDDLPQEQAARRLGPWNWKALIINVKNNGPGVSDGVVNL LKKADFVTGTSKKIKEVKDKTAGKVVKDKVTKGK*
>scaffold_44_R818	MRFSYVLLVAATALLATANAIEADPTSRSLRAHKSHKSQDEERAFTYTFNFSLWDDLFNSLPEQFQRMKEPWYLRIFRSWRSGMTSDEAVFMKSQGLSKAIDQFEDAFLKYRAHKLAKGK*
>scaffold_29_F663	MRFLNILLIAAVLTAAVSASEDAKPIQIKGANNSPNEVSAHRLRAHKESDLDAEERVIPDVYHLSKLEKVAIYP KIVGWRRLRKFDVNAAMADLGVMAREVGTKNMMAIRWYQKIVKNGGFKHTPKGKYITQRK*
>scaffold_111_R171	MDSTASTAVLIAMITALVDAQRLLRSEASTATDSTVSTADFSTLSSGSGFTPGNVEAQSNFGALDKEGSTGGN RPGSMQGGAGRGGSGHNGMQNVGSGEGLAHGSGSFGGHGGFGRHGGHGRDEGSFSGGDFAGSF GGHDFGGSGFGGNFDRSGNGEMPSIGSGEVPTSGSGEAPTAATTSSTSSVTASSSA*
>scaffold_443_F6	MRLSVILLVVAFAVVAALSPTAAASEANTVVAVPNVHESIATGRFLRARPEDDYPVNDERDEDDSDDEKDE ERMFSFFQEKKTALKAKFKLVSESDDDLVMAVSALSKEEFVALFNQGQAAMAKMVPGRPGMSLDEFETV VGRAGLRRDMEDALMVGYGKPAYVM*
>scaffold_29_R1159	MFPRLYAIALIVVALISTCGGTRMDEKQPLALLTRSSRHLRSIDDRPRDTKTEERGPNIDIMISDAAKVKKAT WWKVKFAVWKHLLRKDPHDVRGILGMRNMREALQHKNDEQLAAFWNFYGKGPLKYD*
>scaffold_52_R3	MRICFALLAVALTAVVSGSSVNLRASQKIQSGDVAQYKTSGRELRGDLNTDEAAEERVNFGFLKPDPDKSF AKQQMKYAYNERIFNDLLKFDDPDALYTTLNLSKMKNL*
>scaffold_57_R893	MAFSSLKILAMITLITFQVNAETSQRARNLRIQDDDFPQEQAARRLGGNWAILINVKNNGPGVSDGVVNLIK KVDFVSGTSKKIKEVKDAKDKVKDATEVKVDKLTGN*
>scaffold_592_F19	MRLNLVLAALIALVSRCTAVSAASSTNQINLSNLNQALNTMQNHAGDGFLRSAKIVNDDIDSDDID
>scaffold_73_R384	MRLSYIVLVVVASVFAYCEGASSVSDSERLAVPQSRVQAEQNSNRFLRVHQDDEERAGVEKILTQSASKLRT VSNVDEGGLISKLWSIRKDNDALFHIEVMRLDPGMLLRMRDQGNIDLKKHFTKYWAEKYPDWVSKLP*
>scaffold_18_R458	MRLSLVLAALIALVSRCTAVSAASSSNQINLSESNQVLDTMQDHAGDVNRFLRSANMVDDSVDDIDSGD AIDDEERGGKTLTFAWWNALGKTADDIYKKYNLKLVRDAYKYNQVGEKNNKNYQKWAAYSAYLKDRGN*
>scaffold_16_F1314	MSLLLWSRSRLLSCLLSNVSVLCHGFDIVDFTRCRLRWFFGHCCFSGRHLRFLSCNRSFLFVHWLNVF NLRFNCSLHFFHRLNFSFRVNLVRFSGLLDLRFVLNCIRLHPFRKWESEGFQPVVVKFKHRLPFRIVEKRVEISV EHLKLLVTGRQSVVKDLRFQTRDLVRAHHHEHRQFA*
>scaffold_63_F287	MKLCHIVALTVALVACVDPASASKSKLATQGDNIPTKRLRTTEDEERGISVSLGLEKISNAFTSGKTKEFKN LVKAGQSADDSFANLGLGHVLNFKEKGGEVRTKVMFKFTSDKFKAWSKYTHKMNQKDADSAMLAGLTRA YGEKNAAVIILGKDSLSSHTVAKKLETAQFTKWYTVDKIPNADELVKVLDVPRSKLRKYPREMSIWDNYSK FVGKYVLNPRPGPPVRK*
>scaffold_25_F1242	MRLSFLLPVAMIAIYCATIONAVASNQNKLMSLQARLNDEAGGTLLRVHHENEEDREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWAQKGYTLTQLSDLKSKTRGKYDRVNGYMTYRDYV*
>scaffold_12_F708	MRALKLLLVFLLVSLDATSGYNKLQGTNTVEVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK STSGATNWLKDCKLAALKLRALPMNLDKSVQKTMKDGDIDPDRVFTLNLQKKSNRKINGFRTGEYNLWKE LTVEWTKTYPNWVSNISKKPK*
>scaffold_25_R673	MRLSFLLPVAMAAIYCATIONADSDQNKVSMVQSLDARLNGQADGTRFLRAHHENEEDREERGLTDL FKTEKAAVKKMAKAIMADPSKADDVYQKWAQKGYTLTQLSDLKSKTRGKYDRVNGYAIHLDY*
>scaffold_74_F614	MAGRLCVVMAITLVAASVTPSQSLTERQENDTSGRFLRVASTLKEENDEERAAPVEAVTSLIKKNVA NAAVEAVEKKYPLVIPGVTDLIKARMKRWFWKRTLPTKVFKKLGLRGHSEDRLKNHPFRKFYAEYLEKWD AQAHNVWDWGKAPAALPGTTGNTVLKT*
>scaffold_1_F5431	MKTATAFATVLALIVATNGAHVSPNTPTRLRPTADTPSVEDDKEDRKDHVVKKVKIAIPVVPVVEVPQ YIPVPSVPSVASSNNA
>scaffold_37_F302	MALKIATLSSLWLLFAETFYWKETLSLEVSFAFRPQTSLACVDSTPSNRRSTYRRWLRVNPWQRLRTQS QCETVQADAQSRFSILGFCMEPTAIWSG*

>scaffold_7_F13 70	MRLTYLLVIATTLFSCNNATASGSSAKLVISDSADPLHNGDAAEIVGKRILRGNSNAEKNKYFDEEEVEEEERA LPNLSKLDDEANLKLVDIFKYFDDAGVPGKVFKEAMDLDDLASMMTLYQRYRGAGADDMDLMKKKI GVAS*
>scaffold_2_R34 10	MYKPLLVTVLILLASYTAASLEASETKALDAVPRDRRLRANTEEGVDGTETQHEERGITTGVGNLIKDWKIK QMFKNGKSFEIFNKGVTSVDWDAFKIAKLQNKMSTDLQNPKYANWFHYDVWYAWRTAK*
>scaffold_16_F2 450	MRICFLLPVVALIAAVGSSVNLRASHRIQSVDVAQDNTGGRELRGDLNTGEATEERAKFDFVKKLVSACKG DPLESFAKKQTRYVFNGRVFNDLYKKFPDPDALYTLKLSKYENRWDKGQATPSFKIYGKFRNAYVEKFD WTSKLNWL*
>scaffold_24_R1 723	MASGIFKVIALLVIFAVATTVNAEPVRHLRVEPDHEPVTRKLSRKVPASMLKNFWGIVRTNLENQPTINKAQE MYAKLSHAGHVKKPAFHVKPAGEDVLKHNYRFLDFA*
>scaffold_21_F1 267	MMVVAVRALLVASDLLFLCLHENLGADDWVRNDRCCLSNSQDEGLQCRQRTHALGIASRHLRVLQLEL QFVEDSVVDRRVTHQYE*
>scaffold_11_F6 83	MIATATEVLLISGLTHRGAvgTDNAEGVIDTLADLTPCKSQSFDRVGVGGLAASFPTITKGSSSTAP TRSLRSTGLHKPKPTRASNAPTAKGPKPRQKTAHRDDSAETSTNTKSMITSSRLGGDLRAFRDTVGTKRA DVDGVLDRCEASFAKRTRALGASDELKQKISSICQAASSFGGSIMKVMLFLREEN*
>scaffold_475_R 2	MRLSFVVLLVATISLVTSHALPHEQSTVLNVAATNPDSVTEGQTSEKRFRLRSASLGSQLRSETFKKSFK DLLDENVSSFTFFVDMKGKKYRGIFYNEYAAYRKAHNAYP*
>scaffold_50_F3 14	MRFFLSLAVMILALVADGDAKGSQVLVSNFNFQYQYCASTGVVSTRLLRARSDEERGGGISVPATDKLSKLL KLPKTDEQLQQWLSKGKAAESVFYRMDLGNTITKLFENTQFLRWLQYADDLSDSGKGISAISILSTQYGD DTLYKMIDSAKTTNTTEALGLRLQSEQLKHVKIGKDPDEVFKLYDLNYAGRRLILSTSQFSAWTKYVDDLN KNEGAFVLIVPTLRKYFSDELHFIALAAKRSGTEAMGAKLEDAFVQFWTRRKDTPDNVLVELGLKKSMTL LESPLNILTQYTEVYNVNYAAKKTTVIETLRTFGDDKVARMLLAGRTESTKKIAKQFQTDQLEMWLSSGQ SVDDVYKLLNLPSSRDLSDFGNQKLFNTWVTVINTVSLKNPEKTSAMLSTASSFRDKPMMQILEAANKFPS MEKAATKLQLEKAQNIFSTGISPYEAFLVALDSVGESVLSPLFKWMLYVEDFNKKNPGKEESWYMLR GSYQGGVNIIDKAMKDSTMKIASLAQKEQLNEALTRWKYSPEKLFRFLNVGEAGEKVSAPKFELWKYL DDWNNAYPSSKTTMIDGLVDNYHISALVPMIAAAEKVPSTKKLASQLQDALVEKWVAEKKTMAYLKPRLK GVPSSDDMLERYTKKLNSA*
>scaffold_111_R 154	MKVFSILASAALVIMTALVDAQRLLRSEASTVIDSVTADFSTLSPGSGFTTGNEAHNNFGALDKEGSTGG PCRAELDAVSVGRALYRIADLDIDTTASN*
>scaffold_79_R5 54	MSRFETLSSLLVWRSMCMLVAIPTAAFPPLWRSLSWTASASGYKPINAAWRTFAFFFTRAPPSTRSWLNISV QALISFTHRSLSKMQLSSFRTRCLPASSMQRRNLWSRENLSLPHVPASTLHPPFFDRPRSSAVRLVRPSTTS CCITLPHHQQRRLASIL*
>scaffold_22_F1 370	MTPLHLVLLFLDLGQIHDPPELLVADGAVKDLLQLPIDRSKSACRSFPVAASRALRMLLEHSVLLQSSLPPLS PLLDSAHRPIRQLSGRTRHARLHRQKAASIQRHYRQHPFRSYSFLTAVYLLHPFPPRHLPVPQVFQHSQQRTS RSSPQRCSLPPSRPPARTNLATSAC*
>scaffold_48_R1 250	MVKQPFLILVCLDLGFCRVIPVWSADAPAVTNGSVARSLRAAHLELPLHWILLSEPRTLAHCAFCAQCSTPAVL AKLERSQTWNRWPFSRK*
>scaffold_12_R1 094	MRLSYFVLVAACVIFACEQVAAVKDSQGIAATHDSLRKIDNRFLRGREDVMGDESDEKDSEAENEERAL PKLDVAAEKLMTKAAAKLRSKSLSSLAKLDEAFHRAVVTNNFFTLEIEKMGYNPDKMFKMKEMGNA PLTDPQKVLEYYSMFWKIKYPSWVSEVPHMF*
>scaffold_16_R9 43	MRLSLVVLAAVLFASGTAVSSADPASVAAVHSSRVLSDEDKRFRLRRHPTEHDNEERAFGQNMFAALKLSKM KTDAEYRVKVFHRWKKHGTYADDVAKHVPKALADQYSHVNTFTSGWVMNPLRIFSP*
>scaffold_1_R45 69	MRFPIIALAALVAFLSTLDASSLLVQAASDKLNSGRSLRIAAGTGLEQDEERGLSLITKIRTTFSKKARVDEWAKA EKSDDFVRKALKLENVADDHLWAMKNYPYLLRFLEKTEAKKINDWLIGDVPTYGAWVKLGLGSIDDIEKVKT TAAFKTYEKFKYEDDQAIALWKDYKIPVAKATSPTEMNARMIDLAAKRGDDYAKVLGLDNVTGSALIS HVNQYQFEHYRRTVKRLQDPKKLNRLPTITER*
>scaffold_24_R7 35	MRLQAVVLLVLSAALAVADISAATSGTGGHHVNQNDVSANRLLSEADVSDGEERAATWLQLSKNLVKPN TAPKTMTKLERENPDVAKLVLGKSLDGAFRSLKLNKIKSVDDLFTSKNFETNGFMIYRNSQSRNKPSTIKAFFT KKLGDEQASKLFSQAACKSSNTVKKMGESYQTQLRQWGKEGTLKQVTKIDASLQTRYQTVLDDLARAAK KKAEQAAAKAKAAANTAT*
>scaffold_1_R25 2	MKTGSIMLTAALLSDSVCTAYYLRTGIGIDEDHALDKARVATVQVRYLRKSVEPSFLDSSSTINTSEHHVDNT KAINSSPVSATKLPDVTLSLDDETERTSTVSPNVEIGISSGNMVEFMVHPSERSSVVDIERRVVPAGPN

	RARPTERNFWLGSHPSPRYHMKPQHFFMNRQMDFPVNLPLILPRARQRQHRYFVGV*
>scaffold_29_F1 478	MRCILYIALALAAFARSDAVEAFTNADLSSKISHDFAANELVNNESSQKRFLRVDYPKSVDTEERMKISSLKKIIE LDVKDAKAKKLAAIKDLAKARKDAQKLATNVKSQAQRTRTKFSLEYDNFMRMFKSNAPEKAKSTGKIKT PEQFTRYEDFYETAKAADILTRMAHA*
>scaffold_93_F1 27	MRLHCIVVLAVIAFATNGNEVSAGSRVAITTGALDPTTLLRTQYTDEERAFLGNLLPGSKKISSFMTDKK LSKYLNQEFDDVFIFKLKLKAGDKLFENPKFLAWAQYVDDFNQKHQTQNSMLPTLVRQFGGDDLSIMLE KAKQADKTYGVVALRLQGEQMQLWRREGLTDMFLKIYKLDGATNLLENPGIKIWMRYADELFPGDSTLLF KKLQKTYSDDEALKILINGKTVASSEYGEVGDGFAASSLAEGPCASGESFPVAVTQQGGRCVW*
>scaffold_13_F2 760	MLVERDLLATEASQTDLLLLLGEPERRVCNFHWCASTFLRCLLRRCITSTRLLLFLTSLRWFLRLSLLW SL*
>scaffold_8_F32 13	MCLQSIFCGAAVLCGEWLVIRAFAVKCLRSHSLPPAHAALSRGVLPRLKCLLRYYSYVSTKWHLYVVQRR RWRNLRHQSRILFLVRLPLCDKSPRDNHQFLVQVWDSIPLRFWSRSQTTPL*
>scaffold_18_R4 28	MIHICTFILLLSGERDFGVNLNKPNNHLPLDLPFGSNHADLLIVCLHKIVSGYEKLNSVNCFLTIICNISPYC KKLMNVSAVRLLRLFKLFAQTRYLFDNEANHHLVFFLLETFDNIIQYQYEGNQQVYAMYIQNKNVFYQLNDL QLPPIRASTEKGKEATEKTEEKATDGFVPTVEWLSAWKKLPLTTSRLLQYLPQLEDACKKAGGSLDEDA MLYFLRTTMVGLPVPHPIVIRKYQINQFTHLWFTFTWGVIFLRNQVLPFDGGAITLFTISVL*
>scaffold_8_R13 70	MRLSFVVLLAVAVSSLVATGHALPHEQPTVKTAANLDQSVTEGQTSEKFLRSAAGIGRQMLRSESFKKT KFGEWLKDNLSFTIFVDKLGKELYRRLNEYAALRKASGQYP*
>scaffold_2_F23 3	MMGSVSAFVVWLKSLLNLPCTCSCPCDVQTCYRRELLLLPVRACLRRSLLRLVPRRLEWDLLRFLR ADLDDLS*
>scaffold_4_R23 6	MRLTFLLPVTMAAIYCATCNATAVSDQGKPTVHSLDARLNDTGDRRSLRAFKEKEEFDEERGFLEKAAV KKVGQGNYERP*
>scaffold_29_R9 03	MRLASILLVSALLCGSVFSTEGKQAHTVKNTDGKRLRSDDYDEERGFNWPWLKKIKQLDETTAIKQLATDF ADLKSLNREALDVFLHMREGGMSPKKATYISNLFAKYMENPRLYH*
>scaffold_50_R4 62	MRFSYFLLLTSAILSNSNATATVSGEGHVMTSADAPARALETNNGKRSRLYYAADEDDKYDQKNGKYDDD EEEERNFTAALQAKWTEKAESWVSKNRTPAYIKDKLTGMNGLMTAENRKYYELFTAAYGRANPHALDR*
>scaffold_25_F4 87	MRPILLHAEFWTWLVLAWKACCLREPSGLAFTTYWSMTKTSYRLTWITSFASTRLSLLRMTTRRLRLLL HLLQQTPRMPLYLKMRVKRALYRWLQLTCGASTDASARFS*
>scaffold_7_F31 0	MRSLLALAHHLICLPHPAYFHTLMSASRPLDTSMGLFWLPSSTSRSKIQSVMAGDGGTRPLRSILIADS TPPPPEATLPGDE*
>scaffold_50_R8 12	MRLTYILAVAVVATLHSSATAFSSVKDSNAADSALAEGGRMLRGEYGNALIADDAGKKVSYGDDALDY DDLDDDDDDYFEERTLGDVLLKLNPVRAVKSAEKTKEAAVKKEALKDAADYQAMIAKEMVNKD*
>scaffold_30_R3 24	MGVWFLLLDVALVAATSSVRTALFSLPVELCSFVFLTDAGRGLRPVEASLVLAGDAVRGPWSKFCDPFGAK LDSTVRGGNGPL*
>scaffold_111_F 98	MKVFSILASAALVIMTVLVDAQRRLRSEASTATDSTVSTADFWTLSSSGSGFTPGNVEAQSNGALDKEGST GGNRPGSGVQGGAGRGFGGSGHNGMQNVGSGEGLAHGSGSGFGHGGFGRHGGHGRGEGSFSGGDFA GSFGGHDIGGGNFGSGNGEMPSIGSGAVPTGSGGAPVAATTSTTSSVAASSSA*
>scaffold_25_R8 04	MRLTQVLVIAVASLFLARDTIVATSNQAKISKMEQSTPSQRLRSNKYPIEEEDEFEDTMDFEEGDYTTDKE DLEERSPLSSAVAKLDDIASGWGTWARVAMGQSSISEDKIKALLALRDAYISGSKKAKQAARMAVLANN S*
>scaffold_81_F2 85	MRLNCIVLIAAVASLSTAATSATTDIKPVRPAINLQPPPLVVGRLRTVQDEERGFTLPGAGKLADLFESTALK LTQSARINTWLAKGSTDDAFIKLENTAGSRIFENPKLLTWAVYVTKVEKKNPEEIIARLSKQFTEGSLAKMI ASAKLDSKTEGLATILQAQQRQVWVDAKGSSDEVFKLLQLDEAGTKLFKNQQFSTWSFVDAFNRKYPEEA VSIFSKLAKTYDDFTLWKMLEAKKVPKTEIIASKLQAQQIDAWLDAGKSTDVFNLKLQRTGDKLFKNSQF LTWVSYVEKFNKKDPDQIAIFSKLAGIYDQVTLSSMLEAAKHPSTKRIASYLQQQQNQHWLADGKSTD FKLLKLNTPSLENLIGPRLDAWTSFMRAFNMANEGKETTLIATLTHYKDRLAQLLQEGTKFASTKKIAEELQ TAQFARWLQLGKTEEDIFALLKLKLTPTTDPEAIVFYRKYLFMDAHM*
>scaffold_50_R1 069	MIVSFKKTLFVSAVALAVSVEGSGAVESTYETKRLRLEAQDVAEEEMADDSECGLSLEMAEDDSECGL EMAEDDSECGLSLEMAEESGTGDKGGDNGGNTWTQAPSTGDKGGDNGGNTWTQPPSTGDNGR*
>scaffold_29_F7 68	MRLVSLFLVIIATLVATGNTLAVEHTEGKITSSDTLLQPTATFNPDVTQTRRLRTPESPDKGEERRLVNLIAIK DAVHEVRKLTKWLQFAVWKEIGKKPKLQIWEWGMKYHM*
>scaffold_10_R1	MRLSVILLVVAAFVVALSPTAAASEANTVVAAPKLHESTATARFLVRPEDDYPVKDERGEDSDDKGND

133	ERMFSFLEKKTALSFKKLVSESGDDLVMAVSSLSKAEFVALFNQGQSAMAKLVPGFRPGMSLDDFVTVVKS AGLRRDMEDALMVGYGKYL AHLMD*
>scaffold_50_R9 34	MHLLYLAGVAILSFAAGATDARVLASDDSNNQYRAHAANANIVSTTRLLRARSVIDEERAGGISASASDKLAKL FKSSKVTDEQLQQWLSKGKAAESVFNRMDLGNTTRITKLFENTQFLRWLQYADDLSASGKGTS AISILSTQYG DDTLYKMIESAKKTNTKALGTRLQTEQLKHWVTIGKDPDEVFKLYGLNHNHGSHILSESQFSAWTKYVDDL AKNEGAFVIPIITLRKHFSDDDFHIALAAKRVDET KMGTKLEDAFVQFWIHRKETPDNVLVELGLKKAETL LENPLLKILT KYT DAYNVKYAAKTTVIETLRTFGDDKVARMLQAGRTDSTKKIAKQFQADQLEMWLNSG QSVDDVYKLLKLPSSRRDLLVDFGNQKLFDTWLTFMNAISIKHPEKTS AIFTTLAPTRFDKPMQMQLEAANKFP SMEKAATKLQLEKAQSIFSTGISP YEAFKLAALDDVGDSVLSSPLFKWMLYVEDFNKNPANKESWFLPLRG NYQGDRDLDEIIDKAMKDPTVKA LAKSVERERMKEWLEKWEYSPSTA FRKLHLKTAGEKVFSNPNFELWVKYL DDWNKAYPSNKMTMIDGFRDNYNDLNVLRLFTKA EKTPSSKKLASQLKD ALADKWVAEKTL DYVKSWS HVPSSDDMLDRFTKLLNNNA*
>scaffold_2_F36 47	MRLLFWTLLVGLATSGVQPIAALSKVADTSETSTLDHGQSTIIEGTARHLRAPHEDTSITDVLVERGLKVHDE ERGNFAAAAGWTANKAFNGLGKIVGHDNSNKLTKVAEFFFSLRKKGKTAGDIFAHARKQTDPQSRAKFET VATMYQQYLTKV*
>scaffold_644_F 7	MAGR LCVVMAITITLVAATVTPSQSLTERQESDTSGRFLRVTLKEENDEERA VAPVEAVTS LIKNKNVA NAAVEAVEKKYPLVIPGVTDLIIKARMQRWFWKRTLPTKVFKKLGLRGHSEDRLKNHPFRKFYAEYLEWKWD AQAH NVDWGKLKATAALPETITTGNTV LKT*
>scaffold_104_R 67	MRPNLVLVLA AVATLVSRCTVVSAT SSTS NTQINLSNLDQAL DIVENHAGNGNRFRLRSANIVDDSDDDIDSEDEE RGGKTWAQKFAKWNQRGESANDVYMRGFL EPLVRKAYKYGQIGQLQKNEYRKWAAYSAYLKDKGLN*
>scaffold_36_R1 210	M RFFMVAAVA AAFASSCETASA EAPNAFLIDDNTNAVFSRYLRSTQTDSEERAIAQV LKSE DRTVANVMIK YQSLYRAK ITPKQAKLMLGVSDNMVEFTK MTRPIQRFYTGYYSYFTTMEKRKKWKKELKNQVMV*
>scaffold_379_F 2	MNIAVLLCFFAFPSGLFFSFRRLRAASLA PFLLSDIRWHV LGGPSLQHFTPQKQQQLHALPLQAAYLTSSAAFCS QPLHRRALISLRL*
>scaffold_20_R1 237	MLGFLLLVM TVGSTTEL ILGGWSSA ITPDTRAVLVQALSTTNVCVDSIISVRSQV VAGTNYEFRINGCRGLR GGNCVRASCVA PRTFVNVFDQPWTR TTVMSVSEI*
>scaffold_24_F4 23	MRLSFLLAATFALLSSGTVASPTNDEPISSPNTVLSSET EGRHLREHKN SIDDTEERGFNPAKFDR LMNERS YRQTRFGNWVDKPYTMDVYKLLRVE NP NYRRIFHYYQNYLQEFA PHLITP*
>scaffold_2_R38 00	MKFVSTLV FMTVALQSSCSTA QIDFVR SYDH HAIDFSTSSKPTQPSVADVN TVEANQPLRSGNRRRLDGSA GIVIVGRNGGGFEPLRNGETPD*
>scaffold_11_F1 235	MKA KLLVVLSFLA VTSVAERHLRV ADDANPLK PYEALNAVIEKFKNKAKGKPERRLTGEKA FDKFVKAIA VLKEKD HARRVEEAN*
>scaffold_55_R4	MRLFIKTLVALAAVLLATSTEAKAV QTGGNEDVVQSSHILSRESK RVL RSEYDEEKLLEDDEEEELNDEEERK NGANLFTDAKMKKMLASDWYRYQVARRWKERKYTWQNLPPNVDPLNVAFYKGFRERHG*
>scaffold_55_R6 23	MFFLVKQLSAVN HMYRFSLASFLGLFKTLATKMESSSTKDRILRIP LEHKVLMFVG RALFKEHRPMFGMH LVHGMHPECFDKNEYEFFCGEV VELERGSGGHSTLPEWASPERKEAFTQFVETLPRLTQLCKFESHDMWIR WSKSMCEQNHF PKMDKSGSAGGLSPFQKLLV VQALRPDRLQS AIIQFICGVMQLKSLPPSLDFK VIGTEE ATNTTPVLLTTAGADPSKEE VATSVVGKHYFEVAMGGGQ QEKA NLKSTA EHGEWLCLQNLHVVA WLPVLEKEFSA LNASHKFR LWTTEPHDAFPLV LLEQSLKITFESP PGMKKNLQRTYAA WNPAFIAGNSAR AQLFLLAFFH ALLQERRTYIPQGWTKF YEFSGDFRAGSNV MELACQTSGAA IDWQTLHGLMENAIYGGRI DNPYDLRVLRCNLTEYFSQ DLLSGHKS LTRGV KLPQSTQHADFLDI DRFPDIDAPAMFGLPDNIERSMQRS SGQVIAQLKALSSNEAATT FDR EKWR AQLGP LETW SKLTTGFQLEG TSLSSTGKNLQAMTPADAFVALE NEYALDLVQQVNSILQALKV IYGTGLLTPAI QTVA KALLKGIVPTEWAA*
>scaffold_325_F 4	MLS VLSL ALMLA VVTG AFETS RDVKNGV VTPDSR LLGFTADQ EERKFGG PATG TDHSN SHAWEN FKAWF VRTFFFWRKREQT RRLR NYSSM*
>scaffold_50_F4 3	MGLFYLVCVAILAFVARDATGDRVIVSDGFNQHRENAARASV VSTTRLLRTK SVIDEERVG GIPV SATDKVAK FLKSSKVTDKQLQEWLRKGKTAESV FYRMNLKTSQ LFENPQFLRWLQYADDLSASGKGTS AISILS NKYD EK LYQMI RWAKEKLNTQDLGMR LQTEQ LEHWV KVVGKDPDEVFKLYYLNHAGGGI LSSQFNAWAKYVDDLS AKNEGATVIPIITLRKYYSDDNLIKIA LAKEVDETETM GMKLED ALM QFWI HRKESPD NVLVDL GLK STGT LLESPLFTILT KYTEA YNAK YPSMKT MIETL RTFG DDKVA KVL LAGR TEST KKIA KQFQADQ LDMW LNSG QSVDDVYKLLNLPSSRRDLLGFGGEKLFDTWLTFMNAVSIKPEK TSTIFSKLATS FEDR PMQMQL EAANKFP SMEKAATKLQLEKAQSIFSTGVS PYTA FRMVALDNVGDSV LSSPLFKW MLYVEDFNKNPGKEDSWFLSL

	RVNYQGNRLDRVIDKAMKDPSTMKLAKFVEKENMKEWLVRWEHPPSVAFRELHLNKAGEKVFSAPKFEL WVKYLDWDNQAYPSKKETMIDGFVDNYHTLDLIPILAAEKVPSTKKLASQLDALVDKWVAEKKTLAYVKS WLNGVPSSDDMLKRFATKLNSA*
>scaffold_11_R1 430	MRSYHFLLATAALLSCNATAAVLCEGQVMTSADAAPVRALGTSNSKRSLYYDTEEDETDKHHQKND KYDEDEEERTWSAAQIDKWTAKADEWVDLGKTPAYIKAKLTAFNGVMSDKNRKKYELFLAKWGRANPDEF GRR*
>scaffold_22_F2 08	MALCSCICTGIACFPVSSHFAVYFHLSPHSLLSLRRLCHTDTAERLLRRVLLIARRTQEQRRAEWLQVATQR AERSGSRTIQHGHHITGAGWTCLLLLHRWVISLFPRLHSSTGHQLVSL*
>scaffold_29_F1 480	MRSVVVILAVLALLCNDALAATSKTDLAEALTKNERNRSLRGEVIADEDSYAIDEERVQTSGISKVAKELKLK KANLKKVSAALEKAKSNPSKLAVNAKKMKIAEKKILAKLSSMENDNFLRLFKANFTPQRAKSTGKIKNPEQF AKYEEFYKLAKLIGLK*
>scaffold_3_R39 1	MLPTELVLFVFVGLRQAGYAPPATTSEWPLFSPPLGGLVNWHTFAFLRINRFLRIVHLRPLSDQLQRFLYD RRLKRLTPGICYLVRALDFLLGTHWLSCFYGVSYLAYDDGEMSWLTPDMIAFGDGVRDLADIRKVPLLQS YLRTYHFSIGAITTVCYGDIIPMNAQETEVMAFISVALFSMLSGGFYKYFDMEGRRAEYEERVAQVGHFL KFHRFSPDTWRQMQVYFALSRESRGRRELLSGLPPSVRQDLAQHVHASLLKNVALFTRCDPTFARAIIA ALQHEFFVRNDVIQRGDMERSLYIVESGIVLISAVRKRVHQAGGAEEEQASNDPVGADDNRSPGVSVN EWVRASVGASLRLRKAKKAEGQANTPRQRQKSLVALMSPTGTIDINNQTEREEKIYKGPFDYFGERSSLFG TPRNATCMALCVTSFLVLT SARFE AILDEF PHERNSNSVSAWVMT RPSLPP DERPIDN*
>scaffold_7_R68 1	MKACVYLPSSLFLPCPSTSAAEPTDPDEDNSTPHLDALERLLDRERARQRLVNRLRRPDQEIDLHDSIRWH VRYGPERIPSPQQSPPTPDFIETYGDSEYAVSDDNSSDSGTESEQC*
>scaffold_3_F35 36	MALVVFIRLLVCLQIGLDEEAIEDHPTRKTRRQLQAGPHEASSEERVTEETHPEPMARHLRQFPTQHFDIV QSIFGRIGQQIRRAGIDS*
>scaffold_61_F2 09	MRLLFLLVAIATFLIASEAFSTTGDSNQIYNVDSPVGPRQRFLRAHAEVDELEKEMKMMKMMKRKVTKEDF AKTLKITDQIDDIINKHAPGMHEFMQTPKYQORYSNYMFLNDMAKKPEYAALVEEIKASRAQVALKFRKP TTSQRNWQIIASFALKGKRIGK*
>scaffold_23_R4 90	MQRMRSTPFFLMITTIGDAQGEVEWMIPADNSPLIKSDNFNSRNCATRTRWATGVALPTSMSQTAPRA AGGRAFPFPIRRCLRFQMICLNRSCSGVLPASSIWISILTSSVVALTSGSTAI*
>scaffold_25_R1 22	MYRILLTTFALLCGFSNAVNSDARLLRVDPTEEEKGGFTTYVKDSLTKWRINSKIKSWVKNNKTDEYVLKKLG LSTLTGKDVLNAPKYSQFQDFKVGWMWLKEATPTTVFNTLGLNKVEGAVENADDFTYVKYVITLGEKADN YPLSQWPRLFGGGSLEQLELKRKILRLQKRNVLEINFMLG*
>scaffold_640_R 2	MRICLCYLLSLTAVVSGSSVNLRASQKIQSLNVVQARMSGRELREDLNTDEATEERVNFGFLKADRLKSFAK QMDETHIRTKSMNHKRN*
>scaffold_38_F1 173	MRSYCVLLVYTAVMSTAGTESTAMDSTQPSTQYDIVLHVRLTRTGEYTVSEERAAGASAVESLGKATTSDV DVETWLKRGDTANKVFALKLQKAGDNLLDNPQLSTLMRYLRLFNEANPTKKTSLIATVTKHYRNHGLTKIE AGLASTSKKTVATAKHLQTEQIHYWMAHGRSPESVFGVLIQYTKDALRRSLLKLDKLLPFTWISSDLFEKPG KTWIRYLDEFNVRNPQKTTLISLKRNYNDKTLANMLIAANKAESTSSIAKRIQSEQLQWLSKGKKTSDLFT MFQLQKAGATLFQNPLFPAWIKYADDFRVIHRDTQLETMTLMKHFDVVLAKMIMTAYDVPSTKSLASR LHAELLRGWQWRQETPDDLYILLKLC*
>scaffold_25_F1 656	MRLSLLVAAAALVANSDAAPQSTSSLTKFSTDVAPVRSLRGASKTKTEVDEDSDFDPEEEERGISMGYGKLDK VDDILKQLNLADDTVLKLIAGKGKGGVEQLITTDKGVKTLVITSKKTGKVTFTNIDENIRKIESTPAIKKQISQ WKKLQLSPLAVSRDLRKKGIPRTTDNVWEAFLRYSATSGRKFNSLRPTGL*
>scaffold_77_R2 49	MWMTQYHLLVWAPALSI LT SARDEM GT QEQTAMILKEYLMAKSYTVDDLKD KPSRC RRLDGKTVQDHGP EAYQDVARALRVLQVWEK VYPKASIE DELAFL DRLRPQLNSTKS*
>scaffold_1_R40 7	MRTATAFATVLALIVATNGAHVSPNTPALRGLRPTADTPSVEDDKEDRKDHVVKKVKKIAIPVVPVVEVPQ YIPVPSVPSMVASSNNAVGPSTNVAGPGAAAPGAPGAPGAVTPAPTTLNGRPAATPAATSTTRSRPTPA PTNFAGARPSGAQLPAAPVQAGAPGMAGFPNSVIGDNTNNNAFGAGIGAAGGFPMTGNGLAGGF GG NPMMSGFGGANGMGSGGMFGGAGPAMGGRLNGNDFGGNFGQSMGFGM QGGNGLGGFGGQGMMA NFGQTGNNGFAGGQALTQGGFGGGNSFGGGFSRQRR*
>scaffold_43_F1 201	MRWLCIKQQVWLQSHCLMVSPFLLVCCPCLSDNSVFTSGWTMMLLGSQLMWKSSLQNLEWLAAISSACL QMRP RRLRMQMMLFALVSTSLLS*
>scaffold_6_R23 12	MILLRAIVIAVAFVTISLVTATGSEASSITVVDVADRSTQVKRSLLRNLESVDEDRGPIAGLEKVDDILKTEKAS RKAGKVP SGLKNLLRNIDEFAEHSKLAKKSLGLKLYDAGLEKMSLSTLRQLDDIEVKRVDIKNGITGNKDTP

	GGMRRKMDHVGVDVAPAKYLTSHIGRGDQLYGADGSRLSSAVSRPAEQGGGKVLLISSSKPEKGDWLLPKGGWDKGEDIETAALREVMEEGGVRPVFFLGLSYQYSN*
>scaffold_405_F4	MRLAQILVIAAASLLFAGETVAVATSQNQVKISKVAQSSPSQRLLRSNKYPITEEEDESEDPEVDFFERGEEDLEE RSPLSSAIVSKLDIISRWGTTYARVAMGQSTISQNKKIDALLSLRDAYVSGSKKAKAAKIAVLRAND*
>scaffold_9_F155	MPTLLVVIAVSTAMAIDMTTRLRIRISICWYTRSIEFLSLPLRWCDSRLLINNSILHVHRDRFIRCHHGRMV WVILFLFSYRTVSHWRSSGRRVSRRVFIFTMRNSIHR*
>scaffold_42_R191	MSKVFLLLVLSVFAVLSCDALSAVGSKLSSLKTDELNAQPIDAKRMLRAQEPTNAADEERGMTELANKFK AWAAAIAKTWVTNSKLVQSMNNKLASLTQKGRVGQIEKLLQDNVNVLVQNKVKPDELFLALKLDPKLKL IADAPAAWANNPGLSMFYQYATYYAKMTTKA*
>scaffold_73_R270	MRLLYLAGVAILAFIAGDATGAKVLVPDDSDHNRAQAASAVSTTRLLRTSVIDEERAGGISASASDKLAKLFKSSKVTDEQLQQWLNGKTAESVFYRMNLENTLYTRVFKSPQFPRWLQYADDLSASGKGSPAISVLSTKYG DEKLYQMIGWAKKESSTKALGTRLQTEQLEHWVKVGKDPEEVFKLYDLNYAGWRFLNSQFSAWTKYVDD LNAKNEGAFVIPIITLRKYFSDDDFKIALAAKRSGDTEAMGTLKEDAFVQFWVHRKDTPDNLVLEGLKQST KTLLESPLLSLLTKYTEAYNVRFATKTTVIETLVRAFDDETVARMLLAGREKSTTKIAKQFQADQLEMWLNS GQSVDVYKLLNLPSSRDLHGDFGGEKLFDTWLTFMNAVSIKTPEKTSIAFTTALPTFKNRPMQMILEAANK FSSMEKAATKLQLEKAQSIFSTGVSPYKAFKMVALDNVGDSVLSSPLFNKWMLYVEEFNKKNPGEESWFLP LRGNYQGHGLDRVIDKAMKDPSTVKLAQVQKENMKEWLVRWKYSPSMALRELHLNKAGEKVFSAKFE WVKYLDDWNQAYPSKETMIDGFRGNYHDLQVPMIAAAEKVPSTKKLASELKGALVDKWVAEKTLAYV KSWLGISSSDDMLERFTAQLNSV*
>scaffold_300_F13	MTGSTPDHWLCFVISSLCATTGIKALHCCVSHSLTFQLDFQRLRLQESEQSMGVTTAPLKSPMGSPAVPY EPVQTMWEARPRKYKSLFEV
>scaffold_39_R1000	MMLVLLAICETTIHVAAGTSLFESDFAFGVDSTSACAIRPFLDPDKHLLYSNLTNNLIALARCTVLERLEVS CKDMQDRNKFSLRLRSYLLHVPVALYSTERTSYHQGRNHIHYHGGL*
>scaffold_7_F1494	MSSKAGLISCALWLLLVSTLGVDAHKRLLSEVSHDNLEDRVLDNWKKLSSLVKVGSRTKADKLYAKHVG KITGNFFDTNAFAAWFVAVQMAYAKTPAKAVKDMVSSLTARYGDQALAKMLATTEDDKFIREMKAQLDN WQKDKRTVGSVYKLLKLDKEQDELLQSPLIATWIAYATKLDNEPPLGAVFSTLKTQYNGKDFATMLLNVKDT DDSFVVAEKLETLLMKSQREDKSVDVYKLLNLDNEGDLFFQHPLIDTIRYATVVDKDSFSGVFSLLQAR YNEEKMTDMFMTMRDWPRNI LTDQLEDLLKWTQRQEKTMDDVFKLLKLEQQGDSLFSKLLSTWVSY VAKVETNPYNVVFSKLKSTYGEETLTSMIIQARDMPTADYVGTGRRIRESSVQGLGEGQVHCP*
>scaffold_17_F940	MQSVTSLACVNGAFLSSVAVIAGSDRSRLVLDVGAGGGGRTMRVVRDAHSRAAHTVALPRPTCYTHPSNF YDLLSSAPDSTTHLWDIRADNCVMRFCEHVNVRVHTLGVAFSPCMRYVATGSEDRAAYIYDIRTGRRLVKLKGHTDVVTSAFSPLHPQLATAACDGTVRFYSSTRSD*
>scaffold_21_F492	MSSRLNLLFTGSFLLMARKEIVRIFFHLLLVCNSAIHATVSSSGKSITATRHLRRQAKPSVSRARVDG*
>scaffold_2_R2430	MGLPLMSSVVFPSLAAGLGGSHATTWNLADFVSLTSELPISLYGVMAGDATHLHHVLAHIGFALFSGV DVPRVTRPSIRTWSQDLVGEIPLLRRRSHALPTEVLEQVTGSNERRSALGNELLRVVEPFVSELDFLVRATSAS RAAAFGTNSATFLRTMTQIVRQLRVYARGESTESEDDSDERLKRLRGLLVWLMNENMARFVIDSLVCWAEGDNSSIRRGRTRQREENSTDSPAIKRQRE*
>scaffold_30_R1833	MSDSPAKISAVCTALLLLAPFQWQSTYIPLLPSGLDFLHSPVPFLVGCHSLSETAEWSDVCFYDIRDRIAVP AVTRHLPSSIPNGVELCRLRKARERFCSLRPTSKPWYELSEEQDTIITLTMQEAEIFLRDLGFDISSLQDLASI SGKSSVVLREHEVYSRIR*
>scaffold_40_F1027	MVVKKVGRLHLFSLLELGIDCFADFHCRLLTSRVHSIASSASLRPSRWLQSSRSSLTWTCRLTRSSGSTRTR TCMTNCNRCWRTIERRSRTLCVRRSRYHCRLRDSAVTLV*
>scaffold_5_F2581	MRFHYLVIAAFVLSISDTFSTSATNTVPSRGLRRIFDEERAGGAISVSTSEKLATLFKSSKATDKQLQKWLQKRKPAEDVFYRMNLAKTGTGIFDNPLFIKWVQYADDSATTSGKGKSAISTLTAQYGDSDLYKMLNVAKQDSK SKELASRLQSDQLEHWVTIGKDPSEVFKLYDLNHVGGSLLRNPOYNSWTKYVDDLNAKHGGEVSMIPTLRK YNYDEDLFAIVGAKSVDALKSAGVKLENQFVQYWINDKQTPVKLAELQLGATPKTLESPLFSLLAKYTDVY NVKFPQSKTTMIETFTQAFGIEKVKMVAAKETEGKAKKIA TELEAAQMWMIRSSKSVDEVYNLLKLPPK TLVIDLGSSPLFSTWIAYMKLSIKNGDEMLQLIKTLMQFADPMMQLLQAMEKFPNIGSTATSLQLRKAD DIFATGVTPFRAFKMTALDTVGSVLSSPVFTKWMMSYVDDFNKNPTEESWFVSLRSTYEGDFMDKLIET RKSPKTVKIANTVESERMKDWLTRQKAPEHVFHFLLNKGGEAFSSPNFQLWAKYLDDFNLQYPGEKTTM IDSIRANYRDIELMPILNEAAKIPSTEKLANKLQNALRDKWVDEKVTVTQLKGLFGHMPPSSNDWIQKYAEKL

	NKLS*
>scaffold_39_F7 0	MRLHCFLAVATTAVLNNNGITTEASSLRKVPASAPIDSINAVQPETRRLRSAETVYRHEDSYKRRPFIEEKLH KALTNPKKTKRLYARWYKSGFTKKQVAKGLDQSENRELDVTENLAKGYAKYVDARRTQQQSV*
>scaffold_48_F1 199	MDTSGACVLAVLLFDPNTDTPQKIVLNVGDCRAIIREAPESSPTGTLKGKAASGMTFALSEDHCAANTKE RMRALRSGAYIQNNRIAGVLEPFRTIGDIDVKGPDMKDWWIPTPEIHQSELLVGRSILVIATDGVWTVLN RAMAHAVKELNGGRSSAESAAQAIKEAREFGSSDDITVVVSV*
>scaffold_77_R1 78	MLLCVFTVIAIAVSDTTATNAIDDDGVLQTTSRYLGFNQEERGRVGGPATDGTDKVSDWWPNFKAW FKRTFYFWKKKETTRRLRDQADY*
>scaffold_15_R3 220	MNNLGRSSSVFTAFFESPCASFNSINAASSASLPLAVTASCPNLENITRKLRFNAKMYSPLLERTPSTA KI TADLDWITLQQ*
>scaffold_18_F2 149	MRPNLVVLAALIALVSRCTAVSAASSTNQINLSESQNQALDTLQNHAGDGNRFLRSANIVDDRVDDIDSDDAI DNDDEERGEKNWPKKFAEWADGKTAEDIYKKYALSHVMRYAYRTGSIQWLKKNEYRKWAAYSAYRKD RGFD*
>scaffold_9_R77 1	MRGSQSSQQLCMVLLIRILIVRQLRSLAHQEALLPRAHETSFLEHPARVRHRIVAPHSPQAVGYFGYE PLEW MTVLWEIRSV*
>scaffold_42_R4 07	MRIYFVLLLAVTALIAAVGSTANLRSQTIQSGDVVQDNTSGRELRGDLIADEATEERMNFGLKPDPLKS F AKQQMKYVYNERIFNDLKKFEDPDALYTTNLSKIKNRNSNVKTSRYNLYENFLISYLDKYPNWRSTLK*
>scaffold_5_F20 28	MRLSYTLLIAAVFVSSGNAVATTNGRTTELSAMASPNAVASVDTAVGGEKRSRLRYHNNEGLED ESDEALLEE EERKYTNMFSTKLDEMNGTKMMSRFRWKARGYNTYNLP AVTQKDKYTWRQKYRDFLYHN*
>scaffold_73_F9 5	MRCFYLAVALAFVGCGDTKTTVLASDFSNQH RDHSSTISITRFLRTADASTGDED RVGGVISGTDKIAKLF KSSKVTDEQLQRWLSKGKPAESVFYHGLGEDSLYEVI*
>scaffold_42_F4 1	MANKHDACQSKLLFLFLFSQLPLSAQSPSCPTESPLPPPSAPDWQAAPPFWPLFSIYDV VTSSPPAHWT SRTLPSATQPQP*
>scaffold_5_R21 32	MRLSYVVI PAIAVTFASSGNALAAADGSNTGLAITSPNVVASIDTAVGGEKRSRLYHTNEDLEDDSDDEGL DDAEEERRGNMFSTKLDEM LDGTQLMSRFKKWQELKYNM YLPDTILASKYDELKMYRRFLYNN*
>scaffold_44_F5 78	MRFSYV LAAAALLATANAIEADP RTSLRAHK SHEKSQDE RAFTYTFN SLPEQFQR MRKE PWYLR RIFRSW RSGM GTSDE AVAY MRSQ GLSQ KAIDQ FEDAY IKYRA HKL AKGK*
>scaffold_6_R32 03	MRLAI KTVAL AAAIL ATSTE AAKAV QTGG DVHV VQSS HILP GENK RLLR SEH DEG KL LED DE GED DE EER KY G
>scaffold_11_F1 232	MKMTALL TVMLAF VSADNS PRQLR STLTT SIQD TDGN RLLR SEGL ISNLL GGLN SLLNN QQST NP PTT ST DPTT SEQNP DEID LLRS RRVS*
>scaffold_31_F1 104	MRWYRV LLIAV VSF VISA EQSE SSH IRSL RGN AVN YSFK D EDGN ADER AGE KEV KKLT SSK TM KKL RS V TKA ANS FSE K IS PAL PI AK L Q V WS NT G K SV Q F V R Q EL G L T D A E A A L K K T Q S F K Y Y D D F V T S Q L P I W A K D L D K L T G K E I L T H P N Y E Y Y K Y F V K N R L K A V A T E G T S L D D V A R V L A R V K F A R E A V N F L G Q V L K A H P N F K F L E T N F D G S R C E Y E I L S G S K T V R E E I A G S Q A A M A
>scaffold_12_F4 82	MRAHF LLL AAA IFL ATSD VAS AN QAE LSK V A SP Q S I E T A N I A P K R F L R T N K Y E L D E Q E E R G I G A L D D V I V K A G K T K M S N A E L K K L V T P K V L E N A L S D R N M K L F K D L Y A G K V S L R F A K V M K N N P N K E E V L K S Y R W F R N Y H I S Q A K T A V
>scaffold_30_R1 705	MV SA A L A S S G S L G S L V A E P L L V V L V S R S D S V A L D V V D L P D P D P E L L D V V P P L V L A L S S S P L V T A M I P T A A P M T R I P T M T R L R A A T I V F
>scaffold_5_F17 18	MQLRN LVFL ALFAG VVA AVQ EEQ PQQ KTHL RG LLD KIV SSD P T P A P V R V L A S W E D G K V G E A R K L P I A E K I V G G E T V V M F N D R R L S T A D L E S A V V N F D V R S N P C V S C G E L T K W Y L G S K E Q C F D L R I A P K S A D Q I E N G L I R G V S A A Y S G L R P V A K N L Y L K F D E T N F I D G S R C E Y E I L S G S K T V R E E I A G S Q A A M A
>scaffold_25_F1 308	MRLSF L L A A A M A T V Y C A T C N A T V D S D Q N K V S M V Q S R L N G Q A D G T R F L R T H H E N E Q E S D R E E R D L T D V F E T K K A A V K K L A K E V M A D N R R A K D V F Q L W K E G Y T L D E L N T F L K S A K Y Q H V Y N Q Y M I F R G Y V *
>scaffold_53_F9 1	M RFF L V A A A A F A S C E A A V E A P N A V Q I D D N I N A G F S R S L R S T D S E D R A I A Q L L S E D R A V A S V K V Q S L Y K A K I T P N Q A K V I L G I S D D M V E L T K T R S L Q R F Y T G Y L S Y T L M E K R K K K E L N Q V K W *
>scaffold_11_R9 69	M I L S T W L R L S A A F L L A A I A C C T T Q L G T P R P S M W I N V T C S N T F G R S V R K I M P D T R M Y V R T G P S G C D A Q Q H K W L I D R I R A H T R R P V S R T K E I S R K H T S R P S E F D G G F H S D Q R M A A P R N G L S V D G S P V R A V A S A P P A P

	SAHIRSGPDTRLPLPACRTGRMSSTWPWTWSCCRGGAPH*
>scaffold_98_R3 65	MRLSFLVLAATFALLSCDTASPMTKDECNPSPNQVLSSETEGRRRLRVHKSSIDDVEERGFNPEKFNRLMNE QSYRRKRFPNVSKYTDRDVYNLLRVDSPNYKRIFNYYQTYLENFAPRLISS*
>scaffold_66_F2 74	MRWTSILLVAAAALTGVLDASTSNTVANSVMESAFSPVIHDHRSRLVDDDTVDDDEDESDEVNEERT WFSNAKQALADKLALTAVASNFLAGKSTDEMGEVLKKLSRDQINTIFDKGEDSIRQILPGFKSGMDSKKFDDLI KALPQEQQGVLLSAYGKYLFNNGLL*
>scaffold_5_F17 02	MTLLAFFLEFGFCVLAMAACKPTFFFSCCTVGPSALLKPKRELVCTRSNATSSSPSSSELLSSSESSLPSSPSSRRA LRFCSSCSSRSVSISSRDVTNLRAVPFSRFLRPLPPAAGPFPPGLPLSLIIVF*
>scaffold_10_F1 990	MRVSILLVVAFAVALNPTAAASDANTVVAVPNVHESIATGRFLRARPVDDYPVKDERDEDDSDEKEE RMFSFFQEKAOTALSAFKKLVSQSGDDLVEAVSGLSKGEFQALFNQGKAHMAKMVPGFYPGMSLGEFGTVV RAAGLSDDMENALMVGYGKYLAHLMD*
>scaffold_26_F3 52	MERVGGLSCIAVAGATVALVEALLSGLEQVRVGHHSREVQQARKAIAYVLGARRVEAQHSSASVRSHGL RPRLKAEEGLANCAARSLRLARSLLLRRGRRNCEPRRERGAERRHCKAAGWDQ*
>scaffold_96_F2 12	MRLSHFFVVAAAFLATGAIADSEPNHRNLRKHHDTLVDEERGIPKILDQNIPIERLNSLLKKMDQRKKAELA KLIKLYESSPKFGRTKPSNS*
>scaffold_92_F1 4	MRFSQLVVAAVSLLFASETAAVATSQNQAKISKVSQSSPSQRLLRSNKYPIKEEDESEDSDVWEERGFATPD EEEELERSPLSDLATVGKLNKIAKGWGTTYGKVAMGQSHISEAKAKALLRDAYISGDKSAKAAARMAILNA NHR*
>scaffold_853_R 1	MQRMMSTPFFLIMTTIGEAQGEVEWMIPADNSPLIKSDNFNSRNCGATRTRWATGVALPTSMSQTAPRA AGGRAFPFPIRRCLRFQMICLNRSCSGVLPASSIWISILTSSVALTSGSMAI*
>scaffold_42_R6 7	MRCFCITLFVATTILASSGNISAATVLNRSVTTSIDAVQPIEAQTNNSRSLRRNNYKDEEVAAEERSGIESLA AQLNKKLLPTIRQVAHLDLGRAALTQQMSLSFDERQAIQALLQLSKKDRKAVLMLIK*
>scaffold_7_R19 13	MRLLLWTLLVILVTILSSCDAAVNENKSLQRKLYTKVASHALAADDGFEHKRALRGASNGVTEARAATVST KFGSRLMAFFRSIKDKYLAWEKILVPGFEKMAKKGTTYTQVREDFRTRLNWGLWGTPSGFKRYAKLYRT WLEKNHYSQLAV*
>scaffold_8_R16 99	MRVLGAIFSALFLAAGIDVGSSLPNSGVAFPVNPHSLAVETVESKRLRSYEADSEDEERAGGQLGLVDKFAA KVMQKLYKNPSDVFKRLKLDVNLENNKVFEGWLVYVNKFQVKGVENFPDQALFNVIHQSQYYQRDLVP LFQSLTHVQGMKDARTMQFKLFEATPATRTLMNKAWLEGFDTSDDVFHILKLQDGVDHSKDLIQWLK FSDMYKKLPTSQSTS WLDELNVLTKPNQQETKFGLLFQALKEEKGMETIAGKMESQLFERWMKMDSM TPDKVGGMLGGSATTNWKRIFEKLEFTDDRYIFLKAYTEAYAANRGANVLKSVEKLFNAEKPVAALERAIK*
>scaffold_87_R4 30	MRLSQVLVIAAATFLFASDTVIA TSQNQANISKIEQSIPSQRLLRSNKYAVKEEDESEDSDVFEERGFATPDEE DLEERSPLSAAIVEKLDIASRWGTTWGRVAMGHSSISDDKIKALIAMRDAFISGKKSIRDVARAMILMANK S*
>scaffold_15_R1 164	MRLTSIAFVAASIFVCFQEASA VSDSKGPESRVQTANSNRFLRIDQDEERAGPQPSKSLLEKFTSKVTATK LKRSKAVADLTKLDDVVYLKATPGYNVPLFQRIEKMGFPDGMLLKMRERGKIDQTLKHYTNWKGKYPT WTSNPSV*
>scaffold_17_F1 758	MPVFDALLTILRAIGCDRV CANQKRRMADIVKRMEDGAFRTGKAWRILRKL NKS EMKND AEWSTFLEL SNVLDPRAFNSRRGRALMNEAEQVISVQQL EAA VRGP IV DVK PEGEN VADELW DIAEQ IVE FIER KA APVA EPQDTTNDPTFSFDVVLVTLEDYLKVEARTAPS VNGMSDRE THIIDE GIELS VT EQLS RMCS LKEE YVIS SDILEKGFAELSAI ER LD TVSI QVS FLL GMQM RL KSC IDH CESAK SE FERR IKR VFN QL AGKG QD MD HA ASLI VLSAFC PGQVIREC IRG ART GVLH HDL SKV LQSS P LLE WREP SED GSG T LFE EL QM T VLD ISRN QSS FDRE SHNV SFLL SV GIDN ST SSI QSS LMTV SKL ITVC IN PVW C CPD QS VEM QL NL LTI QQL FHF VATS ADIT DTLQISFNLA FR AL SS A NSN DTKV GALI RE KV LLL K SIM E LMP DP V STIR LE QFD SPN APL W TLIS LFS DDLG NEELCKGLEDVTA VE AYM QTSS VDEADPLPLPIVISA IQ ALL W GLL W DS ILSEN VVESTKAETW RLL II AS FE FCGSESPDETIGGSSLIESAVA EMMI LEC GN VL FR ALL C NI IP FL EYEL TEN F QEEK LLIP QWA IDK VSEN PEIKE QIPCR LVSSHVIMRYVAKCW CLSGVANRQLNAQSDMLV LEAL THV VTA HDQ AIT AS KES LSGT LFCI QWL CF LVSAARELHLDL STW ST VRT QLE LSFL RLLN QL EHLKA IS EAE AKF SQV FVA AWL GYL P DDQF VQF KYI STK RSN*
>scaffold_22_R9 74	MILTTNLATWIFLTWAATSSMNLSKCQCTCLNK CEN PRPLRGHLLSSW VSGKQSGWPYQANLLHVHL EH PRQIRT VGSRK*
>scaffold_65_F3 81	MRIPCSLLVVI ALLSTTNAI STEADAGR RSLRSMKAKVQDNQVEERGGTGSSLKAFVHDFDNLLDNI F PADFKRMTKEPEF LRHMM ASWQMGFMSVDDIVLYMTRLN M SEKAINQFKLAYTAYLEYLKAVAKA EKAA

	KLAKASLN*
>scaffold_25_F1 318	MRLSFLLAAMATVYCATCNATADSDQNKVSMVQSRLNGQADGTRFLRTHHENEQESDREERDLTDFET KKAALKLAKEVMADNRRAKDVFQLWKEKGYTDELNTFLKSAYQHVYNQYMFRGYV*
>scaffold_184_R 6	MRLSYFVLVAACIFACGEEVAVKDSQGIAVSHTDSLRGKDNRFLGRQDVADEESDDEKNTNVEYEERAL PKLDVAAEKLMTAAKLTKSLSNLAKMDEAYHRAVVTNNFTLERIEKMGYNPDKMFLKMKEMGKN APLTPQKVLEYYSMFWKIKYPSWVSEVPRMF*
>scaffold_1_F40 0	MFVLFNLLSCTRLELSFLPPRVAAAIAAACADGGLGVTRHSPSCRLELSVTASFVLLHQLRILRST*
>scaffold_6_R20 44	MRVLLVALAVFASSVNTGVAGIQTKRSLRQYDFKSLTVADSKEERNFVDKVDIARVTDDFTVKVKIPTN MNAAVEVTTMADDIAAVAKTVAKTYPEGLSKGTLAQIKEVEQLRLKDIATYTKTGDGMRRKISPPGMKI APKKYLESHVGRNMQLYGDDGSRLMSSAVRSRAKDGDDVLLISSNPQKNDWLLPKGGWDKGEDIQS AALREVVEEGGCFFIYLM*
>scaffold_60_R8 42	MFLSPTFVMFLVVPIVMKAEDWSTEGISHSLPNARSRLQERVVDNSDVDSKAQRISGHVATVLGHSAAMD EVGVVVGMLLG*
>scaffold_8_F32 65	MPQTPRCLLKVLFADVVHCAALYDLLGHSNGSDLKYL MILGH PADLQLSGIQFVRLRGQSPVALPVAHQL ST*
>scaffold_10_R1 425	MRLSVILLVVAFAVALNPTAAASDANTVVAAPNVHESIATGRFLRARPVDDYPVKDERDEDDSDDEKDEER MFSSFQEKAOTALSAFKKLVSQSGDDLVEAVSGLSKGEFQALFNQGKAHMAKMVPFGYPMGLGEFGTVVR AAGLSDDMENALMVGYGKYLAHMD*
>scaffold_81_F2 33	MRFHALVLLSTALLATDAFKLTADVTAPQSRLRKSEPKTNEERVFEGLTSTKLQKYNLLWSKLQLGDDLSAV LKSPDVAKIAKFNLNRAPGSQVSMIQRITAKYGDVVARTLVSIERDASDNPILLTMVWQLREDQIANWLKN SETVPGVVSKLKGTDSEIFRSRALDVEDFIKKYNTARNGDESLLKTLLTIYGGESELVKMIAKTRVLGRPPYSN HPASIEKANNIESSQLIWKWSQNLPDFRVMNKLNFDDVSMALSAGKVGVLKYSDSTS AFRRSLAKYGEA EVAIAFAKAEGFLPEATALYRMQMNGWLSKGDTAGRVSILKMKDTRFYKLDALETYVKFLKTKNPDDV TDVFVKLKKFGAGENKLALAIVRPPEITGYHKSLFQDWARDLPLSVAVKVFKMSEADAAKYSDEL KPIIKQYTKFFKETVDPPEMPAVRVGRS*
>scaffold_6_R31 91	MRLFIKNLVALAVVLLATSTEATKTIQTSGKVDVVQSSQILTGENKRLRSEHDEGELLGDDEEGLLEDDEER KYGANLFSAEKMKMLGDTDTEKKEEKSQQ*
>scaffold_67_R4 06	MWLLVVTLVFGSIFTYADNSGSDSDNSIDDDYVESTYYSSNAGLFVGMLVAANTIMHLRCLRTAQLLVAE LVNAGVDRLOQPPVVMLATTNPTSAV*
>scaffold_636_R 1	MQRMMSTPFLLMITTIGDAQGEVEWMIPADNSPLIKSDFSRNCGAARRTRWATGVALPTSMSQTAPRA AGGRAFPFPIRRCLRFQMICLNRSCISGVLPASSIW MSSILTSSVALSGSMAI*
>scaffold_12_F1 261	MGVWDWILCFVCEEVGAACPCLTPSIVGRVLLHGKTKKIVSKPPTSNSQTMFCSFILDLIWTTYTTMSKPIQ SDS DLDALRKLTQLRVAKLVTDRDLKQTDRKNATQAVMRKWLPLSTTVLKMVTRVLPSPIAAQVKRAERL CTISSEQLEQSPQHAQVFRSLQSCQTSEDAPLVYICKVISVEANVLSDYHQSGLAATDEVYVGVRVSGVLR EGQPVYVMDPKFQGVSGDMDVDTIDPSTVKHVARIDSGLIKTYMMMGRLHKLDRVPAGNIVGVLQE HVLKTATLSSTLACPSLT KMPYQAKPIVRVAVEPEDPRNFGALEAGLQRLYRSDPTVEHVQETGEHVIVALG ELHLCRICKDLKERFAKVAVQVSEPLVGFRESIVDTISSLQENIVFKELLNPDTKDDSEKDEKIRKLENSIVTFQ KVALGTTPDGTTLKRALPLPLEAKLLEESASLLKRIA VSKKANDDMKDKN TVVELEAAASEDVSFKKKLEK SLOQSESSFLKALPLDQIWC GPRRVGPMLINSIPTYRATGCLFPSDSAETHDEKDEKIRKLENSIVTFQ MASSAGPLCDEPVWGVAFIIEDVVVFHDEKSEEDKSEEKAEMSKYGPLSGQV SIMRTCLMSFVKQPVRLV EAVYECTVQCQAEQLGKLYSVISKRRGDIYSEELSDGTALFTVKAHLPVVESFGFATD LIQTS GAASNPQLIFS HWSIIEMDPFFQPQTEEEREDYGERVYEHNYVRRYIEAVRKRKGLSRDEKVVVHAEKQRTLKR*
>scaffold_81_F9 0	MRFPLLLLVTFLVNGKADSQSTELRLRDGNSPREEERALPPPKVSLFADWLKATGLKVS DKVRARYWL WRKQSAEDVFRLKLDGGLKLLGSRKFNTWTSFVN IYNNKKNPNEKVTMDGILSKTYGDLELAKALEVAMGS MVKSERKMGT TLSLQQREGWQAAGKTADDV FILLKLDKVGA DLFTTPQLNSWYKYVSMQADSKSLMASV LRNHYSDETLSKIFREAKPEIKRMRIIRVQLETAVAKSKPKQILSPEEYF KMLK LDDGVDFKFLASSNLDTWI KYV ARYNTKNPGQDVSTIKILT KFYGDEELAKVLEAGK MATTEKVATEL QSAQFNQWLSLYSTDDVFRLLKLDN DLD SLLTNPNLITWITYLGQFNANNPGKGTTMIK TFKYGEIPLAKMLESSL VPKTEKVATQFQTAQFKQW LRDGKKPAEIWKALKMEKATWMQNPDAQVWYKYKD WYKLNKPQ*
>scaffold_6_R31 37	MPLLILIFIASILATTIYVNTKLLNIPGNLSLGVSTY NHSHLDWKM DTA AFETSVG TSIHPTSSFY RRGKTFDRSA SRERAVMVCLHDAMLNMGSL RELRCLGNHELIQVYHCGNELSDRS VELLFS LDNRVELDVCS DLSRGII

	SKEMATKFRSW/WIKPLAMYHTDVRHVMLLDVDDVILKDPAVVRTLDGYKNTGTTFFYDRVISNKRFLTGN TGEMYVHKLLRTFNYTRFNVSEGPNPSQHMLNTFAFNGKSIHEMDSSMVILDKQRAGKVVMDILLWFITEE RFRFTYSWGDKETFWLAFEMARPVYFFSPWGVSVDSMPNEDLKNHPDTLCGSILQFMPVKGMNDTAEV LYMNGKALIDPYPQGIGFIRKAQQNNLFNTIPTQMTPPQKRRQINTKAYPGKKFNTECLVGMGAVPLPTHFS PNLRLRRVHFLGLAMGVLSLDHCETY*
>scaffold_2_R11 14	MPVCYFMFLCVSYFASSSCQIKQHVAFIGDLADAFVPVEGGVLVQRRRNLLTQGLVARQVHTHELEVPAH VARYLRHEEVSRLDADLDAREGAMYDVRREHETELHELREQTCREPHEEQLQLDVALWPLQ*
>scaffold_5_F94 7	MRTTFLWLLLALVLCVCAAEKPTEPTSSANARDNDPVVQEIRGLRNSGMKLNDAKDFKGAIKEKLREAITLE HNRVFGGEGRHAITDPSEISQDAALYAQILNDYGSVLIRAKQYDEAIEVLEDSTVMVEKIYGDSHPSLGLSLRS ADAYMAKEEYKLAIKKYKTLRKHVKKGLETHEAYIEASLRIAEGYKKLGNNKKNLKVLRDAVKAQGTEINGLT TGIAELYMELSTAHTAVGEIDDALRAAETASAIFLQRDGEETLSYAFSLNALAGVKMRQQVDEAVKLEHAH KIAVKIYGENDPITQASAKTLKEVKEYRLDMHAQKDEL*
>scaffold_48_R1 407	MRALCFIFSVLLLLEIDASSVFNSGAAVTHPHLELTQVTPTRLLRTSHEEENDERAGASVLDGIVAKAMQLI NKNPEDVFKKLANTNLQNNAVFEQWLQYVYKFRAAKGEDKFNNHRLFNSQNHPPDDLVPLFQSLT HVQGMEDLARTMQLKLFESEGYQSTRNLMNKAWLQGLDTPDDVFHILLEKNALESQQLPQWLKFAEMY KTQNKISSWENELSLLKTPHEKETQFGLLFQSLKKTEGMETIAAKMEAQLTRWIKTDTMPDKVGVVLAS PTNTNWKRIFEFLPVTPEPRHLLESYTVAЯASRGKVLKSVEKLFANNQPVAALERAIV*
>scaffold_59_F7 11	MRLTYIMIVATAAILACCDGASAVSDSKRLTVPLDRVQTGNNAFRFLRVHHDEEEEEEEERTGPQFIKSLSEK FAKAAETGNFAKTVEKLRSKSLNGISKVDDVVVPGRAKTLTEVLKIDDVAYLKAAGKWPKNLDDIQEAKLV REISTLDDQATLKLITKENAAEFKKIEDMGFPDGMLKRMKQLSKNDKDYFDLVLKHYTQYYMAKHPTWA SSL*
>scaffold_29_F1 645	MRVFSILLVAAATLVASASAESeskQRDVsPAAPQWRTIAENEVPTKRNLRKKEIEERAITAISLFDVVKA KGLQALPYPELANLDSKLRTQYLKLLVNNLDRKQIVEITGQVPHYILTHGDKKARRIVQYNKWIWRKEPD WVLKNYPAFFKGYEFFNNRFTRGYKA*
>scaffold_2_R17 01	MRFSHFVFIAAIILSSENIAIDTPVEGQALMTETDAETPVRALSSNNDKRFLRSYKEEEAYLTEDKYDEEKKR EES*
>scaffold_29_R9 98	MRLLYIAVVAILASTNVPAAADAEVSSFTTERPHWDRSLTAATEGDNDGKRFRLRKRTTTNETIRERSKQ GSENTVSYCTACLHYP*
>scaffold_9_F34 50	MRFSHVLLATAALISSCNATAIVPGEGQVTSADVAVPVRALETRNGKRSRYYDAEEEDETDKYDQKNGK YDDDEERVMAAQIAKWTAKAGEWEQGKTPAYIKDKLTFANGVMSDKNKEYRLFLATWGRAHPNEL GRRYLRPEAESSPRSFRKLIVFISNHLLDVLFCVTR*
>scaffold_6_F30 32	MSAFLVAFRCCHVVFAFWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV*
>scaffold_32_F2 41	MSRAGSPVALVPPKTVLIVNVLPVFERSETSASGWSVSWATGATAMLYRTLVDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVSAWCDPVSWNPAACQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRRLRTALVTMYLHPFPSSEIFRCLAMRTEILRAMLCADHGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCHVEPDVTHLHEMLDNVAEVNGAALEFAAHVKNAIMAGRDSSEQRRIVIGSVDRLEGTLAI PLKLRAFDRLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEINRRYAPPGVVVYFEEKRKTTAE RVLWRMCDIYLDTCVRGGLSLPFYMAQHRNIQSAGTGWSSPRGTIDVGGLDRSGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNKFVVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFDFRVMRFESGVFLDVKVLCANTSRLFIFDYGGLSSTANILDEEGARFSHRPG CVEQSAENADSSDRSGGDCTRRSAARYVDGKVRKPISDETRASLRTLCVDPNCIVFTSNTQRGALEDQFDSI PNLNIAENGLFIRMGGQTRWECVCEEDKRSFGWKDDVKRVMEAYAARTNGSFLVENAASLLYDYNRNSDQ EYGEIQSLELCAQLRKIVEVCIKVILSSR*
>scaffold_399_R 2	MRLTCILLVAAASLVGLDASAATTGNTVVANAAMVISPLAPESQGRRLRVYDDEDDSADEKDDEEDSA DKVDEERGWLSDKMALTSASKFVGKSTDDEMGEVKKLTPAQIDTIFDKGEDSIQKMLPGFYSGMDFQKFD DLIRALPQEQQAVMLSAYTKYLHQNGRFS*
>scaffold_7_F17 75	MRLNVVWSMLGLVVLVMVTEAFETTKGAKNGVVTDPARRLLGFAADQEERRFGPATGTDHSNSHAW ENFKAWFKETFFFWRKWEQKRLRS*
>scaffold_18_F2 220	MQRMMSTPFFLMITTIGDAQGEVEWMIPADNSPLIKSDNFSRNCRATRRTRWATGVALPTSMSQTAPRA AGGRAFPSPIRRCLRFQMICLNRSICSGVLPASSIWCSILTSSVVALSGSMA*

>scaffold_50_F2 79	MRFLVFLALAIVALVADGEAKDSRLLRARSDEERGGGITVPVADKVAKLQFSKLTDEQLQQWLSAGKTAES VFYRMGLDNTFVTVKVFKSPQFIRWLQYADDLSASGKGTSAISLSTKYGDEQLYKMIGWAKKESSTEALGLRL QSEQLDHWVVKVGKDPDEVFKLYDLNYAGSRILSTSQFSAWTKYVDDLNNAKNEGAFVSIPTLRKHFSDELF HIVLAAKRSGTEAMGMKLEADFVQFWIHRKETPGNVLVGLKKSMETLLESPLLSILTGYTEVNVNYAAK KTTVIETLRTFGDDKVARMLLAGRTESTTAKQFQTDQLEMWLSSGQSVDVYKLKLPYRDLVADIGTN KLFGTWLTFMNAVSLKNPEKMSAIFTRLEPTFGNRPMQMILEAANKLPSMEAATKLQLEKAQRIFSTGESP YQLFKLVALDDVGESVLSPLFKWMLYVEDFNKKNPSDEVSWFVALRAGYQDDKLGPIDKAMKDPKTMK LAKLVEKERMKEWLEWKRPSSMAFRELHLNKAGEKVFSAKFEWLVKYLDWNQAYPSKTTMIDAFVD NYHMSTLVPMLAEEAKVPTKKLASQLKDALKWIAEKESLAYVKSWLKVVPSSDDMLERFTKKLNSA*
>scaffold_7_R19 03	MSVLCLALMILAMVTGAFETS RDVKNGVVT PSSRLLGFTADQEERRFGPATGTDHSNSHTWENFKAWF KKTFFFWRKREQTRRLRN*
>scaffold_24_F3 57	MRLSFLLLVAAVALLSSGTVSAMTNDEPISTPNQVLSSETEGRKLRMHNIASDDTEERGFNAQKFEQLMD DTGYRSKRFA NWVTKRYTDIYNLQISSNPKYKRILNKYQTYIEHFAPGLISP*
>scaffold_17_F3 59	MAANSCAVLVLCGLPAGKTTLVKQLVATACSISSLHERISFDDLYEQHVTAEKGPGEDPEKWKMCQ QDMILKRVSNRLKEQNNDPVHRNECNQLVLLVDDNFQYRSLRKRRFFHLTAKRRRPSGFTLC TTGMPLMTRSE LWFWCALCGCSFRYLSGTECWSKQTRSTQ*
>scaffold_8_R31 93	MRLSFLLPVAMAIIYC ATCNATADSDQNK MSTMVHSLDARLNDEAGGRRFLRVNQEEEDVDAEERGYFEKA AVKKMAKAIMADPSK ADEVYNTWAEGYTL SKMSEYL KAKKYD QVYNGYALHLDI*
>scaffold_83_F3 73	MHPPSLVLT L VLSPAALKSSTL HVAALELQDT VEPSSH LAE PSFSPCR RPLRV ASSTCAKAGADPL LAPNGH LK*
>scaffold_25_F1 083	MRLSFLLPVAMV AIIYC ATCNANVDSNQN KASMLQARLNDEAGGTRLLRVHESDTEERGFLEKA AVKKMA KAI MADPNKA D E V Y K K WADKG Y TL QMSNFL KSKTAG KYDRV Y NGYV I HLDY*
>scaffold_22_F1 823	MRLTFLLFAAMLA F SSTS D AAS VDQTTGVRSLR RYQA EDEERGVTSV STSIDDVLNNL IKQKSSLSKL ISQKK LDNM LTKNFI KLLRSN KSYT D E VFT KATLN KMLTSEKFA EKKF VEWY ALG LTD K LILQ RLNG VGEHFGTLHSQ YVTFINRIHGVA*
>scaffold_157_F 14	MRVASI ALLAVV TALAS VT DSSA ATT GTV LAKV VS NEAAPS VENE HATRFLRKHD HADTEREERNGISLLQ GLKSTFEK VADLPFDRAWHHLQMLNLSWDKREALLKHLRSLAKDREAVLKLIT*
>scaffold_7_F23 10	MISRSFTLITAGVMAS TTLI HASPLQYDPYTPVNISTPLTSSHPAYGAQTEG CIEPIV P EDPNQAKAESMIIQAD IYRK LRS MEDTTNSDIQD LETYFGTKMEVN FQTLKQQYSSGHAPATPWASSY WPTFQDSINYV WKTGEPSA SEKYATAYGLN VTEFKDKISER SGVDSRRSTRCTADSDCKDG SVCGKRD GISSGYCIPGWFGICHAWAPAAI LEPEPQCDVTKNNVTFH VMDIKGLVTSIYDGAEIKTVFTGARFGLDSPAN KDQYGRFTAARRDLGPGYFH IAITNVMAKQNR SFV VDVTAGSEVWNQPVRSFNVQ SMDLVDTRV ASM QYFGVPSYPFNDK MVRLAYVKT TFSWVSE SYK DGPLVSSGRIDRYTESKDYEYL ELDADY NIIGGEWVGQSKDEHPDFLWLPTAKPSASAVTST GLKYADVQELLNLSQSC*
>scaffold_39_R8 34	MPVPLRLSLAGSSCSGAQVLSSLL ALLDASSAQYASLPSFDEL FYPLYLLL HALVKQLED TKVSEVNAISK LH NRLET CWNARRPLRLQT FAPT ILPTFAPQFDEN YTVR KDKTAPKDTAQLKQLQRQV KRARKGAARELRRDA EFIHREKQKEEEARLSAKEEKQKEIRR WLEE QNAT FNQ QV RKG GHMLKGGGSARGPAPRARTPRK*
>scaffold_104_R 20	MRLTSILAAA VV TLT HSA TAFPAV KDTAAIENG AVADIV DSSV TKG GRML RVN YNNDDDD L D KKK KTRN SE DPLD NYDEE VRGIVE ALK LNPV TA AKNSAKL LQSTL RRSR K*
>scaffold_61_R5 21	MKVTKVVVALAACVALWT SPTD SEDIS NILE VSASRHLR QTS AEF AAKPQ ETG KKR D STNPL QRR DQAL VS AHRVYDPVSGLACSLVGE C VACP QSER DESFC RETG YRQ ELD C PRP ND PKDE ALLT K PEDER ETR FK AC SPA DSARPGVAVV K FELL MA A VLA ASV L RERR NH MSSF DLK D PRQ RTG L LGG NSD K SSD*
>scaffold_90_R9 4	MANLYKLFV TLLV L WS GSKV SVCC L TS RFL RTQ NLKTRY LAK STS CFQC GSGH HLHPLR HYAVS VSQL AS LN L RRLACC*
>scaffold_25_R3 32	MYR VLL TVF ALL CGFS NAANSG RLL RVDSA EERGG FT SYL MDSFT KWR INSK IN SWVN KQ KTD EY VLAK LGSL LTG KELV KA AKY POF QDF KVGV WL KEAT PTT SVF STL GLD KV EG AVE K ADD FG TYV KYV M ALGE KAD DYPIT RW REL FGG GSPEQ LK LKR QLL FLA KRNA IDIR IML G*
>scaffold_18_F2 130	MRPNL VVLA A VIAL V SRCTA V SADS STNL NQAL DTLQ NHAGDGNRF LRSANIV DDRV DDIDS DDAID NDE ERGEK LGLRN LPNG MRAER P QTT STK GSH SSL TYA KRTN M A KSDG SITT STIAS GP ITWPS*
>scaffold_74_F6 12	MVFIAPKLT AEQAL VIIS CLLA ARG NRK RFL RTM HF LIASH ASV FETH LLG STQ ATAS TPL T RR HG HAS ILR YLSYDC*
>scaffold_37_F1	MGLLQKVKT MVM IPTV ALSLT ATRIG VPSV LAAR AVPT QTHQ PLLP RRA KLI RE LRQ GIN VSR LLV AKSP RIRL

084	QGHLRAVVRYRTRRMIL*
>scaffold_11_F1 319	MIRGYVTLLATTAILASFGNVSGDLDHTQVVVASLDAVLPTNTNRLRGRKNNEEMEERNGFEALAAKL DKSVPAILKVANLDLGRAALTQQQLRIPFEQRЛАIQLRLSKKDRKAVLLIK*
>scaffold_11_R2 801	MKWCRAASSASLALLACSLAGLQDQAADRWTVHTVQRTRSAEELVHEGTSRWPAGAVASVTCPSDLL ALQCREADGQRELRRDDQDLRRLTRMCMCCADHRIKSGGLFALYMKASVRLTANCdryELDAPLVSTLTC RSVDKDIHLLPDKEPL*
>scaffold_11_F2 657	MQRRMSTPFFLMITTIGDAQGEVEWMIPADNSPLIKSDNFNRNCATRTRWATGVALPTSMSQTAPRA AGGRAFPFPIRRCLRFQMICLNRSICSGVLPSSASSIWMSILTSSVALTSGSMAI*
>scaffold_50_R1 378	MGLFYLVCVAILAFMAGDSTGDRVIVSDGFNQHRENAARASVVSTTRLLTKSVIDEERVGIPVSATDKLAK FLKPSKVTDKQLQEWRNGKTAESVFYRMNLNNPSTKYLFEDELFQTRWLKYADDLSASGKGASAISVLSAKY GDEILYLMIDRAMQEKSALGIRLQADQLAHWVKRKDPDEVFKLYDLNYAGRGLNSQFNAWTKYVDD LSAKNEGAFVSIPIITLRKYSSDDNLKIALAAKEVDETEAMGMKLEAFVQFWIHRKETPDNVLVDLGLKKSTK TLLKNPLLNILTKEYTEAYNVNYPSPRTVIETLRTFSDEVMAKTFLAGRTEYTTKKIAKFQFTDQLEMWLSSG QSVDDVYKLLSLPPRNSLVDFGNQKLFDTWLTFMNAVSINKPDKTSIAFTTLAPTFNDRPMQMILEAAKKFP SMEKAATKLQLEKAQSIFSTGVSPYTAFRMVALDDVGESVSSPLFKWMLRPDRYHLMASHLCCRPAAQ HFLKRPRRRDVCGGNSCVLPYVRGRGVAGVALLCRKPNELQLLLSTTDC*
>scaffold_17_F2 587	MIVYTGLTVTTLVAFVATVCLELTEHTACVLSNDLGISSSTAQVGLSLWSGVHLEQVEQSPFSRQLHGHVVA LDQLEDGRYKGHQVFAIRLLRGALSRNLLHNAQRATDHVLVLVRHTLHQYW*
>scaffold_25_F1 282	MRLSFLPAAMAAIYCATIONATSDQNKMMSMVQSLDARLNGQADGTRFLRAHHESEEESDREERGFTDLFK NEKAAVKKMAKAIMADPSKADEVYNTWAAKKYTLTQLSNFLSKTAGKYDRVYNGYALHLDY*
>scaffold_68_R3 1	MRLNLAVLAALVSRCTAVSAASSTNQINLSNLNQALNTMPNHAGDGPNRLLTAKIVDDSEDDIDSDDDI DRDEEERGGKTWAEKFAKHARGESADDVYQRFALEPVVRQAYKYGQIGRLDDNEYRKWAAYSA
>scaffold_11_F1 227	MKLSMLVLALVCISQLGGSSANEATDIMRRQLRVGKAVASFENHQSTRELEENIMQDEDNKPNEVQAEP TKFRMRRRLRSYDVELIE*
>scaffold_117_F 62	MLRSVGGAHFDVHVISLLHGFSTHTLELLVANVVSQSGIETSECEVVDDQNGNTEPNVARALIDREALEL SFALSISSVTRELRQHIGT*
>scaffold_53_F9 79	MRLSVILLAVFALSSSSVSATRNHPGETATANTAMQGVDMRTDTNQMRFLRTEADDDEERLAGKNMFN AEKIEKALQDTSYAKTLFRRWKRYEVEHGAADFALKIKNIGKDDVFGLYKSYSWLEKHHPLGAETGGGPNL FSKA金陵KAMKDPKYENTMFGRWKRGFESDAYNLLAFNLASDADVYKIYTKVTLNIHHPLAKTRKT TAKDFLFNVDRARIARAKKDSEFAETLKWWKTSGLDEKPVYKLWDMGLKTDDELYKLYKNYVKWLDIHYPPL AKAT*
>scaffold_22_F2 097	MGGLRRPALLWIPHFLVADANCNAAVVEIDFKSGRILRDIDYAVISDEFARVAISRLSTFAVVSRPKVFRNDEE GTGIVDRVRLLMLHPTDLSRSSRGDVLSFTLVRDVTNYCREAEVRLRQAREEIDGQLPCLTMHFSFQLNIRE NFSNLNLHISEGHRVVFWRWPPTAEVLLTHVEIPLDNLTQESVTPAFMEIPIHAVVGKTPAQKRTAVNSADTK TPNTEQFLFQRDWAVVPILRDSKAELDVAPPSPSIERTSTYLVLDLKTPANEGDPTRMTVEQKEFTPYFYVV EMAIFSPFWCRYDQTWWFDKTKVLDGMYQVVRGFDTKVMISTSAYAGCVRVARCSIDCENTVHLF FCHHYPIKAIICLVKISQPTLLKWMRPAADCITYLKSFKLIIPS*
>scaffold_13_R9 5	MRLLFWTLLVALVAILSSCDAASVDENKALQRKLYTKVASQALAADNRHGKRALRGESSKIAPISSVTEARAAT VSTGFGSKIMAFLRAIKEKYLKWEQKILAPSFKAKEKGTTYSEVLANYRTRLNWSGLWTSGFKRYARLYE TWLKNNPKYSHLAV*
>scaffold_73_R3 02	MSSRGHFFSTAFPIILAACKFQSVPSSPFPHVALPFGSFLMSIANTFHAPGSLQYLVASLVSAGVMSSGTATSG KSGSDASMRWLNSMMRMLRLRASLITSRTSR*
>scaffold_25_F1 407	MRLSFLLSVAMAVIYCATIONATVSDQNKVSMQSLDARLNGQADGTRLLRTHHENEQESDREERGTLDF FKTEKAAVKKMAKAIMADPSKADEVYMKWEAKKYTLTQMSNFLSKTKGKYDQVYNGYVIHLDH*
>scaffold_60_F3 66	MLIHATTICHWWLVQLANGFATFVTMEMKTLITRLLRATAKETRNTCTWIASSAGITIWMRTERTRRR CVPSPIQMAWTCAVSARRPT*
>scaffold_5_R26 54	MRPYFLLLVAVLLAISNQLRPADQSIGPLKTRRSLRLAKKSDTSNEKEADERGLSSLKTEIKKPLKVKLAWWQ HFVGPSEYVKKLGANHPLYLKRYHTREGLEMWSLAEAIRHTKSGKGKGLMAWSHCMNLDQIARQIKK PEGTEPFRVYKRYAKEFDGHRLLSDTYFIDESASVAGRYARAHIAESKVDKEYVLEFLGLLYMKPIYVKNPY YQYCSASSSVLVNAMVIKFVEVHGRVEA*
>scaffold_13_F1 424	MLGLPLAILELLHLCHVVLAHMSG LTSQNILVSNTFCKKQKLHVC SRLR LPDTNSRGGRIGPRTARAQVCAG ADQEQQEADPEPTGADPSTYQQAE TLGC GEGDLT*

>scaffold_7_F13 78	MSVLCLALMLAMATGAFESTGDAKNGVVTSSRLLGFTADQEERKFGGPATGTDVSNSHAWENFKAWF KKTFFFWRKREQTRRLRI*
>scaffold_24_R3 31	MVLFSFCLVYIIILCCYAEKIRPLTLGQDRLRCAVIAFPVHEKQRSEDTEVDTIPKVFLAKGSSVEMLWHFCAIS ADRLTFLQRGSRLRLRISRPFLFECAWGIFTFLDVPAAPARWRRRTSRRAQNRAIRPRREVAVRLDQGGL QIGFKEKIKPSPVARTRAKGNDEKECVLHDIRDKFHIQLLHVLO*
>scaffold_11_F9 7	MPTSRPCSWLHPAALCPVVFPMMPLLDCREFSNGPFNLAYVIAGSDSEPPTPIPITTARMCRSGQTPFCP KFRLGCPFFISLNCRGRPHRHRALFHRCRLHFLHYPRRHCPRRWQCPRSVRTS*
>scaffold_6_R29 10	MVSSFLPFFLFLAFVFFTSCSSPHDNEPPSRDRSTPRSSGQPDHLPLRLRGKSGSGTARRCILPGQTPSD ASKDVPLRPPPHAS*
>scaffold_25_R1 205	MRITYILAVTVAATLHSSVTAIPSVKSSKVATENGAVPAVIDSTHTGTGRMLRWVNKYEGDLDKYEGDLDNN DDLDLDDDDLEERGFSDTLKKANPLKLVKKGTKLTAEQAAKVKQALKDAADYQKMIENANKLIRSD*
>scaffold_12_F7 09	MRASKLLLMTVFLASLDATSGYNKLQGTNTEVVDTGQTQGVNSELRLRGHGPSTVDKVSADDEERASLPTFLK KSTSGATNWLKDKKLAALKLRALPMNLDKSVQKTMKDGDPRVFTLLNLQKKS
>scaffold_12_F2 959	MRLISIVLVAVATIFACCQGAPTVSDSKSISVPQHHVQTENNANRLLRVHQDEEERTGAQLVKSLSKFKKA AEKLTRSKSFNDVFKLDDVAYKENFHSATQALYQRIEKMGFPDGMLKTMRKRGDVLDLLAEYTRYWMK KYPTWTKNQ*
>scaffold_5_F36 83	MFCGVIVALVQSLFFNFLDSPSEKKVRYLIEWWWEKATRQNAAKLQTAWRSGNLRRGTDIGDQCHLFS LMRTARRLRIDKPAAIELSVEDQVAE MEATILAEVDRMEAQKVVLQRIQAKATQLATLKHKLQMKKK*
>scaffold_495_R 1	MRLSLLVAAAVALVANSDAAPQSTSSLTKFSTDVAPVRSLRGASKTKTEVDEDDSFDPPEEEERGIAMYGKLDK VDDILKQLNLADDSSVKKLAKGKGGVEQLITTDKGVKTLVITSKTKGKVTFTNIDENIRKIESTPAIK
>scaffold_536_R 3	MVLMVLSAVCMCSCLLYCAAQRLVNPSQLPTGVMEVCRHLRWNHWISRRDTALQASDTLQDATYAKGA SACEVTKLCELVSSCRRRGDAGSRLCPA*
>scaffold_5_F17 32	MALGVAMAITLGLLSQLSNTTAVGTENAVGPVTRHLSL LDQKVSEHGAALDQQPLAASMEATSRRSVPQ NPGGPLGGGRTMEAIAPRYEKAAPETIAEA VAKAAEEAEEIAEALIVALGAENDLQIAEITEASILSLATNSIT EGLGGEIIVVGIEIEDSDDSSDSQEEHTTRKTKAKKHKDW*
>scaffold_1_F44 48	MNCSQQSRLFLSLAFCQHSRLIFALVFSQQSRLQCVYDDRILPDEHLRAIRR DVSPSLYSRTLRRPSNAP*
>scaffold_53_F9 2	MRLGYFLLATIVGFLACDNATASVSESTSSKLTAREEHPIHGRIGDFTAGHDNKRALRSEDEDGDADDSDDEE RDILISTIHRPKYWRWFKA GMTPYAVQQVLGLTVRRWKPFKRREYKGYVVFYTEQCHKPEYHDFCKKHA DP*
>scaffold_12_R4 2	MRLTSIVLVAAVSIFVCCQGALADSDKSISVPQHHVHTENNANRLLRVHQDEEERAGPQLVKSLSKFKKA AEKLTRSKSFNDFKLDDVAYKENFHSATQALYQRIEKMGFPDGMLKTMRKRGDVLDLLAEYTRYWMR KYPTWTKNQ*
>scaffold_3_R28 40	MMALLVGA PWLLPLMLP RFDLAPPVTL PRERALRFSSLRILPSLISSSLANLFTLLR LPMTAPPRGFLPLR *
>scaffold_68_R5 30	MKTIIASLLTAVAVNAANGDVNALSAISTISDAKQVSIRALRAAHGSHESSGMGSMEDSHDTSSTHESTV AGDDDDDHSDHTSSKSSMAGTAGSAGPSNSTESTQAPDTSSAASITVAAGSIFAAAAAAFL*
>scaffold_24_F1 366	MRLLLVV ALAFLAADVSALNDAPS KRLRSTVRVDEEEERGMWETLSSKVTKLKPNQVAIKAMDDPKI AEVAGTSLTSLSVNPKKFDSDGFLFSSKAFNNLENVVR LNQDINKQTSVAKVSTGLGDQAFHLFFTA TQSSDAVEKSGRFFRDQLTQWATEGKTWTEVSKSVPKGLPATYYPRLENKYFDILFNLAHDTQKRAARLA RLEKARMAANTAA*
>scaffold_77_R3 94	MRLTYILAVIAATLHASGTAISTDKSVKIPAI AADDAGRMLRIVKEKPVSDKEAEREERFANPIKKLG AFLKKK WDKQLTKEAIKRDENRRKWI REQGFEP RD*
>scaffold_5_R20 89	MRLSFVLP AVIAVTFASSGN AVATADGRNTGLSAITPPNVVASIDTAVGGEKRSRLRYHNNKDLEDDSDDEG LEDAEEEERRGNNMFSATKLD EMLDGKQVMSRFKKWKA FGYNTYNLPDAIQARKYDELRKMYRKFLYNN*
>scaffold_12_R9 67	MRLQFAVL FALS VITAANGFSETTAQQFN GELSTGEKQHDEKRILRTEKVDEDDEEGTEGEERVQVSPVSWII DLFTP KTAEQIAEAKKAEAVKF YTKLANS P S FRAERFP NWKIDGMQVESVLVHLKLWGLDGEKF KAIATKY EFLASGKLS*
>scaffold_535_F 2	MRLTCILLVAAASLGVVL DASAATTGNTV VANAAMVISPLAPESQGRRLSLRVYDDEDDSADEKDDDEEE SA DKVDEERGWLSDKMALTSVSKFGKSTDEMGEVIKLTPAQIDTIFDKGEDSIQKMLPGFYSGMDFQKFD DLIRALPQEQQAVMLSAYTKYLHQNGRFS*
>scaffold_29_F1	MRVFSIFLLVAAATL VASASTESES KQR D N SPAAP QWRTITENEVPTKRN L RKKKIEER ATTA ISLF DAVK

837	AKGWQVLPYAEELANLDATVRTQYLKLLVNNLERKQIVELTGQVPRYVLKHGDSKATRLVQYNKWIFNHFKE AVDPAWVLKNYPAFFKGYDKFYQNRFTRGYKYA*
>scaffold_11_F1 676	MRLPHFLLVATATFLSSYNSATVSTEGQMMPADAAPVRALEANDGKRFLRYKKEDMYDENDGDDDV DDDDKYDEEVEERGVMTATQVAKWRLKETLVTIKEKLTMKGVISAKNREKYNLFTAVYGRA NPHVFERL*
>scaffold_31_F1 184	MRFGLFLALLVATFVACSTVANAESVALTGDNEVRRRLRNQQNIAKAAGDFISKSKESATLTКАINIАKTANG DEAAARRAVMLAAGAKEGAKEAKLSDETMVKLSAMIAESAKKNPKSWPLKKFVKITLGIQVGGLAIYGAYKLLF DKGSSTAATTTTTTSSGAA*
>scaffold_40_R1 072	MAFSSLKFLAMATLLMVQVNAEPSQQTRNLRIQSDDLPQEQRRLGPWNKALVINVKNNPGVSDGVMK LLKKADLITGTSKKIKEVKDAAEKVKNSAGVKDKVTGN*
>scaffold_62_R1 2	MRVTQTLFAIAVVLFATTNDLTTASKATTSSKVQKDGLYAEIYNVEDGVVTKMRVPLGDVEQYSDTDDEQL KSTLTTEERAHVLPFGVDSFFKGLEKFVGFFKFKGRRRLRIEEA*
>scaffold_4_R32 84	MVKCCAFVVGVLMALASGENNVGDVGNGVKTAEAVRVNATVESLRFPVLFDAIINRSVLEHSRETRD LRVLSIERGVVEVEVEVDAATVSSCATTAKCPPEVNDTEIDENSVPPPSLQQTDRLVRVDGENVSTLSFQD VIDSAAGSVAVLPLFKPIGSAHVLSGFARVEILSPKLLEFEPAREQVVMWIEEKARLKAEREAREREIANN KELQERLEKERLLAEALAKKEEELLAKLDREEYERTRMTPHNLAAGKRRDGWEFRYEVEFKTKGPIGLNWDL NTRDKAVVSHLEPELPAQQLNVIAPRDQIISLNGVDTSKMGPQEVSVEVYLSAIPRKMVFVQMSAERAACK SAEKNPVKRVVMNWTALFADAEVLRGWEVRHLHASWSVPPQINETNASLPLQFELPTITGCSSFPVQSS NETAGVYVLAYRGACTLVEKAKNARTANGSALLIVNNANGEGRFTPSGTVVVERDVPVTLYESTFVFIRCIW NLICGVCTAG*
>scaffold_211_R 6	MRLSQVLVIAVASFVFASDTVATSQNQAKISKTVQSSSQRLRSNHYPVKEEDESEDSDVFEERGFTTPDEE DLEERSPLSAAIVEKLDIASRWGTSWAAVAMGQSSISEDKIKALLRDAYLSGNNAKAAAKLAILRANWT RSQQKW*
>scaffold_80_F4 73	MSFLSLFCFTQLLWSCNFQTVKAVFEAYAQTYVRIPPEGTVVFSTALPTCTWKSTDVSFSFITTRILRF*
>scaffold_62_R7 23	MIGCKSLISRWPFFLTLAADIWSDFVACNPQLTDQARHLSRERKRYLREYPGAARHFHRRFKAFFFL QHTLWKRSSWGNCRLLLAR*
>scaffold_1_F54 50	MKTATTFATVLALIVATNAAQVSPHTPALRGLRLTADTPSVEDDKEDRKDHHVKKVKKIAIPVVPVVEVPQF IPVPSVPSTVVASSNNAVVGPSTNVAGPGAAAPGAPGAPGAVTPAPTTLNGRPAATPAATSTTRSRPTA PTNFAGARPSGAQLPAAPVQVGAPGMTGFPSAGVGDNNTNNNAFGAGIGAAGGFPMTGNGLAGGFGR NPMNGFGGANGMGSGGIFGGAGAGPAMGGNGFGGNGLGGFGQGAAMNFGQAGNNGFEGQGGF GSGFNRRERHRR*
>scaffold_26_R1 820	MAAIKISWQPGCFPVALSYLVWSSSELCAANRRMVWIRRSLRGKTRLRSPYLTEHKLCMFHVFMHFFYL N*
>scaffold_52_R8	MSALRAQIYLFVVARAAVDGQVVCVAVQQDLVRARAIRLATTNHVKRRLISTRRPPQSRHQARRRESAH TLQQVQGLFRALRVRHGQVKLERHHNHERRAANRRFALLRVVFYGILVGQNDGRILLDR*
>scaffold_33_F2 96	MGWVFCIAILLIMASMSIPDPYYGDPAMHSIDEEDWTDDQIDSINADAPNSAGKYVVPMMLASFGYLMCE LQQMPWWSSTHNENRSNNVAAFKPPFTPSKRRSLRSELLSLRSS*
>scaffold_16_F1 464	MRFSQVLVIAAVSLLFANDTVAVATSNHAAISKTVQSSSQRLRSNKYLVDCEEDQSEDSVDFEERGGGKS HPLTNRKLRSLKQSLEVGEILLIRELLWVSQASRPRRLMRCLPCERTCFATGLSRQQR*
>scaffold_2_R38 99	MRLQSIVLILVATLVATTQATTKLNAPSSETVDISPQRFLRKHTQNDEERGAASLIEKAKRVFPSKITDKTLQR WANKKKSPKQALTRLKLDIPGRTSLRSSASGPLSWPSATRTPKKRWSRLYWRNTAMRP*
>scaffold_28_F9 19	MDTCLSVLLSLYVSACSSQTALPEVNLASVEMASVTSLYVGVTPLSIRWLRAIAEFYDYPISSQLDDHFCGL WHRFISASHRPVPQEENSTGWMKNSQLAPLPGNHSLLQQFPLCILLGEEPQMTQETHFAKMLDKIVPIG VLHSFISGDMSENGDRFKVDEILSCFPVDDMNAERKIDFGAQLTTLFAFSFMLVYPPDPLKQLAQQM VDIAERAHGNLTQLSHLGHIAASKMARFTVWNIIQGCDEDLVTQSDLWRDALKMMKEKYSAFDWKLLNIA DLLGEFYVLLLRTEHIDPRAVCACECFKIFVEETRDAVAESSVLQQLLSCFQSIKRLAVRNRRRIGRVLSSLLVRE NCEYFMKSTDKGKYLGFVQQEISDILSKCNASSQTGRNSVGALSVSADDQAELEWVIFAVCNELGSGLKH ALDIDMVSDGISSLDKLNALISSLRVSSETSQSEEERKKNLQSAMGTRNQAIQQILMFIQREQSRGAQFY NPHPFSGTASAISRETYVNAATQSPAGSLADVRLSAVTTCIDMVRDSSQSTMKLYGKLAQTLLYLSSSGTFGT SRYNCENDAASLANTLGQLGALHASEYEVSPAEQEGELSRLYWRHFHRGALREIKTTGLVMHENVLTYLSL LFEGTRFGHVDPVTVEETLKLQTVLNLEEGLAALARSKDNELKAFLKFESSPSNWSSYSSSGDWGVKP

	KRTFRQFLKHWTSAEELGFEAWVRSITA CLARESSDPVLKAC SALSAMRV DMAVFLFPYALERILRLDN RDV DNPDEEKSSGVSEPSRIMKAANQGIRFVLTG STGDALSHHVLASLDGTQRESEFSQPPEAVQLVVHSINFLR ETEKAQFVETNGRPQT VSTIKGKGSRSS TYTAASIGRQH LNDLAYGCLVDVDFLAVA KAA VRVKMPYSA M QYVEMWLEKKQGGKITSLSSLDRGMVDTVRDILVEA YSFDS DDDGIYGVNDG RTVKS QLVKYNREG LH ARALPLYDVSLQFSSQQLVSTD ETLNTPP RLVEGILTSQSLGYNHLLTGYLQSLQSGDVAGNKG ST AIAQ AL EHKYKLAWKSMQWEAVLSGLSASGEHSSH HQMIFQGLRAIAHGNFTRLQGITTAKEQVLR SVQLS LHS F STKDSY SALVRLQAIHEIEELAN HIRNSV PPT EPLV FSATT TGS GPFLPS LGGT LAAPR QETLTVPLLEQWHQR RDQIKNDFDKAESLLAEE LVQVAKPSDNARVTKLYLDLASR KAGRIAIA YRALQK LEHL DERG SLGIYER MQCQIQKAKLLW KQQE ARSAIWTGKVS SELTG YL RDT SISATEV TSQ LLLL V KVLTFTGK WIAF QR SESS QVI LEDFFQKATEIMS NMDPEAVSERSRDAAKAHFALAEFMAGM YQQV SRTV TSQEW LTGK MVVQARH DEL QELQSMEQNMQNENRAHIFALNKEV IYDMNERSK VEASV DQFLIGAICSYGKG LTL SQAEL DMV F R VLS L WFNNQHKPDINRVVIEEVIDMVP SYK VVPLS YQI SRISSAS GTF QT ALRK LVMK LSEQHPHHTL IQ LIA KNS GDVEGKGALQFRTNVGDAKAEGAKVYLT ELMKTEQRELLQSLDSIANAYVQLA LFDTSEYHGKKKPI LSTV K IFETNSGRSGGTTFDQCLRARARRGDSVLP A VLT S QIAPQPD MNYSNVVRMYSFEPQFSITDSG IHRP KIIYC YGSDGEPYKQLVKQGDDTRQDLVIEQVFETMNQFLMEEKATRK RKLRLRTYRVVPLS PIAGVLEWVENTMP WGSYLVSR TS KRLSAHERYHPHEWKHTECRQYLNAPDKLP AFLEIEANFTP VFFF LEKFPDAAWVYQRR LSYVQSAAVT SIVG YILGIGDRHSQNI LHEKT GELV HIDFGVVF DQGMALYTPETV PFLR DMV DGM G ISG VDGVFSRCCEVTLQLLRKKSASV VTILEV FVHDPL YRWTLSPLKALR I QEGQGHGKP TRS RSSR SSSG SA EYDG TGSMQDTQPADEMHAEPGST DAAARALIRVKQKLEG YEDPNGSALSIEGQVQ KQLINA AQDPLNLCKLF PG WAPWL*
>scaffold_29_R3 08	MTSSCRLPLWSGVSSVSDAEIPLRRPFLVFLH RGF WYSRCQPVVSSL CFI RTRFLSE VPWPPARVTR HR VHRSSGRLRFTL LSRRVCCSYRARRSLSACRLRAF VRSY RWY S IGR LFST QLVFTV LRMDHPLGDVLL RL LHC RLLCGALDCGLRLSFSILS DRPI A PISHATI RYIP DAGEFIP NSLW VREN CPVG VSR CRR CRI H RI PYANT WYYHGVSGSRPNRYRIRHL RSYPLL PDVLHPLS YI KGVVHL RRLVY RLVKCLRLA SH*
>scaffold_21_F2 030	MQR RMSTPFFL MITTIG DAQ GEVEWMIPADNSPLIKSDSFSRN CGAT RTRW ATGV ALPTS MSQ TAP RA AGGRAFPFP IRRCLRFQM ICLNR SICTGV LPSASSI WI SSIL TSV VALT GSMA I*
>scaffold_494_F 2	MRL LYLAGV AILAF AADAKVLV SDDS VNNRA QAA SASV VSTTR LLRTS VIDE ERAGG ISAS AS DKL A KLF SS KVT DEQLQKWL NNNGKTA ESF VYRMN LENTLYTRV FESPQF PRWL QY ADDLS ASGKG ASA ISV LST KYG DDT LYK MIGWAKKE SSTK ALGIRL QTEQ LEH WVKIGKD PDEV FKLY KLDY AGN RIL GN P QFSAW TKYV DDLNA K NKGAFVSI IPTL RKYIS DDDLF KIAL AAKR SGET KAM GTK LED AIV QFW NRAGGP VGN GPT*
>scaffold_100_R 322	MRICFVLLTVA ALV TAVSGSSV NLRS QRIDVV QDKTSSREL RGDL NTDEA TEERAS FDFVKKL VAKL GDPL ESFAKRQTKYI F SDDIFDE M LKKF PDPD AYTT LKLHV KNR SNK YGV T T ARHKLH QN FLTS YIDKFPN WR SKL N*