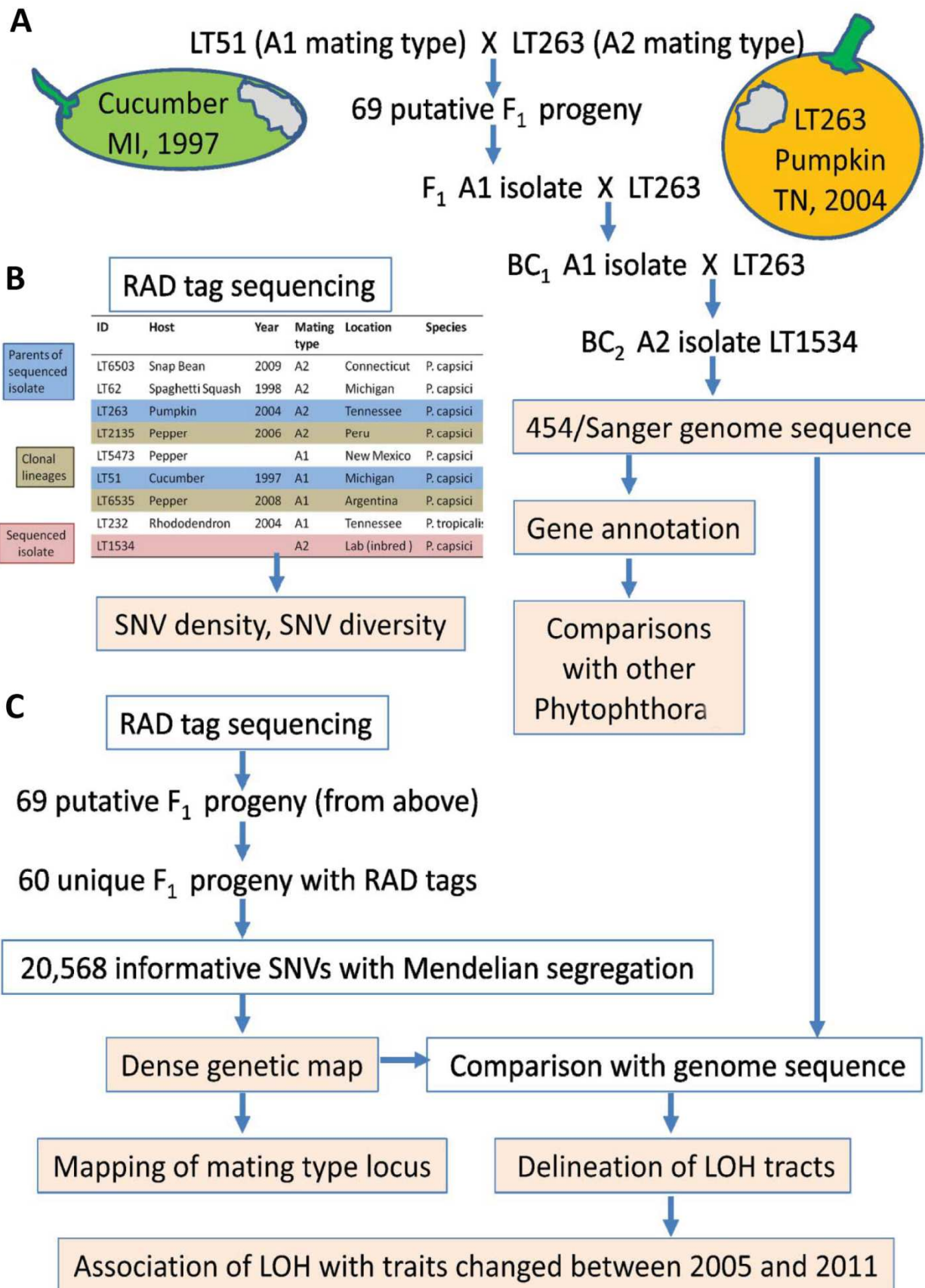


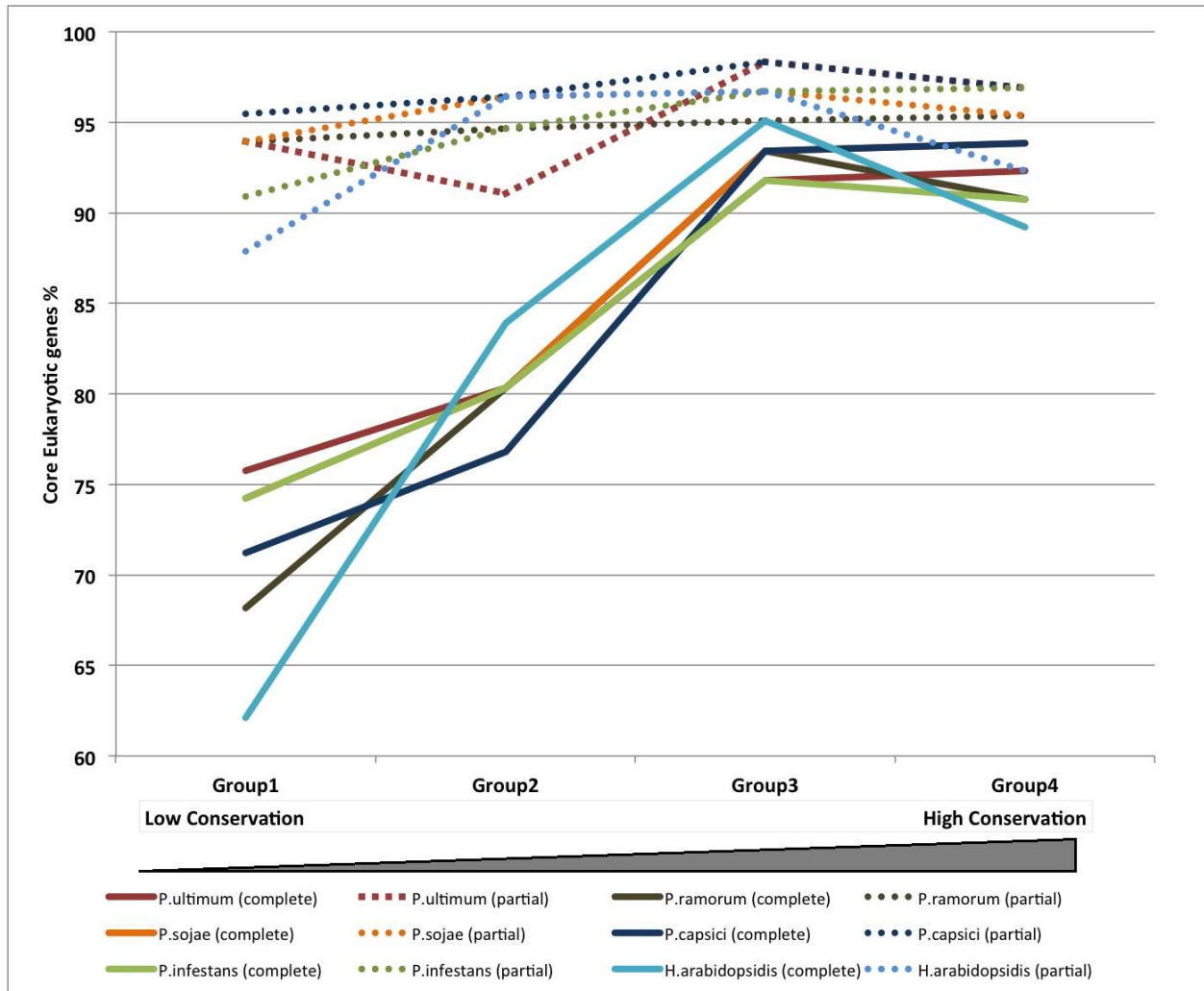
e-Extra supplementary figures and tables

Supplementary Figure 1: Experimental design for molecular and genetic analysis of the

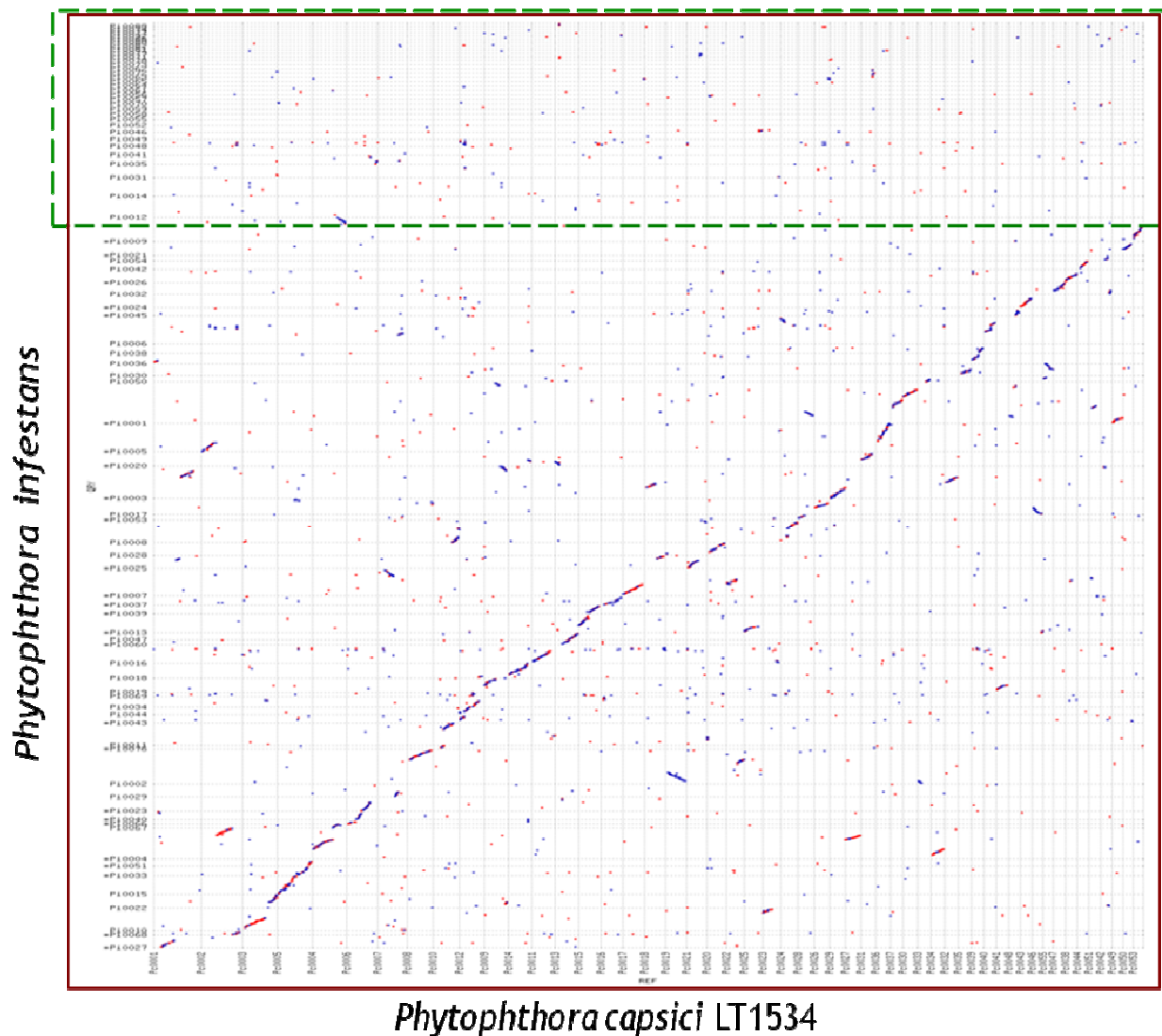
vegetable pathogen *P. capsici*. (A) Two parental isolates of opposite mating types were crossed. Of 69 putative F₁ progeny, one was backcrossed (BC₁) to parent LT263. A second backcross (BC₂) yielded a moderately inbred isolate (LT1534) for genome sequencing. A hybrid Sanger/454 sequence assembly was 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-Associated DNA 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.



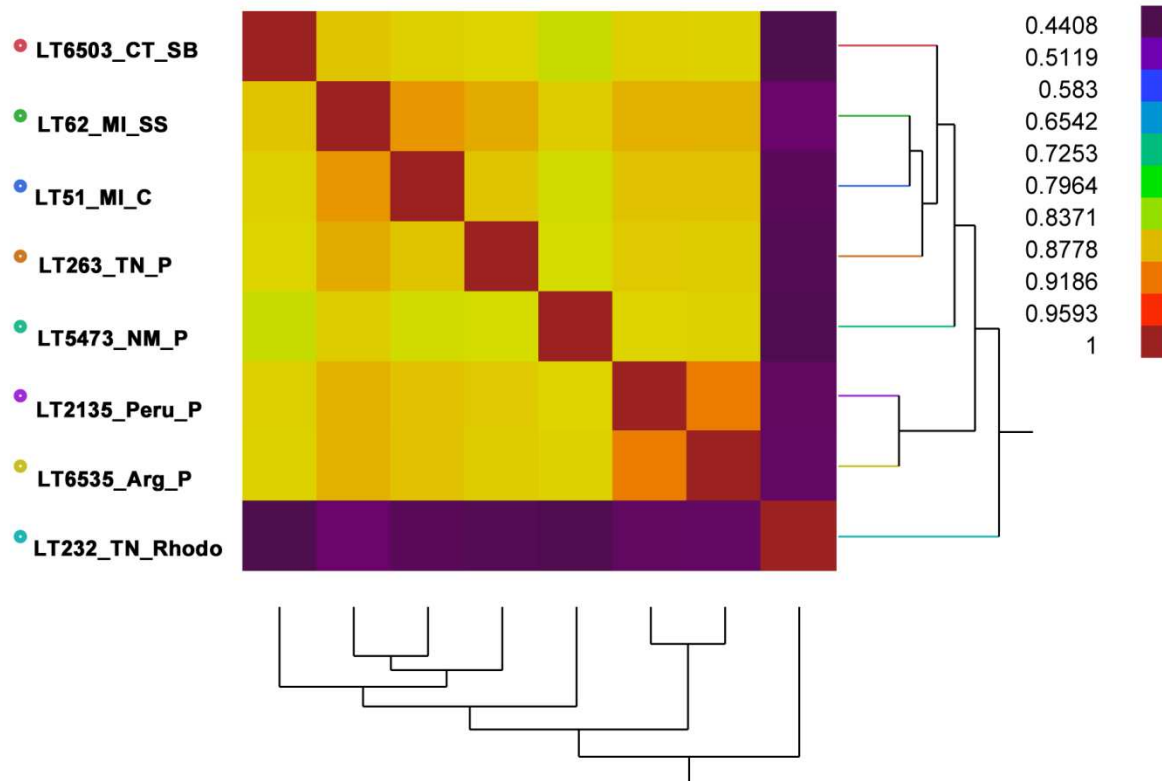
Supplementary Figure 2. Graphical display of CEGMA analyses using standard parameters. *P. capsici* has the highest number of identified core orthologous genes among all oomycete genomes of the *Peronosporaceae* and a comparable amount of fully covered genes. The increase in fully covered genes towards 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.



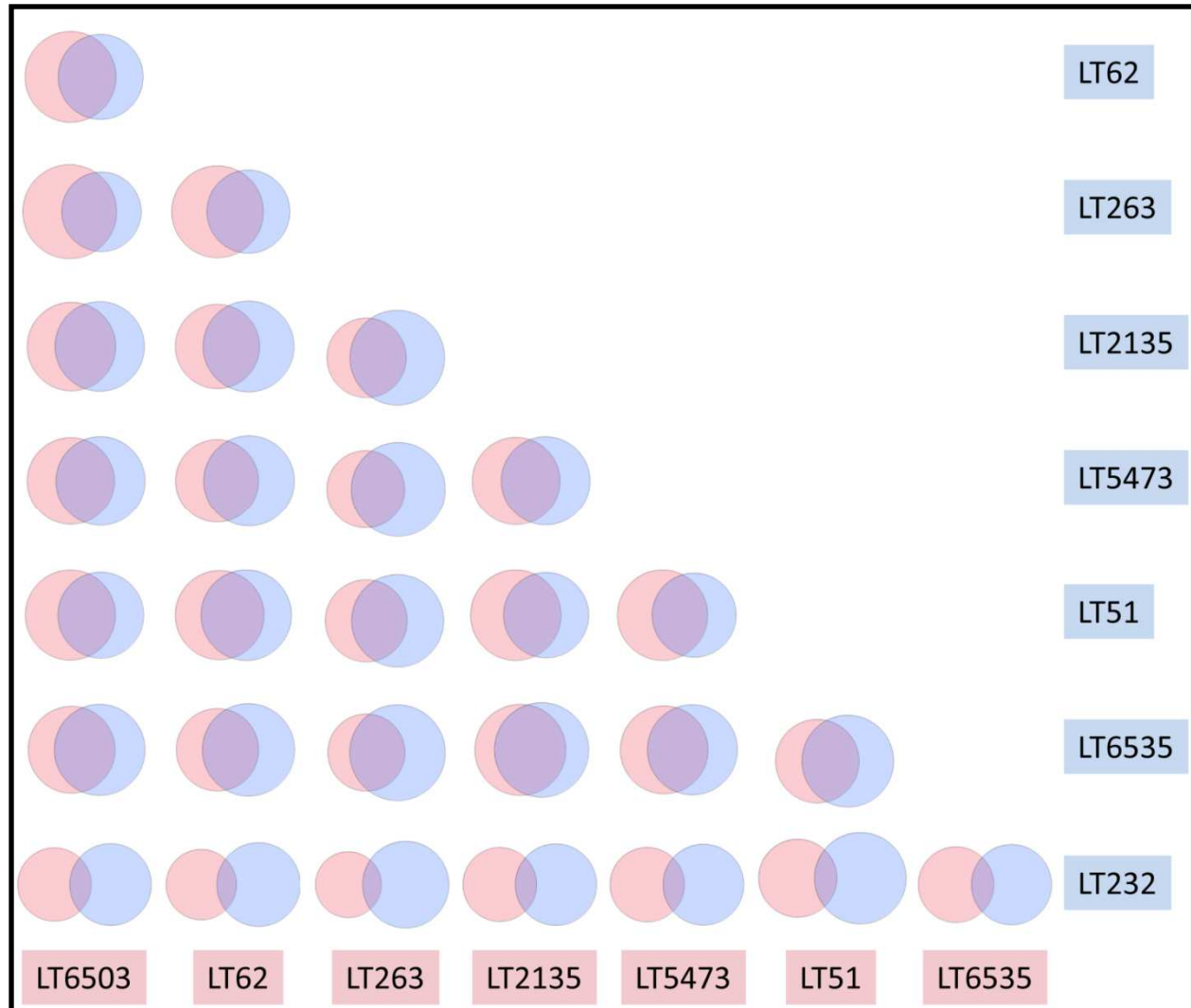
Supplementary Figure 3: Pairwise comparisons of gene models from *P. capsici* with *P. infestans* showed good synteny with some duplications and translocations. Sequences other than gene models were replaced with Ns. Nucleotide sequences were compared using PROmer package of MUMmer, which identifies 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 syntenic genes in *P. capsici*.



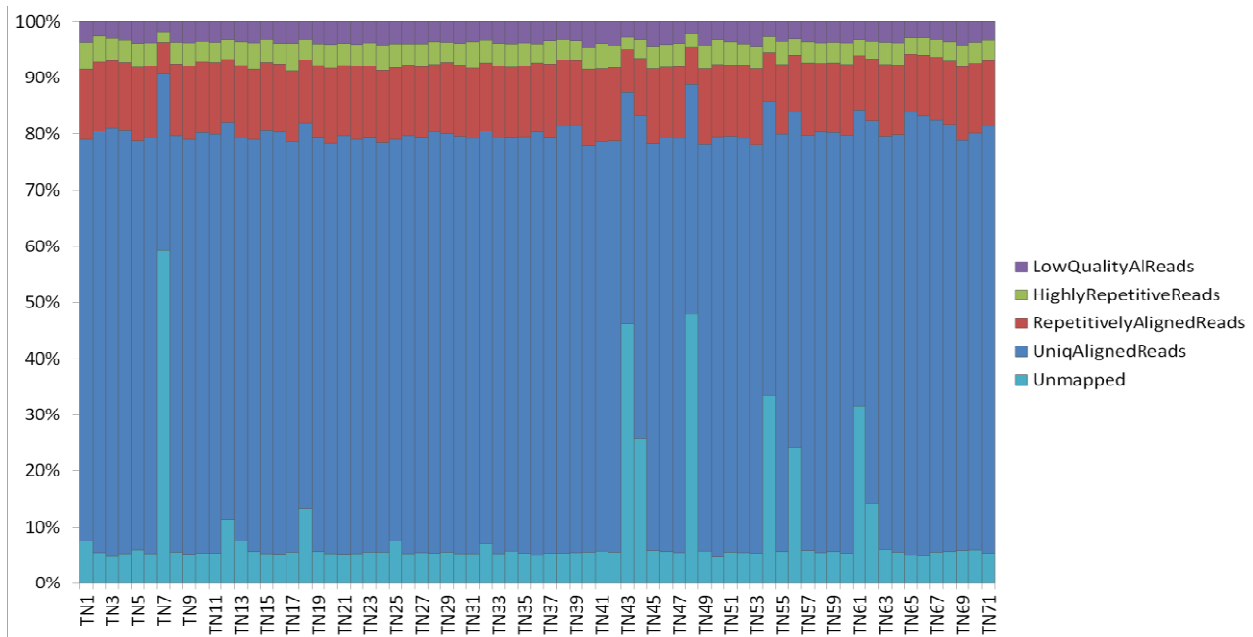
Supplementary Figure 4: Dendrogram and cross correlations for SNVs from 7 *P. capsici* and a *P. tropicalis* (LT232) isolate at regions with 30X coverage in all samples. Left side of the panel indicates isolate origin as abbreviated US state, Peru, or Argentina (Arg) followed by host designated as SB = 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.



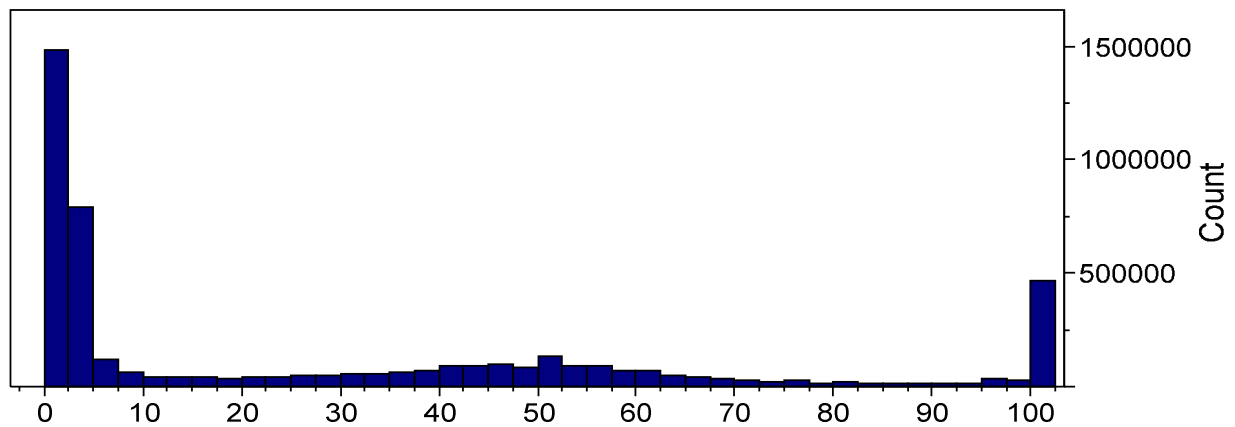
Supplementary Figure 5: Overlap of SNV content of 7 *P. capsici* isolates and one *P. tropicalis* isolate (LT232) sequenced by RAD. SNVs were called at sites with 30X coverage in all samples if supported by ≥ 4 uniquely aligning reads of Q score ≥ 20 . Of note, LT263 (recurrent parent of the LT1534) 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).

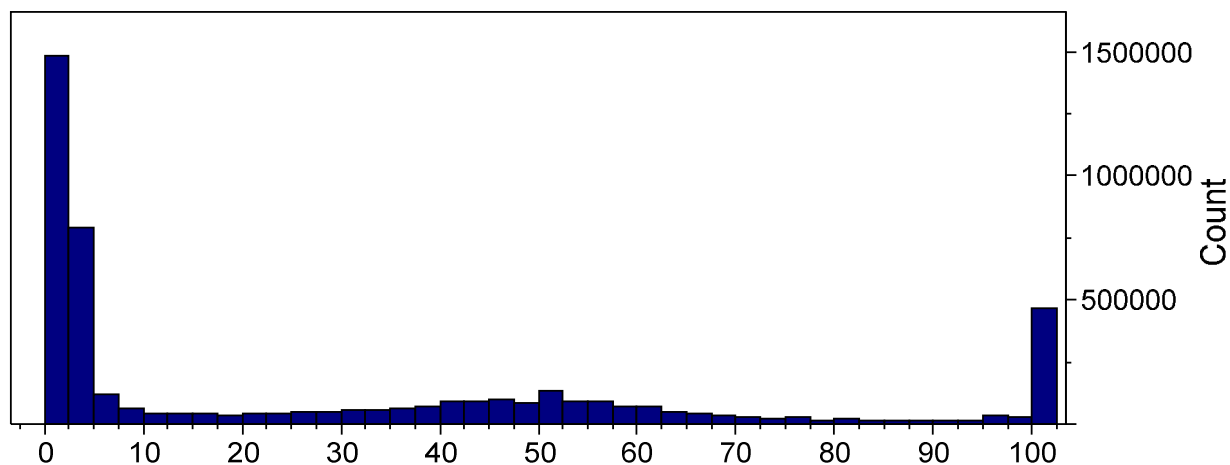


Supplementary Figure 6: Relative abundance of RAD sequences from 69 progeny and two parents based on alignment to the reference *P. capsici* sequence. The X-axis shows isolate ID's and Y-axis shows relative abundance.

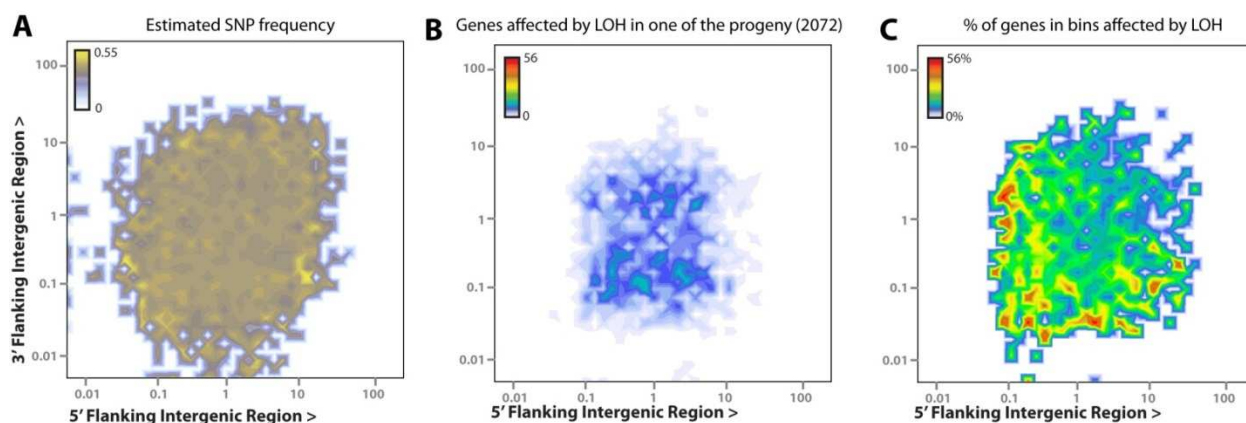


Supplementary Figure 7. Frequencies of alleles based on the requirement for 10X and 20X unique coverage and an average quality of >20 for the alternate alleles. For 10X unique coverage (top) this includes 5,849,663 SNVs and for 20X (bottom) this includes 4,449,193 SNVs across 71 samples.

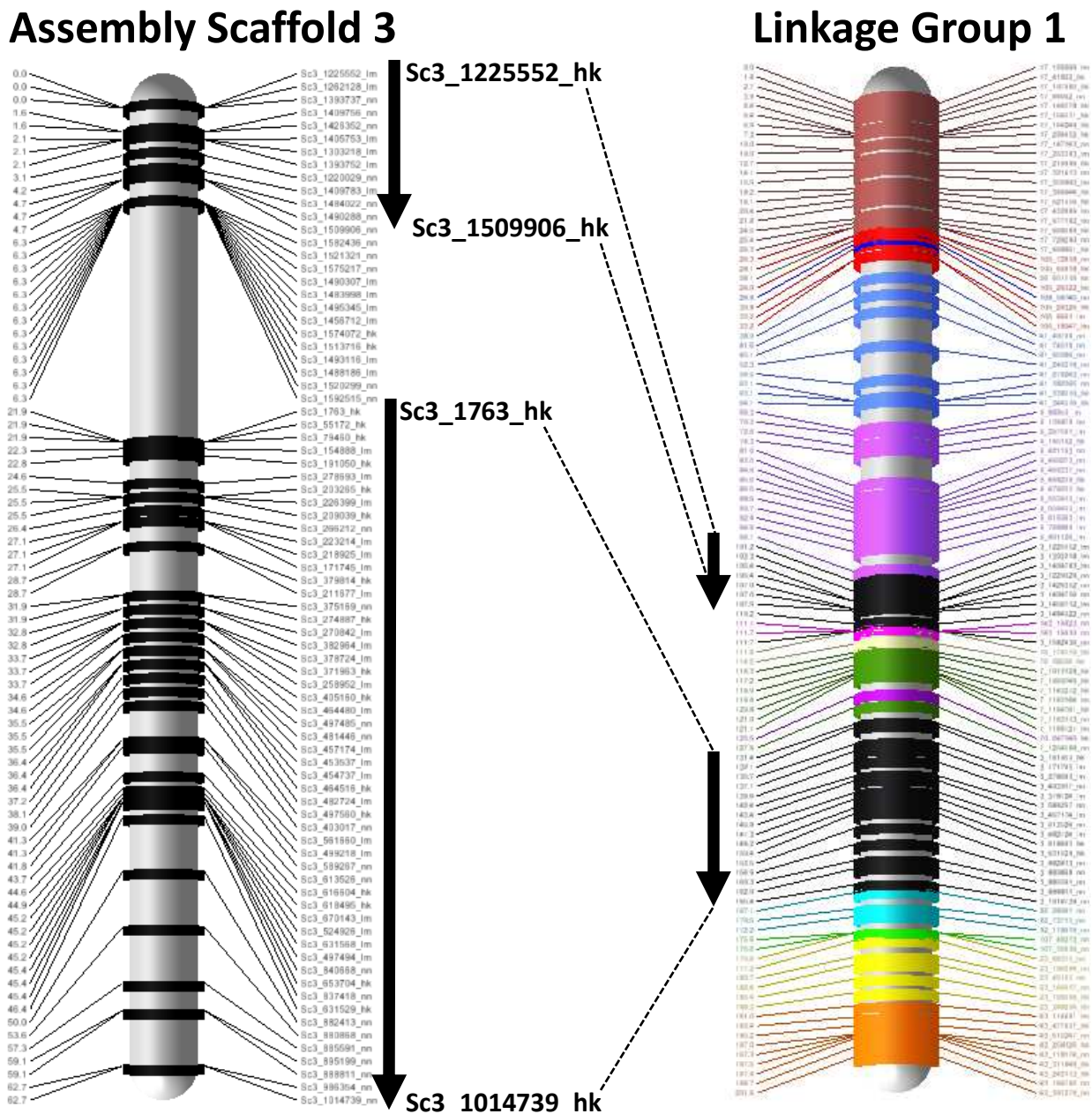




Supplementary Figure 8: Gene-poor environments in *P. capsici* genome show reduced LOH frequency. *P. capsici* genome architecture illustrated by the distribution of all predicted genes according to the length of their 5' (X-axis) and 3' (Y-axis) intergenic regions, counted by two-dimensional data binning. (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.



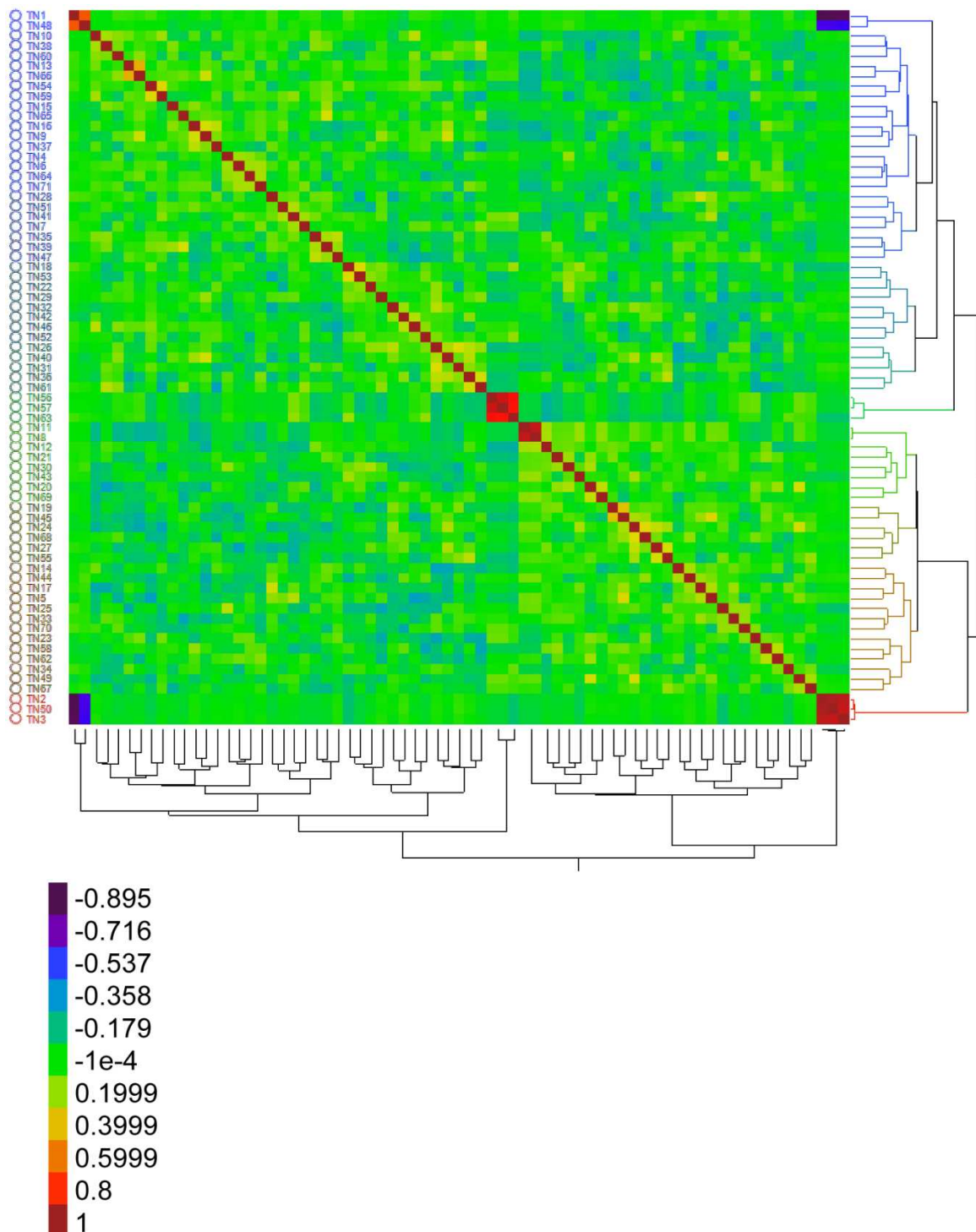
Supplementary Figure 9: Comparison of the genetic map of *P. capsici* Linkage Group 1 (202cM) with the physical map of Scaffold 3. Two blocks of Scaffold 3 comprising 1.6Mbp contained correctly ordered markers on Linkage Group 1. However, an assembly error occurred between these blocks.



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_fgenes1_pg.C_PHYCAscaffold_20222 |
| 2_1452313 | 3 | 2 | 0.00 | 0.00 | 0.49 | | 3 | 3 | 3 | 1 | C | Silent | | fgenes2_kg.PHYCAscaffold_2_#_197_#_4100895:1 |
| 2_1452392 | 3 | 2 | 0.00 | 0.00 | 0.59 | | 3 | 3 | 3 | 1 | C | Silent | | fgenes2_kg.PHYCAscaffold_2_#_197_#_4100895:1 |
| 2_1452394 | 3 | 2 | 0.00 | 0.00 | 0.59 | | 3 | 3 | 3 | 1 | C | Silent | | fgenes2_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_fgenes1_pm.C_PHYCAscaffold_20199 |
| 2_1486428 | 3 | 2 | 0.00 | 0.00 | 0.58 | | 3 | 3 | 3 | 1 | C | Silent | | estExt2_fgenes1_pm.C_PHYCAscaffold_20199 |
| 2_1486435 | 2 | 2 | 0.00 | 0.62 | 0.00 | | 1 | 1 | 3 | 1 | C | M | K | estExt2_fgenes1_pm.C_PHYCAscaffold_20199 |
| 2_1486458 | 3 | 2 | 0.00 | 0.00 | 0.58 | | 3 | 3 | 3 | 1 | C | A | T | estExt2_fgenes1_pm.C_PHYCAscaffold_20199 |
| 2_1487541 | 2 | 1 | 0.33 | 0.00 | 0.00 | | 3 | 3 | 1 | 1 | I | | | estExt2_fgenes1_pm.C_PHYCAscaffold_20199 |
| 2_1487551 | 2 | 1 | 0.33 | 0.00 | 0.00 | | 3 | 3 | 1 | 1 | I | | | estExt2_fgenes1_pm.C_PHYCAscaffold_20199 |
| 2_1487569 | 2 | 1 | 0.33 | 0.00 | 0.00 | | 3 | 3 | 1 | 1 | I | | | estExt2_fgenes1_pm.C_PHYCAscaffold_20199 |
| 2_1487581 | 2 | 1 | 0.33 | 0.00 | 0.00 | | 3 | 3 | 1 | 1 | I | | | estExt2_fgenes1_pm.C_PHYCAscaffold_20199 |
| 2_1487584 | 2 | 1 | 0.33 | 0.00 | 0.00 | | 3 | 3 | 1 | 1 | I | | | estExt2_fgenes1_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> |
| Sequenced isolate | LT232 | Rhododendron | 2004 | A1 | Tennessee | <i>P. tropicalis</i> |
| | 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 Silent | Coding | | | | Non-Gene | Genes Per Linkage Group | Total Genes in Linkage Groups |
|---------------|---------------|-------------|-------------|-------------|------------|-------------|-------------------------|-------------------------------|
| | | 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 | 368,003 | |
| 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 | 130,234 | |
| 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 WLDGAGAAAVTVDEADRVPMLMLDKHGNHHKFKMNP LLWIKNDQHFGENFRPVHV VVVVPDV AHAQTGLWLVTGFVKNALNTKGIRCKLYWMATLRIGY YDPARRTDKKNV AFWYEDTKLCFHVLFETKDAALLFETDLRIEPTLGSPLTNQVVETRVAPVNAVSTELQ RVFYGDYVPDDSKSPQNSVSSISLTTSGSNLDSSTDEFRRFORIEHEKFFLPYGKAESCHLV SRKQSRNHKREFAKYDRDSNSRLALS RDMHGWF DGM SIEVPIVNMLPGSVKENQSIGN RRKVVVEVFKVLDAGCTDRVFSRLKEGSTTTNDPLMMKTFVHVVEDPETFCLCMRWKH DDNAERWRSFWDMPAVD* |
| >jgil106435le_g w1.12.217.1 | MLKLCFCVVVG VAGDAFPVNIENETVGDMMKKIKHEEMYQFPASELQLFLAKVPKEKH DMAWLSSRSEDVKKLKKGEKTP LIDILTEEDQELQAEDPLDDVLRGMDPPSLCQIHVLV MAPPQDSLRSHTLTLSSVLLCHVLT KAPTPTDRNVDFKDDVCNFYGCYSPDESCVRC MLLNDAFPSELV VASHLFRCSNEDVSDVMMQITLS DIDDERNGLLLFKPLKYAFDHFQI SFIRDDTDVFR LKVFDP SILATPIVDLKD RKGKVLSTEQTQLLSRIENPCRNTQTTF GDVDDSA LTFGLERPFYRCLNLQARVARVMAL EKKWIDASYDFQDFWSEVSLDDKM EMFHRSILNS* |
| >jgil106457le_g w1.12.537.1 | MKLFCAMVGM AVGII EVIDIDNNAYVTALRDAIATKNEEIKKSALRLKLFLAKKGNLW TDTEAAGVGGDLESLGFKLMKSVRL LKNPEYFGEDFQTGEGHVHVLVVVPEENMTVG EPVVDVDGVNIYVTSNMTLNPDDL VAFWRAFQAIDTKIEADSVIALPEGTFILGNPKVGS RIYIRPCYPQLWEVCWHIIHETPNL VILGNPGIGKTYFGYLLLLFLARLGKTVVYESRR TKRRFLFSRNVVIKGSQQDFDDILEQDTTYVVDAMEPREFQARTILVTSPDRDVWYTF NKISCQTRYMPVWTEQEIFSCREQVYSTIPKSVVQKCFYRWGGIPRYVLQYAQFDNHQ ALIEKALEVVD FDLWLNAYGKLD DNN SQAHRLLHYRVNERFTCDYFGFASSFVQHEV YQHLHKKEKRKLL EFIGRSVSGDLSVLRDRLAE EHDHHCARTHKKLRK* |
| >jgil106668le_g w1.12.123.1 | MVKLFCAIVGVEGVSFVSVIGEGQTV EELKKA IKERNDDKINVS WLGLQLFLAKKAKG DGDWLTEKDVQEGVYDMSDLQMLRAARAKLRLVKLS DNDVNEEQKVEGGKSVNVL VVLPLGTTTTIKVNERENNAL TDELA YYQRIGQEIQSN CQQHC GPILDKIDS IYEKKPYPM PFICVQSSGMGKSQ LAFALGGEGREHPRPWFY WTHGTVSDYDQRIYRNFASIGSAFDS VVKQDEV RKEEEDDILNCTSVLYMTK KLVWYGFIIELLRYCSRSNVGAQMVRVENQTF YVTKSNLEDVIEVRSRMEKNGEVLPFFILDEMPPSRTK KLSAFQLNVFRACGLV VIGMG TDANISNLVGKPEHSRTDPH WMTVVS CFPPWQSIPFGDPAKEEVWQKVIELHPVVKH IAEHSRGLFSRCFVDAVVKFAMEEVSENETFALADLLDAAFKAVYAELRITKGMFTEE GRDAQLMALS YTIGSNKPPGTRPV TDS DSTSEPTPKRRKLDVDVGVASMR AHFANS G YEGIADVDVLQGD LRFRTSDAWSPICQFP AIEKDVLLYLAVLGGK KFSYNDTSS TLD VFEEFLERNGNREESNALRRDFTNFIFEN VVAHALFCASRRNGARGIALYEF LSGLVSE FQDEYYQRDEFDASALLKEFDGLKEKFAEKKIPFLAPPNARWPDYILEAGGDCNFGHFE RMDGYVMVPGVDG PLFACDCQYWKDDLDSDAMKKIIAGLNGGATGCGDDQ QKPKW SNWSLALVFCRKL EEFESEQREDWEPSTGIATVDCKAWEVNWISKPEAGEKLVIVVQT GRVLCPL* |
| >jgil106736le_g w1.12.192.1 | MMKLFCAIVGVAGSAFSVEVNEDQTV EDLKTAIKNQNRMSVDANDLQLFLAKKKNGN RVEWLTQLDVVKGVMDANGFTHLQFADAKLRAVGLK SSELVEVNEEDVAVGKGHVH VLVKIDQTAPNVELMPFTSTCWLVTGSGVNALNTRGV RGQLYRLAHTELGYD PANV RTDTGNIPRAFWHENNDIQIHVLFKKEEHALYFQSHL VDDFSIPIKSFVSKAPCPTDLQRI FRDHVVADETVSPQLSMFSSEISVFDPSNPVFKYQRIEAERLFGSHGKAESAHLISASHCR NVTSYDEYDKDDNNRLALSREM HGAYDGINCDFPLVNIEVVSASDHPELDHRFKVELQ VSVYSHEYVFLGLR LKDGSTKTEDPLVMKTFVYVQDKNIFCTCIQWKYKNRQLREEF FTTNPRAT* |
| >jgil108643le_g w1.15.666.1 | MAKLSLSCVIVGLMGS AFSVDIDADQFVGD LKKIIQKKKNDLHNVDADKLQLFLAKKD KGWLPDKSEA ALELKKGEVHEDIQVLINGEEMEATKTLNYWLF EKNQMEEQLSSEQIH VLVVPEIRDGKKRYRSSLWFAESEPLKRRVKTDESEDEDDDDVSEERTFFQLCGCPPM AHPARKDSKLMERKAYTVIFAQLVEHV KDCFEFNRTSNNPGLSSNVVVTGNP GIGKSW |

| | |
|--------------------------------|---|
| | FYLYCIFQLIRNREREDIKQLPPYELVMNYDDNFVKYDAARADEFVRLNKEDVDDLMDK PFVLRRLVDARSTKLMGWRGVSVLFAVPDAEDLHDFEKVPGPKFIMPAWSLEELQDCNQ VLPDDLKLADELVSRFDAFGGIPRYVFSKNKTAIENKLRAMASFSVKEILSYCKRGA AVKESDQSDCVLQMPSEANFRLKFYLDLSSDICEKIVFQAEGEDLTMLAKFAMGK* |
| >jgil110274le_g w1.18.174.1 | MVKFFCAIIGVAGSAFPVDIDASLSVGLKNAIKGKNDDIKCTARELQLFLAKKEKGAG AWLTEDEAAAVSLDEGGHLQGFKMMKSSLYLKNPKHFGSNFQSDQVHVLVVVPD QAQPQTGLWLVTASVENALNTKGIRCLYRLMASYLGYYPVRRRTGDKDTALWYED KTLCIHSIFKSEENALLFDNVLQDECITQTSPLDGHVSTNVAPVSRQLSELRRISRYA PQDTESPQVSMLSISTNTSIVDVMTDEFKYQRIESEWFGDVGKAQSCHLMSRDHCRKC PSDRKYDNDPNNRLALSSAMHDWYDGRMYNVPVMNISVESVSERPVIGNRYKVNLI RALNARYAKWISLILKEGFVASEDSLEMHTCVYVQNPKVFCVCMEWKRKEIDKQWKS YYDMEPAVD* |
| >jgil112192le_g w1.21.161.1 | MVKLFCAIVGAMESAISVEVGGLTVGDLKKAATDQKFDFAASKLQLFLTETEGGGW LSEDDAAIAMRTGAIPEQVKLLKDEIDPAEEIGDKFGSAPTCKTIHVLVVVEGVTNV TAATVFENNRKRREDDQPDFWMKAIEDDKVLTLPLTCEELKEHLQRELVPKIPLGGRL SQIVESQNVAGKMNSKLFDTPEARHSVTDVTVAILNGVIAPVWIGGESPTATYHFLWNE VIKVLMYVSDGTCSRNSSVFTSTGLFRPNLCFYNSANDKVCVFRGEEQARGEMTVPL TDLRQLTWRYYDDAPYVFGYAAVAQDVCLVTIRESSETNAPGEKRRKVEIIRYNLRDL RGRSLIFLALLNLSTLFRPVVGRIRPLGIAEYKTIIRPNGVKIAFGENCVVKTYPLTMPD KIISDLRDLHWQMKENAVPNVVELKSTNMRKRSVELAPVGRQLPLENVHQLLMAMRD ILRALVALHTIGLMHRDLRWENVLRYPDEDKWFLLDFDEGASSPAVKVDHLKAETHAP EILSSSTHTTMVDIWSVGYLLETSHVHDLPALEDIKTQCLQENPSVRPTAQSLEAVEA LIAN* |
| >jgil113302le_g w1.24.271.1 | MVFASQDNDFNRCNPFPSQLPTAEETDEWLEFPSSLPLTRRRSLYIRPSFKSIAAQALLK VDSNRRKYAVVTGTPGIGKSVFLYYVMWKLKDKRMLFVTRPPIYFDGESVLDYCYQ LPYAGNRNFWSPDLWCLVDATNPCKIAGLPIHHCVLLASEPRDDYVRHFRKLVPTPQV FYMPIWTEEMEKIPLYPASAASVWRDRFETLGGIPRLVLQAVQTDQPEFMRVCFYSLEN CMRLVLFHSKTKTGFIDSLYTLEMPRHVHILSQEPYHEYTLAYASETAMRAVIDAKWIV NRAEMLHFLVMNLKSTDLSLTQTTCHCIFELYAMQLELGGTFSYRSLQAGVEQSSETLD EDENDIDIPMSWREIVDRVEADQNEQDLYVPKSAKDVAIDAWMPEVGGFQIALGKEQK IKSQADELALLGQGGNRLFFLVFPRDFDSFTKQEPLSIEQYALLIPYPEV* |
| >jgil114043le_g w1.25.260.1 | MVMVKIFCKIVGEAGSIFSVKIDDESVEDFMRKIKDRCDGKIVAPWMDLQLFLAKKKD GVWLTKRDLVLEVTSELPLDDTLAPLNSVGLSEEDVRYRMTKEDIKAKKVPVHVLLV PTEDYLRSPATILLETILPHVLTHTTTLTEDNRDFRHNLCNFYGCYTREQSLVRCMLLD VPLPKSLVLASHLFRRSNEYLSFRMMQISDIDDVKNGLLLFKPLKYAFDHFQISFIRDDT DVFRLKLFSTIKETPLIDLVDHGGKVLSEEQTGELLSDVNNDTCLFDVGGKTFGDVDG CALAFTGIERPYHCLNLQARVALMVALKKGWIDESYDFKDFWSEVSLDDKMEMFHR SILNSVADI* |
| >jgil117630le_g w1.33.284.1 | MIKLFCAIVGEERNVIEVDISDVESISALKEAIKTKMPITLKYVDANTLQLFPAKMPDDK WLRDDSAADDLMSGRIHDDIKAMIADSKILRPACRIRDELSAKKTGWKWSLPWSPITP KIKTKQIHVLVKVPDMMTGTLPLCGVAPEAQWRLWQSIVRVTSYGTYSGTALVHRSSR NL YLLTNLHFWLAEEEEIEFFEHMSAGFKMVELYLKRNPRKRSRSGKQKSDGGELQQ TFNGSPVVVDQLLPGHTTLEVHSFVFKSDACWCSSVDYDYAVFKVLAPPPRIQLLGV EPSVSHFPTNVYVFGFHDGHEEKKFGHSY AIVPARIKYHRYKCLILSLSTVSLPESGVICT SRGLSIGYLAGSTIDESGNEVNQWLELGGFFRIL* |
| >jgil128631le_g w1.77.29.1 | MVKLFCAIVGDPSWSFEVDIDENVSSELNKAIKTKNKIKLKCVDANDLQLFLAKTDGP SGGKVLDRAGAAAVETDDLHRFTRMDSTLYVRNPKHFGSGFKPDEGQVHVLVVVP KWAVEAETIGQNVVERVGLPRTTALNDPKKYAEEIALDDWGVNTVHQIPSIWEFMSLS GCTKTGELFWRLEEKQVASLLLDGWFRESSPGSINQFEDMKSIMLGGSPGIGKSTLLCVM AFHLVLKYKKNVLVYRQLKGENCLLYLGYEDDEVVYFTVKRCKADRAVSIYEELGYR QGFPNVWLLLDGFRYKEIPEGLETFRMLATSQQVSLKSQESTDAYCCLLPCWSKKDLLS MGLIYNFTPDMEERFFYSGGSVREFTYATWEDIQRAMDVAVSGVEDYSKLLTTASC MFTDTSQVVRTFVENTNDRSHYFSSRYWEPMDSEYAVLALSRLKADALHRIYTW KMAGHGLAGCAFEIYLHRLAIDNRLEL |
| >jgil129711le_g | MMKLFCAVVGEAESVFHVDIEPGETVSDLKDAIKEINKHDPVLKNVTAMNLQLFLAKK |

| | |
|--|---|
| w1.86.23.1 | GGAWLSGDDPAVLELEEIEHPDILEMMDAKPMLDDKTLQFLLEFKNKLPQPSTNQIHV LVEIPAGARAIPYSNRPPKPFVSSSEGATWDFQNPDLNEQLSNAIRKHYDAWKRGYYDKI LHPLFTCWSGPGTGKSRLLEDFPKLLKDWLWLFAGKSENPD MIRLLQNAFTFNIAFDKETP HEAGSFSSAAELIGTRMLYQLQDTLKWDPFVQEKSRHSVPSDVMKLSKILGTRHKDM CVILCVDGMEKLSHENGKDCFEFYKVLTVLSYLIGTSKCWVIAICSATIYSPVKNFLLSS PQWAYEVPTAILS RPTVEGEDIFATFNQDQLIELLIDDMGGFGRALVHVM MRKARRK GSLEFMSVLTAVLAELRVLYPRIKKMASMQEAFLAVVARRPVDKYSRFGKLSLDDVI STGLVRREGRFLTCPYVLYLLDTPDPSWSKYKCYSSQETRENAPWQTW EAFNYKLR ALKSVACQGVKVDWRDIHRGARFGRGCYRVVIEEPRTYSLDVNRKTAKLDGFGEGNIFR CKYDPDQQGYFFEDAFTGVKDAESRAFHEIHQCKKIKDNL SLEDLLEEKKAAGPHDL FLLYCTSEVEGDIESLENCAVVDRTCEWKYYPFAARALYVSTVSPPDINTSAIQLQLV NGIGPAISKRIVEKRPYSSLEEAHEKTGVSMNILSQTSCRSKVDK* |
| >jgil131973le_g w1.125.9.1 | MKLFYVIVGVAGSPFPVDIEPSETVGDLTKA IKKENKCRFKHVDAYDLQLYLAKKDKG NGAWLTEDDVATVRDDAVFQTYKLMKPTLFLN NTEICGESINQCDVHVLVKAPMRLPT IHQLHYRHFTVGSVDIRTKKSMFTDPPPLVRFWRALQDRTEFKADAVLTLPEGTFLLG NPMLGSRIYIRHCYPRLWQVCLKMINDEAMNTPHLVILGNP GIGKTYFGYVILWLLRS GNTVVYESRVCHRRFLFSQDMVVVQSGSKKDFIEILEQTTTYVVDGV EPRYYSAKTILLT SPQREVWYEFNKDDCRSCYMPVWSRDEVLTCRELMYSDIPESVQDCFRRWGGIPRYV LHYATAGGRQWLEKAMENITLDSLMDACGDL YENPSEESLHLLHYRVTKEFNTDYF DFASQYVLEEVYRRLYNHNKKKLEFIDKSGVWGAAAVLRDHLFEVYTV AASLTDED GDGVAVEQFKANKL* |
| >jgil20879lfgene sh1_pg.PHYCA scaffold_75_#_8 | MVKLFCAVVG VQGSAFPVDIDASQSVGDLKDAIKTKNKIKLKNIDASDLQLFLAKPKD GPWLRSDSDVIRMRS GAIPEQVKLLNEQIDPAAGIGALFGDAKPTMEIHVLRVPDY DSDSEVNQQRKLT SFQKLRKESGATGELPVQGD FMKLFDLTDDDIGKVLDIKAIGDIVG FTGSEFYIRKEILTS DQWGSARLLVAMIAQDNPNWKNIMGELGVINCTQLSLNNDTKV STTV DASQFIARNNQLQVLPDSD* |
| >jgil39322lgw1. 83.18.1 | MVELSLQCAIVGQIGRSFDVEIDDGKKVSKL KEMIQVKNRETIKCDAKDLRFLAKMGP PTTTQLLKIWKKEST DGNSPFSPIPEFGEMVKLFYLN NMFIEEQMELPSTDQIHVLVVV PPPDV GSKRSADIEVAKVLKRLKMI EPKVLTEFIPLVPEKEFQLDNL SMVQQSDDPVM TPTLHEFWKKFGEFPLYFVRMEEVFWKVIKLLFGEDRVVIVGSPGVGKSCFLMLL AFYLACIKRKKVLVIRRLK |
| >jgil506739lfgen esh2_kg.PHYC Ascaffold_21_# _111_#_Contig6 65.1 | MVLVALTCALVGKTGVLGVKIDDSAQVWELKKAITVEKSN DLKDVDADKLELFLAKM EGGTWLDGAGAAAV ALDERGHPQGC VQMDPTLWIKNSKHFGDNFKPGEQVHVLVV VPKDENDRSAAMALGVPSLPPTTFHRH PERLKRWAANEMIRQKNQD GNEKTGTRDTN KKRKNRDIDKSM PYSSLSWNDLEPILSVE DFLKASAVPPNVVEALRDRLMQVRKLYG DVYSGKEAKRQVFIVAIIEAVCLMLGDATILVEEEVKGNV LVHGRFEFVLRGKNRVS IVEAKRDDIPQGIAQNVAGLEALSDVEGLERTLGIVTNYLEWVFISDDDEKIRRMNTTLK VYGAVPSIQELTEIVGMIYGFLANSS* |
| >jgil511381lfgen esh2_kg.PHYC Ascaffold_83_# _9_#_Contig727 .1 | MVKLSLQCAVVDQAGSSFDVEIDDSAKVSKLKKVIKEENPATITCDAKDLQLFLAKKD DAWLDGAGAAAVELDEHGHPQGC VQMDPTLWVKNPKHFGDNFQPGEGQVHVLVVV PEGVVGSASETSKMD FVVVDKVS KLYEHSVLSKRTRYVHSEMSSSKGNKLVKELKIRVT PVDVAPFTGGSPTPVEEF EWIKGRTEEQQSGRYRDYVEANIGDVL RNNKLCVFSVEKGA NILSVEVPGCDVDLAGRTDMIVLSAIVQKFPHYLPHLPGVKMLIEVKREVKSA SEFQALS ELIAMDFIVDESVMALLTNLTNHWEFLWVSNKSNRPIAATTTLTPGEAFEVIR TLLAQ SSTADADIMLPCLAEPVKRRKLNQMLPFIGEASGDGIRESIERYYDIASCLGPDFDMARA VARQVTRSIPTLSYFS* |
| >jgil527369lestE xt2_fgenesh1_p m.C_PHYCAsca ffold_180095 | MELRCGVYGE GSVFSVKIARDVKVSALQKKIASVLSTEQHTVFPRLLALYLARKKEGEE VKWLKDDRHA KDFLRGGTITEYEEMRPSWTLDD EELFGPDFQPGQEEIHVLVELPKAA VESASLVKMEKQLDEMYERIAENKRKRYVHSEMSLIKGRALLQDLKMRLTVVDTVPFT TTDASPVQVP AFEWESICDGRGQNIALTEEQQRVRYREYVENNIGDVLTTKKLCV LGVE KGM DILSVAVPGHDIDL AGRSDILM VIEVKRVLKSGCTFQALSELIALDFL VDDPVMAL LTNLTDHWQFFWVSDK KKKKHTLLAQSSSADAEINLPCIEEPVKRRKLAEVLPSVSEGG ESNGVREAIERYDYDIASVLGPDIE MARAVANQVTRSIPIYSSYPS* |
| >jgil540976lestE xt2_Genewise1P | MTIYCAVVG ENESAFGIDIDEGKSV DQLKQAIKERNEDIHVPSHRLKLF LAKKSGVWLT EIDVMEDVSDTTDLELLEAGRATLRSVGLSDEDEVGEVDEADAAAGEGPVNVLVVPIPEL |

| | |
|--|---|
| lus.C_PHYCAscaffold_50927 | YNVPSVMALFNVMLPHVLTQVPTTESDVVIEFKMELCKFYECYSRHRTWVRCMLLD VAFPKSLVSASHLFRCCNAFMAPLTVQLWDIDDMRNGLLLFKPLKHAFDHFQLSFILDD TNVFRLLKFDPSIYNTRLLDLKDCDNENVLSMEEMGVLFYNTSLTRNPCEFDQTTFGD VDGSALVFTGLKRPFYRCLNQARLARVFALKKHWIDESYNFTDFWSEVSLDDKMDM FHRSILEN* |
| >jgil556686lestE xt2_Genewise1P lus.C_PHYCAscaffold_940014 | MVLVALTCALVGKTGVLGVKIDDSAQVWELKKAIKEKSLNKLKNADADELELSLAKK GAGWLSIEDLAAIQKGEDVPGFERVSLVDTEDEAYSASIRDVLKTNMGPPPQTRQIHVL VVVPKDENDRSAMALDVPSLPPTTIHRHPERLKRWAANEMIRQKNQEGNEKTSTRDT NKKRKNRDIDSSMPYLSLSWTDLEPILTMEDFNLEASAVPQNVVEELRDRMLQVRKLY GDVYSGKEAKRQVFIMPIFEAVCLMLGDATILVEEDVKGKNVHVHGRFEFVLKHGKKR VSIVIAKRDDIPQGIAQNAGLEALSDVEGLERTLGIVTNYLEWVFISDDDEKIRRMNTT LKVYGAVPSTKELREIVGMICGLLANST* |
| >jgil558527lestE xt2_Genewise1. C_PHYCAscaffold_11463 | MMKLYCAIVGVAGSVFAVEIGEDKTVYDLKDAIKTQNKIKKVDAGDLQLFLAKKKKK GKGMWLTEKDVQKGVNNTSDFNLLGTVGAPLKFVGLLKDDVEFEPTLKDVESMNTPV HVLVAIPQQWTISKKTD AKRLEKEENIPLEMLWQYSEMEITTFQPDELSSLLQRPLPFQ LNLQKFLTPKTI FDPSPGFLVCNELSALIDGFSYSCDYRDPMASENTWQRMVDQLL DIS YRLCRAHGFDVVSNRN |
| >jgil559084lestE xt2_Genewise1. C_PHYCAscaffold_21063 | MIWLYCAIVGKAGGVFKINKGDQVWELKKKIKDENQATITCDANELQLFLAKKDG MWLPEDELVAVDLENGTIHPDIDKMMNAEQMQDNKTLQFWLFEENEMPKPSTDQIH VLVVVPKQDGLTNETSVAQTPLQDEEASAYSFSELNSAMRDQIVRKMRLENVDPVKE PEDTSIGGYSWIPKIEENESQRAGYMAYLQQHLKTLIDRGDFLLDDIADDKSVLDIVDP RLPFAMSGTADVLLINRTSKNPLIKLAGVSLVIELKKKVEPDHVPQAIGQLVSCSMKAPL NCYPLSLLTDLNDRWHFSWFSNHTLTQVTLKYPKNAFRLIEAAVLRRTESVSLPSPFIP GPFKIKVDDFLLQPDDGYAEEMMERYELMADVVEPEFLMARRAEYAQHLVQSMPMY AHTFK* |
| >jgil563460lestE xt2_Genewise1. C_PHYCAscaffold_120265 | MMKLFCAIVGARSAFSVEVGEDQTVEDLKLAIKNQNRNKLMSVDANDLQLFLAKKDK GNGMEWLTQLDVVKGVMDANGFAHLLFVDAKLRAVGLDSELGKVNKDVAVGKG HVHVLVVVPKTVGSKDGRSIQKFSAPIEFSEMEHEGGVPSTFRTERKDTVAIASLLDR IPVVFFRAPPLSGKTAMCHLLYNHIVFSKPDALVASVRANRMARNETFAEYFKKMYGC DFEEFCAYRCDRVLLIDEAQITYNDEQLWRGFVKDTLESQIPGLRLVLFSSYGSFVYRK QERPGTPILVPTDNTFGLNVTSPKPLQLSRVELEEMVLNSIGASVSDLIWVLCSGHIGIA RAVLVFLRWKFGSTTPNAEDVEMELRSEELLQYVRASYRGIPTADAFQIRVKNNDLSEE TILKMSEVLNGVASGKVTSLHDPDGGQTPRSQTAVELLTKFGFLYEDQAKQLQFASNM HLKIWLLSSRTDPIGYMVKDISHGDFIVACVQRMSASRLQKFATENTTRVARERQIQME LYGATTSCLPKGVLTPEWRTDDGKGFIDL VIRGSGILWFWELLVNGDDAVCHSKRFE TGGTNYGSLTRNCRYMLIDFRQNMGVKRKRDGFLYVSFADSFTKAHVFLDKPTVSVSE LLS* |
| >jgil567378lestE xt2_Genewise1. C_PHYCAscaffold_240430 | MVNLFCIIVGVAGNAFEVKIDDGASVAALKKEIKQENNIKLGVDAGDLQLFLAKKDG AWLMSKDLLRMWNEETPEEDERDYMSERLDDPTVRIKEKFPSEYPDRSIHVLVQVPRE LLYNQPKPRDSSNEWLAEFFNHKVEPRSLPFVGLKSSFVTQPLPAKIRVKQEWLNEWAL SPGLQEKMFVLDLDDAPCMEFTSLIFNKRTLNPFRGKTENAFISMWDSIFRNVLDVLF QAHIDRDSCNGSSTRQKRPDLFVLDQVCVFRGEEKPPDVNISVPTTEELCSKLWAYGS VPYVFGYAASGYDIQLHALCPLDPLNVGGVVTKNIGTFNLEVKEHLFQIVLVMNLSSL FQAIADCEPASGRDEFDRITRSSGWPSKPLQTMISGKNFSTNVGDKRTTTLKRASVNV DRLTNLHFQKRLAVFKPRGTMVRPSNLLDLFGALKDVLQALVALHRLGWHRDIRWSN VIRQRTGNSWFLIDFVDAATNPQQYPSGQHLVVEEHAPEIFVENGVHTTAVDIWA VGF IETSGVEWLDFAGR TSLYRRLIAKDPAARPNAAEEVLAELKALEEAAKSEKEACENSRRS ETQCRKRKLPDS* |
| >jgil570403lestE xt2_Genewise1. C_PHYCAscaffold_370147 | MEVVKLFCVIVGEAGSAFSVKGQDQEVDDLKEAIKDRSDGKIDVPRPDLQLFLAKNG NAWLSSDNDVVKALKEGVKTTLIDELTQKEKELQGESGLKQVLAGMLTPSTDQIHVLV VIPDQSKSAPSVSFEGLDRCRDSFFLQLPTAGEDDSWLMFPQPLPLTERQKLYIRSSYK SIAAQALSKMDPKRRKYAVVTGTPGVGKSVFLFYVMWKLIEKKRVLLMAEPAIYFD GESMWEIQQLPYSGNRTFWSVDLWCLVDSVDPTTIAGFPIRKCCVLLASTPRRDCIGEF NKLEPTPDVFYMPLWTKEELSTIAPLYPNAQDQWENRFEGGGVPRLVLDLKVTPQE LLQTACSNCLDGDQFEDQNGSDSDSHPQP* |

| | |
|--|--|
| >jgil573037lestE xt2_Genewise1. C_PHYCAscaff old_510047 | MVTLFCAIVGIESDAFAVEVNENDSVYALKQAIHARKMYEFSADKLLLFPKTEGPAW LSNSEDVVKLEEGEKTPLIEALTTKQCQLQAEPLVSDMLKEIDPPSLSQIHVLVVLPLDK KRKRGGTTRLSDLLETCSKEGSLPTEGDFLQMFWDQDCGKVKDITAIGDIVGFTGFR FFVRKEILCVLENLKHFKANFDRGEVGDQFIFLGSFGTGSVCLALLCFYVAATSHPV LWYRSVQYGREMSFTCLFYQKKYRWRNGAEVKIYDRLYDEV |
| >jgil573801lestE xt2_Genewise1. C_PHYCAscaff old_550284 | MIKLFCALVGAQGSAFPVDIDASQSIGDLKDAIKDQKQNDLKNVDADKLQLFLAKKGD GWLASKDLPSIQRDMASPTIFEKLPLVDPTCSIQEVLTENELPDPQTRQIHVLVNVVPQVH VLGKRTRADEWFQIGMKSRVIVDGNETDRITRYFEMAGFPPLAHPKAEYRKILERN AYIVIFTELMKKAKLSFEKGADCSLVVTGNPGIGKSRFYLYCIFTLFFALTWKLRSPPSI |
| >jgil96357le_gw 1.1.1010.1 | MIKLFCVFGAAGSAFPVDIDASQFVGDLYAIVEKKKEDPNLKSVTAKNLQLFQTKTE NGDGWLSDDGAVIAMRTGAIPEQVKLLKDEMDPAEQIGDKFRNAPTCKTIHVLVVV PPAPENERKRKRMEDEVAPDAWIKAIKDEPVTTLPTCEGLKHLLRALHVKIPINRFL QIVSAQNSTGELFTVLEKLFEPQPRNVSDITGAVLRPIIDPLPSGPTTKSSYHHFWDCVI ATLLKVVTGNYHRSTNASASTGAYRPMCFYSRKSNICVFRGEEKANGELDVPMAEL HEKLTWRYDDAPYIFGYAAVGLLVCLVTIQKDEKTSRAKAEKIETYDLGNLKDRLFL LALLNLSTLFDPVVDLIRPLGIPEYITRERTNGVRIEFAEDCVIKTYPKNMPSDGIIRNLKS LHRLMKEHSVPNVVELKNANKKKHVKLAPIGIDRRPVNVQQLMALCDILKALVAL HAINVMHRDLRWENVLKYSTEGDKWFLIDFDEGRLSAATVTHLKAESHAPEILSSSH TVKVDIWSVGYLLKTCCLQDLPPELKRISQCLQTDPSRPTAKSLLAGIESLIES* |
| >jgil100678le_g w1.5.1392.1 | MVKDIKLRGAYGEGSVFSVKIKQADVEALQLAIVNARKGVNRFNVDPSTLTLYLA GKQEGEEIKWLKDEDSLDELRGVYPKQYMKMRSSRILDEDYFGENFQGRHDVHVL VELPEKVSSVQSVVRAVFWLVTGLVENALQTRGVHRLIYRIADAQLGYDPANMLPDN KPRAFWYTNNDLQFHVLFKEGECVHCTCWC* |
| >jgil102004le_g w1.6.669.1 | CVIGVAGNAFVSNIDENLSVGHLLKAIKGENVNDPTLKNVAAKNLQLFLAKAEGGARL SSLTDEASEDATKVKKGEWLSNLTDDVKLKKGEKTPLESVTHENKGLQGESGFKRV LGPFLAKRKEWIWL* |
| >jgil106201le_g w1.12.351.1 | IKLSLQCAIVGQTGSSFDVEIDDGEKVSCLKEMITEKNKQDPNLKNVAAKNLQLFLAKK GDAWLPDDDPAAQDLEEGKIHTKALIDGNMKKEAWTIEDVLVDNNMTGEGRAPKS RQIHVLA VVPGILTTIERERVDENQD* |
| >jgil108281le_g w1.15.479.1 | MKKVSLQCVIVGIGSSFDVEIDDGEKVSCLKMRAIKDRKPLTITCEADLLQLFLAKKGN WLSSTDDVKALKKGEKTFIDELMHEKEKMEEEYPLSDYLANMNDPEVKQIHVLVV VP |
| >jgil108895le_g w1.16.149.1 | MAKLSLQCAIVGQIGSSFDVEIDDGEKVSCLKDAIKTKNKDDPILKTVAKNLQLFLAK QGNAWLPDDDPAAQDLNEGKVHTEIQALIPDEHRAALKLVNGESDDYINALTAGEQIL ASKTIETWLYEKIKMEEPSTAQTHVLVVVP |
| >jgil109378le_g w1.16.566.1 | MKLTLQCAIVGQTGSSFDVKIEEGQTVGDLKEAIAVDQKFGFAASKLQLFLAKQPVEDD DGKEVVPVYHPCAEGMKKESFKWLPDKHRAALKLVKGESDDYINSLTAGEPILASKTL TIWLYEKNMEDPSTQQFHVLVVPE |
| >jgil113678le_g w1.24.344.1 | MNRLERYAALNEIVQEKNAENGKTSNQDTNHRKCKKALDNSVPFSSLSWDEIEPVLQ LNMFLHTAKPVPDEFVRKILAQLADLHQLYGDVSTGKEEKRMFTMTVLEAVCLHLG DVMIFVDEELTGTKIHMHSIEFVLQRGAKRVPIVIARRDNVEQGMACVACVEVLAD AEGLETFGIVTNYLHWIFIRDEDESIELIDQPLNASMPSFESLKVILGMICGMLESE* |
| >jgil114206le_g w1.25.261.1 | PLDDTLAPLNSVGLSEEDVRYRMTKEDIKAKKVPVHVLVVPPADSPATILLETILPHVL THATTTFTEDNRDFTHNLCNFYGCYTRQSLVRCMLLDVPLPKSLVLASHLFRSNEYL SFRMMQISDIDEVKNGLLLFKPLKYAFDFHFQISFIRDDTDVFRLLKFDSTIKDTPLIDLS RYGKKVLSEEQTGELVSVADNGSCLFDVGTFGDVGDCALAFGTIERPYYRCLNLQAR VALMVALKKGWIDESYDFKDFWSEVSLDDKMEMFHRSILNSVAEF* |
| >jgil117855le_g w1.34.341.1 | MVTIFCVVAGPGSVFAVDIGITQTIDHLKQIKENKPNMIRFDADLLKLYLARDGGAWL NSNDDDFKALKRREVPARIKNLMQEQLLDETAKLNDDDYFGKHFPGDRDIHVLEVEL PEDPTEVLHYKSELVCGDCFLAELLVVTNSFRL* |
| >jgil120859le_g w1.42.251.1 | MVTVFCIVGVPGSVFSVKIDENESVAELKKAIKKETPNIFQCNAMEDLQLYLTKKGNV WLTEAHVKEGLRDTSLGLKLLNSMKTCLKFLGRDENDEEGEEEGMGLVDVLV |
| >jgil12221le_g w1.47.236.1 | MVTVFCIVGVPGSVFSVKIDENESVAELKKAIKKPNIFQCNAMEDCLSDGEGRRVAD GGSCERLERHQWVEAVGFNEGETKSSGVEERKRRRR* |
| >jgil124124le_g | MGEELYDPTDKISAKFPSQIPGGTIHVLLVPEGGKRHLSNEWFTESFHPLKRRVGED |

| | |
|--|--|
| w1.52.167.1 | VNKEKRFFDMRDFPSLLEHPKVEFKTIVEREVYVVVIFSQLVQYAKTCFEFVPTSASEPDG KSKPGKDSNIVVTGNPGIGKSRFFLYCIFQLILREREDVALLPPYELVNNHKTNYVKYDA VSKEFVELNKKDVRALQRKPYVIRLVEATSSSELTGWRGVSVLFASPGVDGIDNFSKVDG LTFIMPTWTFEELEDYNSLLSDELKLAEDELLSRYDRFGGIPRFVFSQIMDQTEAKIQSAI ASFSALDVISYCRKNDVREKDYSHCVLEMVPTKADFRANFYLDVFSMHIAEAVIDKV HGDSLAKVSEFAV |
| >jgil125925le_g w1.60.118.1 | MTELKLFICALVDTQAVFYVTINGEKTVYDLKKAINEKPNLEKVDTAMLQLYLAKK CDGLWLTENDVKNVGSSTAGLTLNAAQAPLQDAILGEGLEHPSKEDKVAGNGPVHV LVSVPESVG |
| >jgil127207le_g w1.67.186.1 | MMKLFCSSVGVAGSAFPVDIASDETVDGLKEAIKAKKMYCFPADELRLFLVNTSVKNP DEEEKKAH* |
| >jgil127255le_g w1.67.176.1 | EDEDSSIDPYPWESDLAEDHEGQRKGYMQYLKDNLHGVLSEGGSTTIPPGTSTSGQPKY HLKDTSRMTSLLTCKASSLPFGLKGTADLMIIGEV AHSRNDIFADLQFVIKIKKNQC GPK ERKELLELVAANWKSYSPCAPIGLLSNLNDYWFYFMWFTTDRKIARMKLSCPANGLKA M* |
| >jgil127347le_g w1.68.101.1 | MLNLLCAIVGAQGSVFPVLIGESVGDLLKAIKMKPLTVTCKADRLQLFLTKEGGG WLSSKDEVVLATKGGIPEEINKMLIDEIDPAKEIADVLGAAPTVMVIHVLVVVPRGLY TIHGDHVTARTDGWQSFSPTKLRLTRCRLLEV* |
| >jgil12764lfgene sh1_pg.PHYCA scaffold_1_#_21 6 | MLLNCAIIGGGDVISIIIEWKTVALLKDAIKEKPKIKLNDVDAGDLHLFLAKKDGAWL MSDDLQMRREDGKGERGYMSEELKDPVAKISAKFPSELPGSIHVLVVQRFRIEDVIDIL AELEKIDDVKNMTVILCVDALQQLVNDDAKTRNGQFRRSYHACSDDMAAI* |
| >jgil128012le_g w1.73.136.1 | MKVSQYVVVGVAGSAFPIDIGKNLLVGHLEAIKEKHDDIKCPARDLQLFLAKAGGN AWLANSTDHVKKLKKGEKTAYIEALIHENKELPRKDPISKYLERMDEPQMEQIHVLVV VP |
| >jgil128338le_g w1.75.57.1 | MVKLSLQCAIVGQIGSSFDVEIDNGEKVSKLKEMIQVKNRETICDAKDLRLFLAKKGD AWLPDDDPAAQDLEEGKVHTAIQALIDGNKMKKEAWTIADV* |
| >jgil128353le_g w1.75.129.1 | MVKLFCAVVGQGSAFPVDIDASLSVGDLDKAIKTKNKIKLKNIDASDLQLFLAKPKDG PWLRSDDSDVIRMRSGAIPQVKKLLNEQIDPAAGIGALFGDAKPTMEIHVLVRVPDYD SDSEVNQQRKLTQKLRKESGATGELPVQGDVFMKLFDLTDDDIGKVLNIKAIGDIVGF TGSDFYTFARP* |
| >jgil128403le_g w1.75.131.1 | GSAFPVDIDVLSVGDLLKAINVEKTIKLKNVDAADLQLFMTTKTKDGQWLRSDASDVI NMRSGVIPEQVKKLMNKWTRQMKLANCLVKNHQRRQFTCWWRLQRLI* |
| >jgil129581le_g w1.85.127.1 | MVSITLCCVIVGVVGESEFDVNIAGKSVHQLKEVIKAKNKKLENVDTRELQLFLAKTA DGTWLSLTDHAVNSLRNGIIPTKVKALLKRE |
| >jgil129660le_g w1.86.101.1 | LWNDEVIVAARIPREKVLMMHERISRGGFGEVYVGVYNGRKAIAIKMLLPEIRKRIQSVNE FLVEVKLMAALEHPRIVEFIGVAWDSLTDLCVVSELMERGDRLALLSQFAENHPHGF DHDVKVIALHVAHALTYMHSFSPPIVHRDLKSKNILLTNEFDAKLTDGASRERVDRTM TAGVGTSLWMAPEIMAGEKYDEKADMFSFAVVLSELDLHVLPYTKLRQETRASDVAIL QLVLQGKTQIDFSDACPSSIAALGMACA AKDPTARPTAAQALYELQQTLSNEFY* |
| >jgil129685le_g w1.86.155.1 | MVKLFCAIAGIAGSVFLVEIEEALEVDLKNAIQKEKPLTITCAHQQLFLAKKDDGK GAWLTEVEVKNVNDTTGLKPLDAVRAKLKNVQLSDSDVGGVDEADEVAGKGPVNV LVML |
| >jgil131615le_g w1.108.42.1 | MKVSQCAIVGQTGSSFDVEIDDSEKVSLLKKAIEKQEKISFKDVAVDLHLFLAKVQK DMTWLYSRSEDVKKLKKGEKTPLIEALTMERHELQGEDPLENVLNGIDPPSVRQIHVLV VVPKG |
| >jgil131976le_g w1.125.17.1 | MDVRYESVAHYFKPINGTRVLKSIENCGEDFEPDEGRVHVLVVAPVTVLMQLDPPYCH FVVDGVNIPITDNMAFNLPGLTGFWKAFQEVDTIEIANTAIKLPEGTFLLGDSNRGSCIYI RSCYLQLWEITQKVQDEVKATNLVIDGNSGIGKTYFGYVMLLYLARLGETVVYESY GTKKRVLSSHNVVVEGSQQDFSDILNLPFTFYIVDGVEMPHYQAKTIFLASDHTLWYT FNEKRDQIRYIPVWSWDELSTCREVLYSDVPESVVEGCFHRWGGIPRYVLQY AQSEEK QILLEKTMEIADFFWLLNGYEKLNANNPEAHRLLHYRVNDHFKEYFDFASPYVQQEV YDRAYKKDKRVLLGFIGGGDGWKW* |
| >jgil133238le_g w1.380.5.1 | MQFLAKTEGKWLDPNEDLDTLLTLHRICTCSHPTSWKLSNPDLFGPGVSLGEDVVH VLVVLKDAEAGVASVELSALPTVKQRHPERLKRWAANEMVRQRNQRNEKTSTRDT |

| | |
|--|--|
| | NKKRKNRDIDCSIPYSNLSWVDLEPILPMEEDLKLEECRPSKCSSRST* |
| >jgil13996lfgenes1_pg.PHYCA scaffold_5_#_210 | MVKLSLQCAIVGQIGSSFDVEIDDGEKVS KLKEAIRTKKHLTITCEADQLQLFLAKQPVE GDDGKEVVPVYHSSAEEVKEESFKWLPDEHRAALNLVNGEPDDYINSLTVGKQILGSK AIATWLYTKNNMELPSNEQIHVLLVHHLTQFK* |
| >jgil15135lfgenes1_pg.PHYCA scaffold_11_#_121 | MMHNLHGEAGNLKLTAELESPTRLVTDVVTQWLASMDMRDEEAAPGRTSPKTQME KSKDEERDEEVQKKRPKRKFRKSTINVRKEEK AHLLEELSGIAYQDGGDQSSSICHIHAL MSEYYLFNSPARGKMKTLTCAIVGVAGSAFPVDIDANKLVGHLKDAIKEK KMYQFPAD ELQLFLAKKADGAWLSSKDPEVISMRS GDIPQVKTLMNVEVDPTDDIEDVFEGAPTK TVHVLVVERR* |
| >jgil15224lfgenes1_pg.PHYCA scaffold_12_#_49 | MSDMVFLMLLFEHIVICVSLSSSIFDAITEKNKDPILKNVTAKNLKFLAKTESGWLS YDEDLVNKL LLLNRVDTSNVTATIVDEEREKFDLLSEKKGCLNVLFICATKYGDNLQE AFGPAKSVIYTEAMQSTEKKATTQYIHEVILLDLTTPQNRAAFFGLAWDDTLQTRLENV IHKAGGGNTAE* |
| >jgil20945lfgenes1_pg.PHYCA scaffold_77_#_13 | MVQISLQCAIVGHAGSSFFVVGIDDGAKVNKLKKAIQRENPLTITCEADQLQLFLAKD GK HFGSGFKPDEGQVHVLLVVPVCAVAGSKAETIGQNVEMNGDGEAGVDIFSL* |
| >jgil34764lgw1.83.9.1 | MMKLFCVIVGEAGSAFPVGIKPEDTVGDLKEKIKGKNTMTITCDAKDLQLFLAKTHAG WLSDNEDLDLTLQSEIDSSYL RMRASWKL SKPNLFGPGVSLGEDVHVLLVVPPEG |
| >jgil49590lgw1.51.244.1 | MLELLCVIVGVTEWAFSVEVDETKSVDNLKGVIKNAKRNALTGINASDLQLFTA KTTD GKWLESNDPDIKMTSGDIPKQVKLLKDVDPVKDIGSVFQDAPTTMTIHVLLVVP |
| >jgil502542lfgenes2_kg.PHYC Ascaffold_1_#_2_#_Contig261.1 | MVSSPLPSSTLDALAPPFYPSMTWYPSITDDYLERPHSPGKSGFVIADPVEPMAEIPDEE LFDPAFYPLSAMEMQELEQVDEINEILAE LDMESHQELHYKLSKTRRELRSSSDVDAEI YSMMAKASKAKNFSKQHVHLHKNTSFH SKRNMRSALHQPRSVK* |
| >jgil505958lfgenes2_kg.PHYC Ascaffold_17_#_43_#_4096633:2 | MAASPNK KIREASQLDQLKQFTTVVADTGD FEQINKYKQP DATTNPSLLFKA AQMEQY SALVDDAVAYGKGLSPDLSESERLGYVIDKLSVNFGL EILKVPGYVSTEV DARLSFDT EGTIARAHRIELYEKAGIKDRILIKIASTWEGIQACKHLQKEGISCNMTLLFGFAQAVG CAEAGATLISPFVGRILDWHKAKTGKSSYESHEDPGVSVTKIYQYYKKYDYKTIVMG ASFRNTGEITELAGCDRLTISP NLEELTKSTAKLDK K LDAETAGKAYTGEKLSYDEKDF RLSMNEDAMATEKLAEGIRGFSADIVKLEQILKAKLSA* |
| >jgil508616lfgenes2_kg.PHYC Ascaffold_36_#_69_#_Contig4695.1 | MVKLFCIAIVGVTGGAFEVKIGYTKSVDHLKDAIKTVNKITLKDVDAPDLQLFLAKPKD GPWLRADNSDVISMRS GAIPQVKKLMNEEMNPAARIGDLFGDAKPTMEIHVLKVKSV SS |
| >jgil50894lgw1.79.118.1 | MLKLFCIAIVGVAGSVFAVEIGEDKTVYDLKNAIKTQNKIKLKEVDAGDLQLFLT KKKK KGEGMWL TEKDVQKGVNDTSDFNLLGTAGAPLKFVGLLKDDVEFKPTLEDVESMNTP VHVLVAIP |
| >jgil509061lfgenes2_kg.PHYC Ascaffold_41_#_57_#_Contig1845.1 | MGQGQSSIPTTEVAALVELYDALSGDRWRRRDGWKQPTRDPEQWFGVEVAMGHVVA LELPANELSGCLPVASLARLPNLRVLDLSKNQLRGEIPAE LGQLTALKRVDLSCNDLSG AIPRQIGACNQLQELNLYQNSLSGTM PKELGKLQSLRTLQLQHNNLCGALPETLCELTQ LTKFSVRGNCLTGRIPTDIGRLQSLVFLSLRNNELTGVI PPSLGCCCKALEFLNLSSNQLSG PIPETLGELEDLEYLYLFDNALEGRVPGSIARL KFLKESDFRDNRLRGELPNFLDGCSSLE AVMTKWKNRKASYRHAILGDPMPSPDTPPTSSHQFLQTLEDPPSNSSATFLSQSFDHKG SEGLGDGAPEDDPANNSHLVAEFSRKR VFQLPDSV VQAK* |
| >jgil509444lfgenes2_kg.PHYC Ascaffold_46_#_29_#_Contig2750.1 | MFWSAALDLKTQRAPNRLSYGSFNRSASNSSFHIPQSIPEHQPVSQFTSPLPKRREEYV ERPDTAFARTRASLQLSEHTQTALKMLLQGEREFK KPSLGRTIENKPKPKFAQDMYSMS KISSYSAHRAPMKELPERTPLLKPIARAHGQEEP NPQENSNEHNYWMQELERRRAGLRR SRRRTCTPM SERATTSPLKGVVVMCSVPLVVLVGFVLFIFLFGGSDASSNVMDVMNN LLGIGRGDDAETIARHAQDRDRIASTFARPPEKIQLEESIVMQLDVSSDHQKLRGSQKE KAGE* |
| >jgil509452lfgen | MESVVPCDLLDEL MHGVVILHHEAALVLGTTQFSVETFFPLLAYVLVHCRLPIIHAQLH |

| | |
|--|--|
| esh2_kg.PHYC Ascaffold_46_# _37_#_4103400: 2 | LLENFAITADNANGEESYYVYCVHAAVEYVCNAAGLSGSTNPLGPASSASSAVTTPGG CGLTPVAPMSSTSPPKQAAFSLDLELEMENELDVKVLGLSRDSGEEQPEESGTSALQ* |
| >jgil511091lfgen esh2_kg.PHYC Ascaffold_75_# _1_#_Contig233 .1 | MKLFCAIVGVAGSAFVSVEVGEDQTVDDDKDAIKVKNDDIKCPARELQLFLAKKDKG |
| >jgil52465lgw1. 36.563.1 | MVELFCVIVGVTGSAFVSVKIGNTKSVVDHLKDVIKTKNKITLKDVDAPDLQLFLAKPKDG PWL RADNSDVIRMRSG |
| >jgil52481lgw1. 67.119.1 | MAKLFCVIVGVTGSAFVFLVDIDQAESVRRLLKAIKAKKMYQFPSNELQLFLA |
| >jgil530721lestE xt2_fgenesh1_p m.C_PHYCAsca ffold_750003 | MVKLFCAIVGVAGSAFPVTIDEGQSVGDLKEAIQVKNRETIKCDAKDLQLFLAKQPVEG DDGKEVVPVYHRSAEEMKEESFKWLPDEHRAALKLVKGESDNYINSLTAGEQILASKT LTTCKST* |
| >jgil532971lestE xt2_fgenesh1_pg .C_PHYCAscaff old_90155 | MSISEACTCSLQARLHQVENARSREMKAHGTLVQSLEIELESRRICEGESRLALAKLRGV NCKLEAELQAAHGKARQLLEQLEAQRVETSKCEERLKVFEESDCQLRAAREKSDNLM EEKRQLNVEVSEVKACLEKAKAENKVLHAKYEEKISEANAKTADIVQLTSQITDKQEI EVATERSQLLLRELTTRYVQQELAQERIKLLEKDVNRTMVTRNRALNWRKRQELWE NQFISHRIFLNWKLRSAAQAKFQSAAMARCNDTRSLQLIEMNVAQCSSAKEELNR VRQACINECVQMQLRREICQTELPKLLTIVRNHQRSDNQQRHLEMQLVNQRESFERE RERLNMSRQDFEHCVQKAIEEKRLFQKRQH HAMFQVFTSKKKWELHRRVFGAWKEFY LRSIVGHATHAMVFQSQHLRASTSEVSRPIRRGESHQWTMPAPIRRGESHQWTMPAPI RPISLASQHWRRIDRASS* |
| >jgil533583lestE xt2_fgenesh1_pg .C_PHYCAscaff old_150061 | MMSEEQAATLSEDDGQLQRLETEKRELIDKLLKAVVAGKAMKKQLDDTRAENVLMD EEKQDIVVKLQEVVTRYQAIQQELESKEGELMLAQSKVEAFQVELQAQQEADAKAIKE LTTKFSQEEARVMVFQEENNRLTSQVAAMLEDKQQQEKETQTLMQDNQSLQNKVDDL MQQVAGARQLQEDTEQNYLEVASKLNQVRENDELKNKPGTDASVELQSQLRAVETE LAASADKWRQEQLEQQIETLTAVNKNNTTELADVADIREKLLVHV GIDLTYASIESL LDTKNNEIQMLTEQLAAAEAPQEIQIREADKTVEDLRAELERVAEEHSLSLVTMENEYR LACEELKTEAEKLTGQLDAAQTQTQQLNDDFSVERIHKSEENRLLVEKLEECEAALKA RASEIEKLVARARTASSD TDKSGSSEMAACKAENLRMIFEVAKTADSVSKLKL DHEELL EAHRKKSSEVDTVLVQLASLESANQTLES DLKKKSEELHNHLETCSMQKEDFESVVAN LNNAVIAEKEKSKIKRDLEDMDKTENDVLEEQISELHAVAETKTEPDESQEKDREVEEL RSSLVQAKVDFLEKQQLTALEKLLVAVSKSSGPTCTVNSTSLDAERREFEAALIEIE MEKKLQVAYEAKQGLESTLQERMEAKTDLETRL SIAEDKITELEQQLEQKVALIATIEE QLSRGKLEEKAI EFETFKTTT NMLKDESR LFN EIALLDKDKIAKSEVQKAAMADSQELA NEELEEQNLADRIAIEIAEKQDLLARLDET VYRSEEDIHQRLRERLYMLEEEKSGLDDE NFKLERTIEHLESKLD SLEEQA GLEAANEASSQQLSLEERVEKATSEIATLSAEKDSL VELQQSLEENVSVLQEDKVKLEQTLDETSSKLRDELDRVTEQMESFQAQLAQSTAEKE EVTIALAELRERSEADKLASGEVTAKLEAQAENQTLENKISVLKQMAEKALQSLQSSR DELSEAEARASVLVEERDAVKVLLQEKQTA YEQLKTQREELQLRVEQLSSELESSQKRR TEEAIAAEETIQTLKLTEAKL TESLESVKHDLAEAESAVMVLVEERDAARKEITARDLKI EMM TSQQGELDLAAQKLESELSILRSKSSADSEATKEMLRDLEESKAQEETVRS LQGS SAQAKESLQSIQEQLAESELRVATLEKERDARTSLNENMSTQETLSALQEDLQKKVDA LEAELKELRETSSAELAAANETIENLNKNAEAEAEETLAALRQELAEAEAGCVMVLVEERD ATKKTLSKRDLTLEVLSSHEGELQLKSQSTTTELENLRSKSAADAQAAEEALQSLREEL KRATESLESTQNELAESESHVAALTA EYDMVTKAMADVQSERDLSGLNEELKESVED LKNEMENRQKQFSSELES AEEAIRLTKTSEVKMTASLDSVEQELCEAKSSASLTAEYNS ILATQKEKEAGISKLTSEIDSLQERIQALESELQELRNQRETEAQTAEETIRSLKESESQTV ETLEAIRVKLSETEARVVSAD EEREAAASKALSEMESH RDAQSTEVE SLQERIQALESELQ ELRNQRETEAQTAEETIRSLKESESQTVETLEAIRVKLSETEARVVSAD EEREAAASKALS EMESH RDAQSTEVE SLQERIQALESELQELRNQRETEAQTAEETIRSLKESESQTVETLE AIRVKLSETEARVVSAD EEPLEIGSLQERIQPLESELQELRNQRETEAQA AEETIRSLKESE |

| | |
|--|--|
| | SQTVETLEAIRVKLSETEARVVS ADEERE AASKALSEMESH RDAQSTEVE SLQERIQALE SELQELRNQRETEAQTAEETIRSLKESEAQKVVELDSVKKQVLLSESSSSALD LASLR EQVWAKDESHAADIASKDEVISTLKSLEEVM AAYKRLKGHLHELQDRLTQQTSTNDT LKASYEELSGQKIAAVEELEALKIELASSEHSNTMAEELRQAEKVAVAEAQRESQME NFKKRVLAYDDELAQMOKQHAVALQE QKETLLKTS AVREEAAATKLLEQKHDQAGT SHEEASQKHEAERMELESQ LSSANETIEKRL LASVTSVGLEKEIAELKLQVETETEGANA ARAALETYKKRAHTALKKASSENKLN LKKTSEATAKLEKELISAKSRVNTLETELEDTR KRMAEVESAGDLLAQSTREAVESEKRSLEVTLRLEIDSLKAEVNRLEEALENDRQPLEA QINQLSERNAALNQDVITLKEEIRSQTESMEQEVHTKEEEIRDLSKQLQAALAAAASLAT NEAGRRYSPTYSPTTEKERRSTASSRSFSDGNN SFLHRSSIEEQSEHMAAAVADSCPIP LASKMAPANGVNSQTDDEDEV CRLKLQ LNELETASHLFQKKYEDTSALLEEANQQKQ RLQELCDGSTQAINIEYLKNVIMKYIESQVPSEKEQLVPVISTLLSFTPQEQQKVM AVHR PNDEGAGLFGGVFSLFGGAAAAPPKPLAAPLNFKPSPTTAMSNTTGAALGSKDKNG VLSFGSDPSDDEEFATPLNPF AA* |
| >jgil537103lestE xt2_fgenesh1_pg .C_PHYCAscaff old_740004 | MTVKLFCAVVGFGNIFPVDIDMTDTVEDLKYKIKEKPNAIYFDASLLKLYLAREGDK WLNSTDEAMRAVKARQCPDRIK NMQEHL LLDAMNFQTPDGVIVHLVEYPREDELG SFLNWFPCWSSLQSLFSVEKEKRA* |
| >jgil537210lestE xt2_fgenesh1_pg .C_PHYCAscaff old_790005 | MRPDFLLHYLGMVLLRGEEKSATTEIDE PVKELTAKMSW WNP MFYGDLPYILGYATSG ARLRVVMIDRHLHSDAILEFQSIFQQR AEVIKLFYNLAFFH KMSVLAKRTCPSSLM PPT PDVNRKRKIELMDDVIVRTIKRNQCRDRVDFRRLADIYATLQELNNRVRERTHLQIVRK LRLKRLRLRVQLSPLGTVRPPMNVDEVRVWLQGMLTALKYWHSCDYCHGDLRWSNI VYIPVSSDSGFVVLIDMDESRSNTTTIDWNHEFQGYTLIFEHDLFQLGQLMNSFTFSLP SDLEDVQKALLTAVHTPAGDLPTILLNQLHD* |
| >jgil537308lestE xt2_fgenesh1_pg .C_PHYCAscaff old_830012 | MKLFCAIVGVAGSAFPVTIDEGQSVGDLKEAIQVKNRETIKCDAKDLQLFLAKQPVEGD DGKEVVPVYHSSAEVKE DSFKWLPDEHRAALKLVKGEYDDYINALTAREQILASKTI ATWLYTKNNMELPSNEQIHVLV VQGRIEDVIDILAKLEKVDDVKKMTVILCVDGFQK LVNDGEKTFGSTNADQETVDAVDRVCLNKKSKIAKIIRGMYS DLSDEENL FSEP* |
| >jgil537717lestE xt2_fgenesh1_pg .C_PHYCAscaff old_1080013 | MFEKTMKGFGTDEDALSATLVRYHGV LNDIRPVYKKKYGKELRDRPVYKKKYGKELR DRIHGETSGKLLALPTACAIVGQTGSSFDVEIDDSEKVS KKKAKIEKQEKISFKD VDAV DLQLFLAKVPKEKQGEVERTEEVKDEVQEDMTWLYSRSE DVKLLKKGKETS LIEALTM ERHELQGEDPLEDVLSGIDPPSVLQIHVLVVVPKGEDV VLA* |
| >jgil544930lestE xt2_Genewise1P lus.C_PHYCAsc affold_160416 | MPEPPALQTLDIVDEDGQHQQVVLVGGIEGGKQTVEIANEYGGFTTEEVTAVKTSEGEK LVIEDAQGQATVVEVHELPAEVAATVTEDS QVVNAV TADNQQVQIVVEGEPEHGEQT VQLIDEHGDVTVEQAKAIETSEG VKLEIKTDQGPVS VVVAEVPEEIKEEVIKEQEEHQPG EFGGNPFTEEEPAVLQTVDVVADNGEHQ QVVLVGGIEGGKQTVEIANEYGGFTTEEVT AVKTSEGEKLVIEDAQGQATVVEVHELPAEVAATVTEDS QVVNAV TADNQQVQIVVE GEPEHGEQTVQLIDEHGDVTVEQAKAIETSEG VKLEIKTDQGPVS VVVAEVPEEIKEEVI KEQEEHQPG EFGGNPFTEEEPAVLQTVDVVADNGEHQ QVVLVGGIEGGKQTVEIANEY GGFTTEEVTAVKTSEGEKLVIEDAQGQATVVEVHELPAEVAATVTEDS QVVNAV TADN QQVQIVVEGEPEHGEQTVQLIDEHGDVTVEQAKAIETSEG VKLEIKTDQGPVS VVVAEV PEEIKEEGPVSVVVAEVPEEIKEEAAEDSMQGSKFMTKPEFREWIRKHYEDRLES LKVEE QKLEDEERANAGR VKGLEDCIEQASNKFGY YGVYEQAPYYQNAVDWVEEECWTAQQ WEPESPYEHPNLRR* |
| >jgil547387lestE xt2_Genewise1P lus.C_PHYCAsc affold_240409 | MSERLDDPTVRIKEKFPSEIPDRSIHVLVQVRREFLYNQPKPRDSSNEWLAEFFNHKVEP RSLPFVGNLSSFVTQPLPAKIRVKQEWLNEWALSPGLQEKMFVLD DDDAPCMEFTSLIFN KRTLNPFRRGKTENAFISMWDSIFRNVL DVLFQAHIDR DSCNGSSTRQKRPDFLFLVLD QVCVFRGEEKPPDVNISVPTEELCSKLVWAYG SVPYVFGYAASGYDIQLHALYPLDPLK VLTKNIGTFNLEVKEHLFQIVLVM LNLSLLFQAIAD ECPASGRDEF RDITRSSGVVRLSP TFVEKIFPDISTFGHLELVYGH LKRASVPNVDR LTNLHFQKRLAVFKPRGTMVRPSNLL DLFGALKDVLQALVALHRLGW IHRDIRWSNVIRQRTGNSWFLIDFVDAATDPQQYPSG QHLSVEEHAPEIFVENG VHTTAVDIWAVGFLIETSGVEWLD FAGR TSLYRRLIAKDPAA RPNAEEVLAELMALEESAKSEKEACENSRRSETQCRKRKLPDS* |
| >jgil563418lestE xt2_Genewise1. | MVKLFCAIVGVAGSVFEVDIDDGASVAALKDAIKNQNPKNLMNV DANDLQLFLAQKD KGNVVEWLTEKDVQGGVSDTSDLKLLARARARLRRVGLSGEDIVEVDEQIEAEGGGPV |

| | |
|---|---|
| C_PHYCAscaff old_120143 | NVLVELPPGTNRAPLSDGTDLWLSRFHHARISVLPTLGDLEGEYIEHLSL* |
| >jgil576078lestE xt2_Genewise1. C_PHYCAscaff old_830005 | MVTIYCFAVDVEGSSFDVEIDDVAKVSKLKEAIATNQKFDFAASKLQLYLAKKGGKGG VWLTEKDVQEGVSDTNDLKLGAAGAPLNLVGLSEKDVKFEPTLEDVESMNTPVHVL VVIPSPDVGPKR |
| >jgil576081lestE xt2_Genewise1. C_PHYCAscaff old_830013 | MKLFCAIVGVAGSAFSVEVGEDQTVDDSKLAIKNQKPNLDKVDREKLQFLAKQPVE GDEGKEVVPVYHRSAEEMKEESFKWLPDEHRAALKLVKGESDDYINSLTAGEQILASK TFTTWLFKKNKMELEPSNEQIHVLVVVDPSSNAALEYKDPMSNLRKRGAEVMTDVMM EDDAFVAEENRVRVNRVRCIIVGIAGQSF CINIESNARVENLQGAKKSRQRQLKDVEA DDLIFLAEKADGAWLSADGAF |
| >jgil577802lestE xt2_Genewise1. C_PHYCAscaff old_4740001 | MVELSLQCAIVGQIGSSFDVEIDDGKKNLKEMIQVKNRETIKCDAEDLRLFLAKKGN AWLPDDPAAQDLEEGKVHTAIQALIDGEKMKARTIADVLDDNNMTGEERAPKSRQI HVLVVVPPQWTVKRVTKPDDLSRAKRARIVRLEVAYQGPVPTDFYSVPAETIDTYQQ LKEAFLSHERLVRPLCLLYGPRRFGKTTIGHRLVSLLEDAEPSILVIYCSLTPLSVESEEAF WVALGEFIGEHTRSFQEF |
| >jgil59630lgw1. 67.177.1 | DEMDSVAPYAWMETNEAVHDQRNAYMQYLKDNLHGVLHQDGSTTTPHAIRQPRYH IKDTSMKTLNCTPEALNHGLKGTADLMIVGETAHRMNDVFTDLQLVVEVKKGGQCK AYEQQLTLELVAANLCKEKRCAPIKLLTNLNDYWC FMWFTPDSKKARLTLSCPANGF KAIKDFLAGAVDA |
| >jgil60673lgw1. 22.466.1 | MVKLSCAIVGVPRNVFTVTIEDAASVSALKEAIKTEKKNV MANFAAEDLQLFLAKK |
| >jgil70314lgw1. 16.564.1 | MVKLSLQCAIVGQTGSSFDVEVGESQTVGDLKKAIKTKEPLTITCEADQLQLFLAKQPV GDEGGKEVVPVYHPSVEEMKEESMKWLPDEHRAALKLVKGESDDYINALTAGEQILAS KPIATWLYTKNNMELPSNEQIHVLVVVP |
| >jgil70333lgw1. 32.357.1 | MMLSLQCAIVGQTGSSFDVEIDDAEKVTKLKEAIQPKNRQTIKCDAKDLQLYLAKKGN AWLPDDPAAQDLNEGKVHTEIEALINGNMKEAWTIADVLVDNKMTGEGGLAPKSSQ IHVLVVVP |
| >jgil71308lgw1. 16.631.1 | MKLFCAVVGAGQSAFEVKIDNTESVSFAFKEAIAAELRYKGRPDMLELFLAKTESGWLS NDDDLITKLLQNRIDTSKLLKALWPTWKLNRANKPQLFGRDVS LGENVVHVLVRF |
| >jgil71310lgw1. 13.804.1 | MVKLFCAIVGVAGSAFSVEVDQDQTVDDLKDAIKTKNKIKLKKVDASALQLFLAKKG KAWLSNDKTGDTVLSQEDVASFEQMRGWSRLNDPKLFGTSVLLTEKVIHVLVVIP |
| >jgil72559lgw1. 33.407.1 | LVMVSLQCIVGKEECFSFNKIDDGANVSKLKDVIKEQSGGLITVPPPTMQLFLAKTGR EGWLTGDDATAVVSDDLNHFTLMDPTLFLVKDPKHFGEFGKPREGQVHVLVV |
| >jgil73233lgw1. 1.815.1 | MVKLFCAIIGEEGSTFPVDIDKQTVGGLKKKIKEENEDDPTLKTVA AKNLQLFLTKTES DRWLSSNDSVVIAMRKGDVPEQVKKL |
| >jgil73236lgw1. 1.816.1 | MVKLFCTIIGDGSVFFVTIDEQTVGDLKVAIAAALS YTGRPDLLQLFLAKTEKNAWLT YDKDLVKNL |
| >jgil73237lgw1. 12.699.1 | IMKLFCAIVGVAGSAFSVEVNEDQTVEDLKTAIADNQKFNANSTLQLFLAKKHEGAW LTQL |
| >jgil80099lgw1. 10.758.1 | MVKLSLQCAIVGQTGSSFDVEVGEGQTVGDLKNAIKTKTKLKTVA AKNLQLFLAKQPVG DESGKEVVPVYHPSVEEKKEESIKWLPDEHRAALKLVKGESDDYINALTAGEQILASKT IATWLYTKNNMELPSNEQIHVLVVVP |
| >jgil81603lgw1. 5.1157.1 | MVKLFCAIVGVAGSAFSVEVGEGQSVDDLKEAIKNLKRNDLKGVD RDKLQLFLAKK NGKGVWLTEKDVQNGVSDTSDLNLLGTMGAPLKFVGLLEKDVKFEPTLKDIESMNT VHVLVVVPPKRTSTAIVSEEK |
| >jgil81615lgw1. 2.1184.1 | MVKLFCMIVGEVGTAFSVDIADGDSVDDLKKAIAENQKFGFAASKLQLFLSKKNDTD EWLTEKDVQGGVIRDTSDLKPLVVARQILSTATDVEVKIETKAFEAEETSPVNLVVVPE RDSTA |
| >jgil83070lgw1. 12.944.1 | MMKLFCAIVGAQGTAFSVDIDVSQSVGDLKKAIKHKNE DIKCPHRDLQLFLAKNEKDE 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 | MKTATAFATVLALIVATNAAQVPPNPTLRGLRPTADTPSVEDDKEDRKDHHHVKKVKKIAIPVVPVPEVPQ FIPVVPVSPSTVVANSNNAVVGPPSTNVAGPGAAAPGAPGAVTPAPTTLNGRPAATPAATSTTRSRTPT APTNFAGARASGAQLPAAPVQAGAPGMTGFPNSAGVGDNTNNAFAGAGIGAAGGFPMAGNLAGGGFG GNPMNRFGGANGMGSGGIFGGAGTGPMANGFGGNGFQSMGFGMQDSNGLGGFGGQGMNTNFG QAGNNGFGGGQGGFGGGFNRRERL* |
| >scaffold_63_R3 81 | MVNAMTMAIVLCTIVLRCCSSRLRAALKQEVRLRADAAQAAQTASNISGLGTRLNALIARVSDLEKAFQAS QASHHP* |
| >scaffold_22_F1 619 | MGPLTGISLTFVACSRGSLGSSAFLLISLDSTAAASWSRVASRQRNKKGNACMRISKRLRLSLKTTKTMSY PTS* |
| >scaffold_52_R1 150 | MRLGLFLLVALAALFASCDASAARPVNDLPSGRFLRGESFKESLKILPKTYVKELVDDGDKLKAALQSWSDMK LSRKRLAKSLELSTNRRIKLRKIANRHKVNEVFILDQYVKHLKQLKRSRPLRVVE* |
| >scaffold_8_F31 04 | MSLIWRIKFVAAILLAGCNAFSDAPSHLTLTESALMLRLIDTDPDHIGSRDTGKRFLRRDNDDEALDFDTEEG KSSHPKMDKAVISPSIQGLSGQIPTALYNFLKI* |
| >scaffold_438_R 9 | MRLFPLFVAIATFLIATDAFLMTGDSNQISNVDSPPGSRQRLRAHAEVDFEKEAKEMMRMMKMKVTKED FAKKLIAEEIDDIINKRAPGMHEFMQTVKYKRYANYMNFNDMAKTSEYGLVKEIKAKSRAQVALKTFRK PTTSQSRWQSILASLKGKRIGK* |
| >scaffold_87_F3 00 | MKSIYRVLLLVTFALLCGLSNAVNSGGRLRVVDSTDERGGFPSYKDSFTKWRINSKIKSWVKKQKTDEYVLS KLGLSTLTGKDLVKAPKYSQFQDFKVGMMWLKEATPTTTFVNTLGLNKVEGAVEKADDFGTYYKVMALGEK ADDYPLVTWRKLFGGGSLEQLELKRKILLVCRDEIDISVMLG* |
| >scaffold_29_F3 65 | MRFAVVVLAFAVAVASTDSAVASEQAKLTSVVVAENSLPHTSLTAEQEQRLLRAASDGESEFMERTKFYY WYAMGRTPSYVYEDFFKGMDSIAENPNYKVVWERYKAYYEKRKAN* |
| >scaffold_10_F6 08 | MFCPLCVVYLPGLLLCSSDCIIPMYSSVQESAERLPLENRRRPPYRNRPTSITILEIPQIGRQPQVDRHLRLP* |
| >scaffold_52_R8 22 | MKILRSLTFLLIGTVAQAENSSKLLRGPDPNYDVAGLFPADTVVDTPTKEFEPEVDEPVIPTLLNELDE NSVERLVTEVTTYDDYTSEYNRHLRSVNDPAFQGGIQLAEPHHPKEHMIPSPHDDQVPLNRGWT* |
| >scaffold_27_F8 34 | MQIRVIALALSALGSVAAVQTTDRSLGTLVGCPAVRSKSPCLWAGENGVEVDSRALRELFLERHYVAHSD REAYGRNLQEHMTYIEGEDDKAFIGLDTVY* |
| >scaffold_623_F 5 | MHLGHLLFVTIVILLASSDGMSTQNAKKPIEDTVVGPSNSNGIKKTRALRTDNGSEERGFLSNLVKEAKVR WWLETGKSESQVKAQLLDELGKAFETNTKLFQKFKATKAYANKLNKWLQDDATTYSVWKILKLENVSSKQ LKSSPAYRTYVDYVNVQFDELLQRKWGAYKLPPEMVGSSKEMMAKSSIWGEAKRYEAYVKMALDMEGVTG KALKEHKNYAYNNFLEALKGN* |
| >scaffold_25_R1 072 | MCVAPPTICLIFQLCQGLAEILNDQVCRQNHPARQQLAISIGYMLDAARAVAYLHFSFPAFVHRDINPSNFL VDAECNVQLSGFGESRSTVKRNGRALRPKMKVPELEAKYFTLNATVASPLGTTTRQLDSPCTHMEKNSAEYI APELIDGLSEPFYGEAADVYALAITLWDILHPNGDKYPQANDDYVQVHVFEVCLQGLRPRINSNLPKLCSSII ERAWQQDPGLRPSAQIVKSLEDFQDDL CARLVLDLISDFHDCSSGSLRTTEDLHVRYTYGAFIVDRMIDRR FVRCPAEAIIRLGNALMDSVLHGHANHSTSFENNSASRYFDVDEAQLALPIDSHIERSVRSSSSLMNESSIA MLNVGHNDAPRSSFVGRTRSEPPCRCRQLGQRLINRSTHFHRVQVQLTALEPTVDGNTLTALLGGDE ASPQNSYNGLDMTPGIAATMA* |
| >scaffold_25_R3 36 | MYRVLLLVTFALLCGFSNAAKSGGRLRVVDSAEERGGFPSYKDSFTKWRINSKINSWVEKQKTDEYVLKKL GLSTLTGKELVKAAYRQFQDFKVGWVWLKEATPTTS |
| >scaffold_68_F6 89 | MRLNLVLAAVITLVSRACTAVSAASSTNQINLSNLNQLNMQNHAGDGNRFLRSKIVNDDIDSDDDIDR DEEERGGKTWAEKFAKWHARGESADDVYARFALEPVVRQAYKYGGQIGRL |
| >scaffold_24_F4 44 | MRLSFLLLTATFVLLSSGIVASPTTKDESIPSPNQVLSEGRRLRVHKSSIDDVEERGFNPEKFNRLMNERGYRST RFSNWWNKNYTDRYVYNLLRVDSPNYKRIFNYQTYLENFAPRLISS* |
| >scaffold_11_R2 | MAVWIVIAAATALVRVSGGGRILVNCSWIASICVLPRLRLADVDMAKMAARRRVFNGLAALYDCLTSS* |

| | |
|------------------------|--|
| 784 | |
| >scaffold_50_F8 48 | MRFFYFLLLSAALLSNSNATATVSGEGHVMTSADAPARALETNNGKRSRLRYATDEDEVETDKYDQKNGKY DNDDEEERNLQRHNSPNGLKRRRAGSRRIGRREYESDGGGR* |
| >scaffold_46_R1 328 | MRLSQVLVIAVASFVFASDTVAVATSNQAKISKMEQSPSQRLLRSNHYPVKEEEDSEDSVDFEERGFTTPDE EDLEERSPLSSATVKKLENIAGWGTTYSSVAMGTSSVSQTKAKALLALRDAYISIGIKSEKNAAKMAILMANK S* |
| >scaffold_24_F1 432 | MRLFAVVLILAACLAADVPALNDAPSKRLLRSTVRVDEEEERGAWDTLSNKFVKVILKPNQFAVRNMDDP KVAKAAKTLLKTFKSVDPQKFNVENFFRGKAFNNLENYVLRNLKQDINKQTSVAKVLSTGLGDEKAFHFL TATQSSDRAVKKSGKFFRDQLLTQWAVEGKTSAEVTKLFPKGLGANYYLHLENKYSGILMDLARDSQKRATR LAKLERARAAAT* |
| >scaffold_25_R1 150 | MCVAPPTICLIFQLGQGS�AEILNDQVCRQNHPARQQLAISIGYMLDAARAVAYLHSFSPSFVHRDINPSNFL VDAECNVQLSGFGESRSMVKRNGRALRPMKVPVLEAKQFALLGTVASPLGTTMTQMDSPCMHMEKNS AEYIAPELIDRQSEHFFYGAAADVVALAITLWDILHPNGDKYPQANDDYVQVHVYECVLQGLRPRINSNLPLK LCSIIERAWQQDPGLRPSAQQIVKSLEDFQDDL CARLVLDLISDFHDCSSRSLRTTEDPHVRYTYTGAFIVDRM IDRRFVRCPAEAIRLGNALMDSVVLHHAHSTSFENNSASRYYFDVDEAQLALPIDSHIERSVRSSSSLSMNES SIAMLNVGSDAHPPFSVGRMRSEPPCRCRQLGQRLINRKSTHFHRQVQFLTALEPTVDGNTLTALLGGE EIS PQSAYNDLDVTPGIAATMA* |
| >scaffold_6_R18 84 | MRVLSLVTLFAVSAQLIAANADQTGIAAVDSNTALLPRVLGVESKRTLRRYDPSEFDSEEAVDSDEEADPVE VADSDEEVVSEGEERVGIPGMEKVKASKATKADDMPVNAAGAMKWKILVQKNMGQLVETAKLVKLLKVG SFYSNVELEKMSLSALRQLDDIQQLQKADIKSNVFGTKATANGMRRKMTRTENMKLPPEQFLVSHVGRGA QRLGENGQRLLSAAVISKGGDINSKVLIISSSDTKKGFLLPKGGWDHGETIEKAVLREVIEEGGVNGQLLHKL GEYFPKKGATAYAYMMKASTVYDDWAESIRYIWRVTEMLWKY* |
| >scaffold_10_F2 067 | MRLSVILLVVAAFVAALDPATAANDANTVVAVPNVHESIATGRFLRAHLEDDYPVKDERDEDDNEKDEER MFSFFQEKATALSFAKNLVSQSGDDLVEAVSNLTKKEFVALFNQGHTHMAKMVPGFHPGMSLDEFGAVVR AAGLSDDMKNALMVGYGKYLALHMD* |
| >scaffold_58_R5 67 | MVFKCWRLTFVVLVTAPEILKGEYGTAAADWWAFGAVLYELLTGLPPWYSENAREMCKRVLRTPLSVPEY VSTEAKNLLQKLLTRNPYERLGSLLGGPEIKEHPFFQHIDWEMLSRDAPAPIQPCATSOTVVRPVIIPYVDF VSNFLPCL* |
| >scaffold_13_F2 050 | MVSVSSVYLSAVLAFATLQGTATSLSLDPYTTCTSYENFPGRDTDIKDGGSCTVIVPEDPSTSTSKRKLWV DGNDIADLEAHFGVPMERKLRLNPTSTHSSPPWSSGNWLVNVD SINYVWDQGGQPSAAEKYATAFGLDV KTFMDDVSAQNGIDSVQNATECTEDKECYEDLICAKRAGKSSGRCLPTWWGMGNAAAAATLEKEPKCP VTFNGVTFQPMDIKALVTDIYDSANVSVFTGSRYNGYEDSIDDYGRHTDASYRDLNPGFLHIAATNMLGLL NTTFIVDKDPDYVWNQPVVGFVREQMNMTPAEAAKLYGLDYPWNVNASSIVVNSYLSWVTESLN GPIVTLNNTTETGNITYLLELNDDEEIIIGGEWLYDSNDYHPDFLWLEAKPAPDTATSFGLSYTNVTMLLEKATE C* |
| >scaffold_12_F1 755 | MQTIRAIVTAALTAVAGDEGATRTRLRGAMADNSELDQLEGSGSNNSILWVNASKPEERRPIRPNYHHGH QQLVCQINFPKNCWYVDIFDQ* |
| >scaffold_81_R7 5 | MRSALVLLSTALLACIAGVSTTNPFKVTVTETFRLRKSDPKTNEERVIGGSLTSKVGDFIGSTKLQKYKLRWKT LQLDDDLKSVLKSPQVEKIAKYNLNKASGSQVSMFRRIAAYSDDILARTLSIERSADDNPALLAMVKQLRE DQIANWLKKNKTVPRVSKLKLGSDESIFRSKALDVLDFIKKYNTARNGDESLLKLTITTYGGESELVTMISRA YPYAQLNPQSVEKANNIENQLISKWRSENRPDFSVMSKVKFGDDINEALSSGKVRVFFKYSASKTMALKRLN AKYGEGEVAVAMARAKSTLSGHDAVVVQKQMTGWLSNGYSMERVFSILRFKEADDFYKLDALAEYVK LLKSKNPQDTTNILQVLRKGFNGEEDKLAIALSRPLSEEAKGYQALLFKDWMARDMDPMSVAVSVFKIPEAD VAAGFSKEITPIMKQYTKFYNGAADIQPLPPAVRNGRS* |
| >scaffold_81_F1 03 | MLLFRLLLLVAIGTFLVSDSVLAASDDSTFARRFLRTDDSEKINLSGLEQSASLKVGLKIDNVWDIKLQAKS DTAFKLLNLDQAGEKAFKSPQFKTWSYMSISSEKYPEAIMSTLAARYSDETLVKMIEATKKFEGMEGIAMK MQEAQKKNWMSQSGKTADDLFEVLKLDKRSMENLLTNPKLDIWSGYLNLFNKYSKPGKETTAVNTFVTFYGD EVVAKALEAAKKVPNTKEKATELQALFTQWLMEGAKPHQIWKMLQMEKATWMRNPDANIWREYLAFY KLHK* |
| >scaffold_47_R6 72 | MRLAFALLVLGAVLLSISVSTASTQADKEDDVVPSGEDTRFLRGLTDDSEERWGIRDLFGRKKFEQMLLNN WDELNKFVDTVISKLGKTYGAELLYLNSRPKAVRAAAK* |
| >scaffold_17_F1 | MGVTVLVLSFSVSGGKSSMCPHSDCIFDLHELMYNLQISAISFGGMQQNGSMLTRAVSQRNSNPTFM |

| | |
|------------------------|---|
| 914 | CVPSLSRMLRILSGKSYREK* |
| >scaffold_31_F1 334 | MRLLLWALLVTLVAFVSSINAEFTADSKVTQDSKEEINALTRLLAVDSSDAAKRFLRGDAKDLTTANDDSKELS AEGEERGLIPSSITNLVSKVKTGWANWKAKALEKAFQHMVKNGENPTTLAKRLDIGRTVEGRHHRLYEKFTA WWINYHTVAGT* |
| >scaffold_143_R 10 | MRASIVLVAVIAVFRSGLASASNTAQLQQTVPGATNKVENAEGGRFLREGLKDESDDLSEERGIWGDLPK KAKPYLNGDIMYARLFKKSNSQLWRREITPDELKSMVIHLEKAGWSGEKQLQKDKSHGYEKYFYTPIKD* |
| >scaffold_40_R1 065 | MAFSSFLKFLAMATLLMVQINAEPSSQQRNLRIQSDDLPEEQARRLGNWVKALIINVKNNPGVSDGVVNL KKADFVTGTSKKIKEVKDTAGKVKDKVTGK* |
| >scaffold_51_F6 03 | MQLTNILWMAIVILFSCNDAASRPELPTISGVDVAVQLNPVNQRFLRSDYNSEERAVTLPFAKLDVAVAKL TTKLFNTEKLAALGKTKAEYNAREALFKEVNAVNPVAVRKAFLAKLAKSGDYNWLLPYWLGFKLKGIRKVVET EIKNAVRAGNAVS* |
| >scaffold_2_F40 6 | MVVVGVGVCVASSATYAVPTDPDCAIFSSPQGVWNYKTGILDIPYGRVWNGRCWAVRPIRKRPAKLRKR RSRNQQRHSRPRMLRLNLWLEPLVKTAKARAMTQMKFNLLTERRRSRKRDPVPRQRPQYAPGTARSVEEK AALRPVGVQVICYACSKPGHFAKECPDAEANARNDAYLASRIQRSAGKYNERSL* |
| >scaffold_107_F 190 | MRVLSLVTLFTFVSAQLVAVNADRTGIAAGDLNTALLQGVLAVERSLRRLRYPSEFDERESEEEADSDEEVD SEDEERQIGLPGIEKLDLDAVSKFTKTDVVSQAADDDVVEKAAKASPKWKALMLKTWMSLLRPEDW* |
| >scaffold_1_R39 4 | MKTATAFATVLALIVATNGAHVSPNTPALRGLRPTADTPSVEDDKEDRKHDDHHVKKVKKIAIPVVPVEVPQ YIPVPSVPSVTVVASSNNAVVGPNSTNVAGPGAAAPGAPGAVTPAPTTLNNGRPAATPAATSTTRSPTP APTNFAGARPSGAQLPAAPAQAGAPGMAGFPNSVGIGDNTNNAFGAGIGAGGFPMTGNGLAGGFG GNPTNRFGGANGMGSGGMFGGAGPAMGGRLNGQSMGFGMQGGNGLGGFGGQDMMANFGQGTN NGFAGGQALTQGGFGGNSFGGGFSRQRRR* |
| >scaffold_17_R1 815 | MKACKFLVLLPSQALANHAVLHVVDRLFRYTAIQRELRKLVGDYVSMRTEMNQCCQKWLALRHEHTT TLDRLEKIETRLDSLVEAATISMNKLQ* |
| >scaffold_11_F1 315 | MRIGYVTLTATTAILASFGNVSGDSDLTHTQVAKVASLDAVLPINTNRFRRRENYDDEEAATEERSAFEALA AKLDKSVLPAPFLKIANVKDSERAFHILQQYPISLEKRLDILALHGLSKSDRKKVLLLIQ* |
| >scaffold_14_F2 398 | MQPTLVIFVNVLLLLVGLIGRSDAVHFDATAQASPSDSTIESKTATNERFLRGNDKTIRTTEEDRDITVGLSVI ASAKAVNMRNKINIMLARGLSPTRVLLKLVVMSDKNFNNFARFYARYLDKYSRKKPNLPKTAEDVIVLPK MKDWLAQKLLPFQVEQNLKDLASRNVNRYMQLYFKDADNIIILPRLERWENQKLLPSHFKNLIEIGVKDTT KYM EWYMRNGGEQVVKAKLQKWLROGIDPREIGAKLRKIGVTD TAKYVAVYIDAPVVTKLKTWVNTGLR PPQILDELIRAGVTDIQRYYNTISVIYQQRQVRLYRKNLP* |
| >scaffold_53_F9 3 | MRFTYFVAVALTGLLASNDLVVASTADKTVVHDQVLSRELIDTGLNDNEKRSRLRTNEDLEDDSEDLLKSDDEE RFSLIQLSNQPRYYVWFEDEMTPKDVRRKFLTRHSIKLVKRSIYRGYVYKYDEHCSYFENRKKDFCRAQEY* |
| >scaffold_18_R4 67 | MRLNLVVLATVITLVSRCATAASAASSTNQTNLKLDQALGYQVGTMQNHAGDGNRFLRRAKTVDGSDDDI DSDDEERGGKTWDEKFAKWHARGETADDIYGRFFLEPIVRRAYKTANIGSLKQHEYYQKWLKYAFLKAKE K* |
| >scaffold_25_F8 10 | MRITYILAVTVAATLHSSVTAIPSVKSSKATENGAVPAVIDSTHAGAGRMLRWVKEYEEDLDDSDDLDDDL DDDLDDDLDDDLDDLEERGFDTLKKANPLKLVKGTCLTAEQAQKVEALKDAADYQKMIENANKLIRS D* |
| >scaffold_6_F39 33 | MRFCFVVLATV FALLSSETPAIATAEKHSEALSFINGNNGGTANRSLRSHDKVNAVEEERAIDADKLVKLLNP DDIAAALKNYDDEVLFTRWWQHEDDVVNKLMWPNRMKNMPILTKFNDFRSTSMHYDQVLTWPWLSTK DLDEVATALRASKMNLAKEQFNVWHKSGVKPAEISAAIEKVNPLKRRKGYGALNGLYKMYVSGEAKKTARA AAKKT EFAAKRAQAKKIAEAAA KRAQEIKTQKETAKKIADAAAKRAAELKAKAAVAA* |
| >scaffold_193_R 7 | MSKATPNLLGISWLLGSLVGSYHLLNRPCLCAWNGRLNWPPRLGVGLRFHQQRRTLIYLEMHRRYLLLLQL TPCFLPPTLHWRHLRRQRSTKT* |
| >scaffold_686_R 5 | MRFHALLAAFLAFGSGND AFELTADGAAAYSRLRSEPKTNEERVFEGLTSKVDLIDTTKLQYKQLQWS KLQLGDDLTA VLKSPDVAKIAKFNLRASGSQVSMFQRITAKYSDDLARTLVSIERSADDNPALLAMVKQLR DDQIANWLKNRETVPGVVSKLKGTDSEIFRSKALDVLEDFIKKYNTARNGDESLKLTLLTIYGGSEELVTMVS RARATYPYAQMNPQSVDKASNIENQ |
| >scaffold_5_F19 44 | MRLSYALPATIIAVTFLSSGNAVATTDGGTMDLSAMTSPNAVASIDA AVGGGKRSLRYHNKDPEDSDDEAL LEEEERKYTNMFSTTKLDEMLEDGTMMSRFKWKKEKGYNTYNLPAETQGDKYTWIRQKYRDYLYHN* |
| >scaffold_2_F15 74 | MGLHLLLAATTFVACTNAADQLEMPVKRGLRANEERAFGAGVTEGLSNWVARAAPKLLLDNELEHLA MKVTSTDKVFKMLKLDGLDILRNPNLKAFASYIRKVHATNPQVLLITLITRYGDDTLAKFLFEAKQVRRTE |

| | |
|------------------------|---|
| | ESAKMLQAAQFIKWFDGKTPNQVFNLLGLKHLTAYEDKFHKLWWEYVVAHYHLASKLKKPLPVEL* |
| >scaffold_24_R7 96 | MRFTYFLLVATAALLASCNAAAATSNNQNKLSTMTSTDAVVSALESSNDKRFLRSYRKEDDGDSDDEEERS VMTAEQVAKWTKKVERWVKKGHTPSYIKDKLTALDGTMNAKNREKYRMFGAAWGRANPHELGRM* |
| >scaffold_10_R1 115 | MRLSVILLVVAASVAAGVPTAAANEANTVVAAPKLPTARFLRAHLEDDYPVKDERGEDSDDENDEERMFS IFGEKATALKAFKKLVSSEDDDLVMAVSALSKEEFVALFNQGOAAMAKMVPGFPRGMSLDDFATAVSSAGL KPDMQNALMVGYGKYLHQMG* |
| >scaffold_3_F29 10 | MATLVVCASNFALASTVADDLHQEAEGNVDRRLRAAAPANKDNVAKIAGGFLTKEGKSTLTKEQMIKNA NGDEAAVKKAILLASTAKESAKMSDESIAKLSAMITTAUVKDPKSWPRLQKFKVKTLAGVGGALYLGAYKLL FNKNSSSGPVTTTTGSADLVAASTSGSGSA* |
| >scaffold_26_R1 068 | MLLCVILAFNLTVILVLCPRIGRVFVSFSTVAAVWFDAGDTVLSPLVAPSSWFVIRLAAAFSGEWILGEDS RDLRPDDMDSRSLVTILTEELAEVGGVEAEGRFCSWLVGFLPSAIVDCEDRMIFELLPWGRFDTSVSLATC R* |
| >scaffold_8_R11 92 | MWLLLCVGFVAFVSCVDAVRLGSRVLLCPPSSPPYRTPYCPQQFSHPRPRYEDTPQRGLRAFGAPALDGFVLR CRIVPRSSRLTLQPVLLTV* |
| >scaffold_93_F9 1 | MRVLNLTMLGAIAFIVNMSAVSSADKPQLAVAGALPISRFLRTHYTDDEERAIGNLLPGSKKISSFMTDKKLSK PQEQRNRNRRRFQQTARQGRQVREPEIPRLGSIIRGRFQPETPDPELDASHACATVWR* |
| >scaffold_24_F5 45 | MQSAAYIAAFAAMLAAVKAESGSILQGRQVLRGQLRVHQCADHALGHRGRQAVPAVHEGGRRVLRHLPG SVPVLLHVEDGLPVH* |
| >scaffold_2_R11 64 | MHFHSFIFLHHTSMTTSAASCSEAAARSRTIELLAPVYEDVCCAIKGGSTTVPVTLERTADDWIQLVFPQT NVRKDVACVNGASGCRDFVRLLRALLTLTEQLVVKVAATYVLAHRSADVATLEMRVQVHEEARRGTSLRD FDLVLCHLQATVLSVVSRSKGKKIEARHEALESARSSSCHVVGCRLLHPWDASSGQFNPSIN* |
| >scaffold_214_R 4 | MRLAIKTLVALAAVLLATSTEAAKAVQTGGDVNVVQSSHILPGENKRLRSEHDEGKLEDEEEDDEEERKY GANLFSTAKMEKMLGNDWYRYQVARRWKRDGYTWETLPKDVVPDLVRYFKGFRERHG* |
| >scaffold_7_R12 23 | MVKVVRVAVVGAVFMASVDATSMKCGTTQATPLGAPMND CNPAYGGEIDPNCMPEVAITDAEGCEHAT GPAMDEIIGSITGASAATRSLRRMEDASNSDVADLETYFGESLELSFTTLKEQYSSASVPTTPWPGSYWPTYQ DGINVIWKTGDVSASEKYAIAFGDPTDFMKNISAKTGIDSRSSSTKCTADTDCYRNDGVSACKRKGVESGY CVPTWYGICHAWAPAALLEAEPQCDVVKNNQTFHVLDIKALMTDVYDGSISTVFTGARFNGPDIPEEMDA YGRYISAPRRDLGPGFFHIAITNILGKHKPFILDVTAGSQVWNQPVRSYQVQTMELVDANEASQKYFGVSP YPFNSEMVFLAYVKTTSWIVEAYADGPLVSSGQVDAYTVSNDYQYVLELDANYAVIGGEWVEGSKTDHPD FLWFPTKPDASTITSTGLSYAHVKELLELSLACGSSGGTSENASASTSTGGSTASISGSASSASKSTNESTSS SNTYSNSASASSSDSSASAGSSASTSTSAASASTGTSETASASTSSASGSQTTSSSGYTSSASTSSGSTSS NYVAPGSTGSGSTKTEAPSTAYPSSGTNEDGFLSMGSDGSPARSASGEESAAGQGSTPASGSAGTTDATST YTSLTGSTVEDTGASLPSGKDALILTPSTSPSTTGEESTPATTAAPIGVGYTTPPTSDGQDLADVLDTTSPID SRAAVTDEPSEGESTGTTALAHRS* |
| >scaffold_568_R 2 | MRLVIKTLVALAAVVLATSTEAAKAVQTGGDVNAVQSSHILTGENKRLRSEDEGNLLEDEENLLEDEE RKGGANLFSSAKMEKMLGNDWYRYQVARRWKQYGYTWETLPKDVVPDLVRYFKGFRQRHG* |
| >scaffold_81_R2 17 | MTVRDIVFLAAVLLVQAQALSVPTDSSTRILRSTVDTEDLDFEQRAGLAESVIKKATAALRPNVTPPELANW LNKGKSVDDVFARLQLNKASDKILDNPQFATWLTYTDDFVKKNPNTETSAISTLTAHFGTEALTKMIQAACK VDGTEAIAATKLEAAQIQKWLSSGKSADDIFILKLDQVDDSLASPFDANNYRKVFAQDNPTNLHKPSVR VLEFVPEPLQQGPTS* |
| >scaffold_81_F2 24 | MHLQIALFLVVVSLVNVEAVPAKAQSDSVVSRFHIGRLLRDEREERGVSANAVETITDAVESKINTAQLKSW LESGESADDVFKLLKLSAADKVLGHAKLDEWIEYMKLFNGQKSKTTLIKTLTAHFEDDGVARMIQKALQ VDSTAKMAKRLQFEQIQRWLGHEKTPPEVLTLLKLDINRYDLFEKPELLTWVKYLLDDWNKMYPDRQTTLFA RISPLLEEGILANLLIKAKSVASTEKIALRIQAEQTASWLKAEKTPDDLFTLLRLNRAEDSPLENPIFDAWVKYA DDFREMYPKVSFDPIATISEHYTAAQVATMIVEASKSPSTSSIAHRLNTEQFRDWLNTRQSPVRVFKLLKLE AGDKLFQSPVITWLNATFYNTKREKVSITLLRKRFGDEVLAGILTEAQQVPATKEATKLLTSLVGRWPKS RVHPDNVYKWLVRVQGREKTDGFRFLFYERYAAAYKAARNG* |
| >scaffold_37_R1 044 | MRVASLVLLAAVTLASREVVCATTNTLAKADTDTINAIPIDARPLRNLKKTSSVGYDAEERAAEELVDV ELLDKVISDHAYAKQVFLSWLQNGQTSQDIENRLETGLVQDKYKVVVTQYAHYLTLEERSA* |
| >scaffold_18_R1 184 | MRFSFIVLLAAISSLVTTSHALPALPHEQSTVSKVSSNPDQSDTGAQPSERRFLRSAGVSGKMLKSESFKDAK FAEWLQKQLSSFTVFESKLGLEKRYGLYDEYAKLHKASGQYP* |
| >scaffold_175_F | MWRLSCWAMAETLSVPSLVCSFAVPAVLELPSEVPAAVIVRGRSQVCLRGHRRHRHRRHLRWNCCSSLR |

| | |
|------------------------|---|
| 27 | RLRHPFRLRRASLVVLVESSLPS* |
| >scaffold_13_F2 21 | MTALSAWCTAVNSVDAGKCCLLVLMLYCAAQRLVNPSQLLPTGVMGVCRLHRWNHWISRRDRTLQASDT LQDAAYATGVSACEVTKLCELLSSGRSRGDAGSRLCPV* |
| >scaffold_81_R2 28 | MRVHSLVLLAIAIISATDAVAAQSLPEEQNVSRFLRTNKAKETDSEERVNLAGFELKIGFVDDIIEKMHLSPTEFK ELLANKGDAKKAFKAFHVDDVADDVFQSAQWKEWAEYVYKAVKKNENADEALAAAMSVAYNPDGSLK LAKAAQNPNTKDIATVLENARQSSWINGGFTPGAIFKTLNLDKADDIFDTPAFATWTNFKAYNEKHPKKA MTEFEVFSVYQKQKFAKLLASADDGKYAQAMKKELVQSWIDDGVHPTNMFALIKLDKVDLLTNPVNL WVRYMKEFNELYPKQATTMIKFTQSYGDEKVALMIQEAAKSSDEKVVFAKNLQTAQINQWMVDKKT EMLTVLGINSQTLTENPLGNVWRAYNKEYTKKMANGDFAFQP* |
| >scaffold_50_F1 228 | MRLTCILLMTAAALVGLDASAATTGNSVVANAAMVISPLAPESQGRSLRLVYDDEDDSADEKDEEEESA DEVDEERGWFSDKVALTSLASKFVGKSTDEMGEVIKGLTPAQINTMFESGEDSIQQLPGFKTGMDFTKFDE LVQKLPQEQQALVMSAYTKYLHNNKIL* |
| >scaffold_38_R2 61 | MRHNFAKVLVAIAFVVTCSAEAEYHTTKSDPLALTLSLRGGVPRTRSLRSDKYAGNEERAAGTSAVESLVKAT TNVDDVETWLRGDTTNKVFALKLQKAGDNLDPQLPTLMRYLRFNEANPTKKTSLLATLTHYGNHG LTKIIEAGLASTSKKTVATAKHLQTEQIQYWMARGRSPESVFGVLKLDKLLPFTWISSDLFEKPLKTWIRYLD EFNAENPKQKTTLISILSKRYNDKTLANMLIAANKAESTSSIAKRIQSEQLTQWLSKGGKPEDIFTLQLEKAGD SLFQNPFPWIKYADDLRYSETDLATMSTLINHYSDEAVAKMIMAASEDPNTKMLAKRLQSELQKDWV FSGQTPNDVFIMMNLDKLTKLENPLFDIYQYGLFVNKMSAKANWNPITALLRITYGSEKELSGILMAALK QPSTKAMASKLLQIT* |
| >scaffold_13_R7 93 | MRLSQVLVIAAASLLFASDTIAVATSNQAKISKMAQSSPNQRLLSNKYSVKEEDESVDVFGKRGFATNDE EDLEERSPLSNAIVAKLDDIAAGWGTTYKIAMGQSSVSEAKINALLAMRDAFISNNKKEKELARGMLMLAN KS* |
| >scaffold_29_F7 71 | MRLALLFLLVIATLVATGNTLAVEHTEGKITSSDTLLQPTAISNPDTTQTRRLRTPESPNDGEERRLVNVDLAIK DAVHEVRKLTWKLQFAVWVWGLKKSPTKLIQEWGMKYPYTS DPRWAKILAYKAFWVGKGLPKYP* |
| >scaffold_25_F1 705 | MQRMMSTPFFLMITTIGDAQGEVEVVMIPADNSPLIKSDNFSRNCGATRRTWATGVALPTSMSQTAPRA AGGRAFPPIRRLRFQMICLNRSICSGVLPSASSIWISSILTSSVVALTSGRMAI* |
| >scaffold_102_R 91 | MQQHLVFCVKVANFILSTVLFRRVALPATHADLNFQSQHEQQLRPLTSSVLDQTRA AVARGRHSQRLRRRRLG WR* |
| >scaffold_83_R2 91 | MSFTIGVCLVLLQSEVHIICRTHKKFSKNSFVNFLVIAPKEGYGQPYGLHHTQVLRRLGRVLLRRGRERHF YILDSN* |
| >scaffold_3_F32 92 | MRLTCILLVAVATLVGLDASAATTGNTVVANAAMVNSPLVPESQGRSLRLVYDDEDDSADEKDEEEES ADEVDEERGWFDTKMALSSLASKFVGKSADEMGEVIKGLSPQQINTIFDSGESSIQKLLPGFKTGMEFDKFD DLVRKLPQEQQGVVMSAYSKYLHNNGRFS* |
| >scaffold_81_F7 0 | MRLHCLLLAVALASVDASPLQLNQSSISKANGFLSKRMLRTAESSEDEEERMNMPGLDVTNNAISSTV TKGKMKLWLAKGEDADDALKLFNLDNGIEKLRDATNFNTYASYVRQYNAQNPDPHEVSMIAAIAARYGDDE AAKLLHTMRHASTQAVATKLSQEQVDLWMAAKTPKEVFALLKLNQNVDDLSSPALYSMDDFLNAYNR QHKNQHTSLMKVVLGSYDNAKAKALQDGLSASTKSRAWELKERQFDDWIKAGIKPEN* |
| >scaffold_46_F8 62 | MVMILIFIILETGEGFVPDGDNRHYADTNSAQTLLKQTDIGEMREQGASGKEIIQKLVENSSTWETKTEFSKQ KYLKQKQKQYMPRVRFRLCTAESLCRTYRLKNPAKICNLREDSLGQMLVYGNIFAGGQVLVVDTCMGLVTG AIAERQGGSGRIICPYEQQPAADILRRFNFGTISSFCEQNGWIVANFLFKNR* |
| >scaffold_6_R31 68 | MRLAIKTLVALAAAILATSTEA AKAVQTGGDVHVQSSHILPGENKRLRSEHDEGKLEDEEEDDEEERKY GANLFSKAKMEKMLGNDWYRYQVARRWKREGYTWEGLPSNVPAALIRYFKGFRERHG* |
| >scaffold_93_R2 83 | MRSSFLVLLGATLLQSINSTSLAETHITSNVQLHSEDIVRRLRFAPEQVEERGLTVPKFQDIVNNQQLAWWLK RGKTTDDVFALKLNLGSSIFENPKFVAVIKYVDDFNSKHNGISAIPTLSKQYGGDDVLAKMLQKARQDET TKATATRLQTEQMGIWRSQGISDVTVFATFKLDEGIANLLANPGFNWARYLTFEFPNGKKTIFKTLDEHFSQ NILSQLLIAAQKNPRTEKFATSLQNVQLQGFLERGESPTVVFKLLQLDKGADNLFANPQYKTLWLNATSFQKT KSDAAPAPVIDTLTAHYTDISLVKMIATAKATEGTKNMATYVEKSLIGKWATDGKPTYVSKLLWASATDKKT LEAVYLEQLIIHSTVKVSSNLQLEQISLWVGKKEPTDAVFKLLRLNKGSENVFKRPQFDTWLQFANKFKTQNP EQSKSVFATLSAHYDDLPLATMIKTAKEDVNTKKIAEYVERGLLPKWAADGKAPAYVLNKLATDKQDKERVL TLFMKEIRIVEDKITRAES* |
| >scaffold_118_R 15 | MQTRYRLLLLVLLAVFLASTRASGASLQETTATSENPRPDGPVPTRSLSRESGAVRLLSPSEERMVKEAIG WLWHVVKMKLKIWFVLYRGTTEQVLEKLVVSKTDENYKYYARYFRYYVYKPNKMPKNAPTKTADAIMK |

| | |
|------------------------|---|
| | ARLQDWLDKNLSPPQVFKELGFSGTFASAQGHDPDYKYFEQYAKMWSDLQVRISSGHA* |
| >scaffold_31_R7 46 | MAIFVACSTVATAESVALTLGDNEVRRRLRNQNIKAAGDLISKSKEATLTKVINIAKTANGDEAAARRAV MLAAGAK* |
| >scaffold_8_F29 12 | MIAPLFLIHCILLTVCRGRHPPFGLVVVVLQLEQRRRDVRAQVRNLRIHIVTAATNNNQRYAVLDNIISIN ELTA* |
| >scaffold_18_R6 89 | MSSSTGSQVPSSFLLGSLAGVPPFALIAGTHARPLRAVTVAASTNTPVVSVMNSPRSPFAATAISMVTA FAGICALPAIRAGAVVRPASASLVRNGITIVRPFSPYFTSLRSKSIFAFLSVRKSTPSRNSSVQPEMTHASTQIVS PNTFWNWKRQTPRTSSRNPPIPSTYSTGASRGLFPYCIHAVPATEYLAQLSSLMNFCSPSIITHAVGNPSSAS TYCTASPPGSPCRCTSLTRANSVSVPLSPHFRRSCRCRNCSSASVACFSLTASRVSSKVRFTSLTRSSLERA RRTRLRFLILFLRRLRRRACCNCFSGPSSLPVSGLSLSTNSASSPILSGSAASPSEIACCTSSFAGVVRLLPPS APTPPVASCCAPSPDRASSRTRCSSLAGRCR* |
| >scaffold_3_F37 06 | MPRATSPSLVLLSLYNSIGLGELEPNRSWISVSVGIHAAQKLWWISARRDERVILARKLRPGSVRGLFAGAT RLLGWVVVRVFGRRHGGGGFFLELPIFSRIVERILFRVGVSEVLLRGMILLVESMVLGDREVTLCRETGSSSA NGHCRGCACCRGSRRLRVRWRERGLERWRVHSEMDSVLPET* |
| >scaffold_2_F13 82 | MRFFWVLLTVIIVLAGVDTCTARGQSKLSSDFVRPFVKNAANTQLESKATRLLRDVTKYDNDEERAILPS GLSNMISNVKKWFSNFISKIKLSFGEKRLNAWLKQKGTPEVFEILNLHKGTKGLLANKNLRTWSVFMTRY NRKNPTKMVNMLGLTLTKYYGNEAVVLEVSQRGKTRPLANRLQTQQLHGWARNGLNTDVFNLLRVGE GSVKTLAQNRALNVVYFRQMNMFYKNGWEAELTKLLTVYDDIPMAKAFAVAKTDPGTYISSQLQRFQ FKKWLADKVEPATLLEKLMKDKTKLSFEPTVEVYVAYSSFYKAYSKTVR* |
| >scaffold_60_F7 08 | MSWMICWICLVTKTISLMQCIRDQCRLHELERTRNLRQLSFKLVALKSSTDFFRHYLRCSHDPLLFGAPK Y* |
| >scaffold_24_F1 319 | MRLSFIFLLAVTFSSLVATSHALPHEQSTVLKVAATNPDQSVAEQTSQKRFLRSAGLGSQMLKSESFKKSKF DELLDDNVSSFTFFVDMLKSKKKYRDLYDEYAAIRKARGAYP* |
| >scaffold_18_F2 148 | MRPNLVVLAALVSRCTAVSAASTNQINLSESNQALDTLQNHAGDGNRFLRSANIVDDSVDDIDSDDDI DNDDEERGGKTWAQKFAKWHARGETADDVYKRFKLEPFVRYAYKHRQIGMLKENEYRKYWSITWPS* |
| >scaffold_5_R12 81 | MQFTYVLLLALCIAITASAQVQDNEYQAVVSPFAKRSLSVDTGTESDDEEERGIIPSSLKDIVKKGTSKVS SLLRKYKSLQKAGRTDDDIYKLVWRQRKNTDQIYTRWIRLGKSEEVSQLFLKHGLNAEVLYNLSRQKGSMD DIYSLWKKLLEPGQIYNIWLKSKTDDQIFSAWYKAKMPEDIGHVLRGKDPNTRKRVPFWEKYSEFYRAK KYRTS* |
| >scaffold_81_R2 13 | MHVRLFLPVLVLLVGSDDTSADKSVPTTRLLRSQKLLQITTKSECESTSDPSTTSSKTGSLNRGYP RSDPTK HDTNESNILFYLMK* |
| >scaffold_7_R22 62 | MRLHRTLLAPIAVFLAWNCFDPTTNAQVVAQDLSDKAFANTNALRSLRSQKSTSEDDPIINEETRNTMYT QLFPAWYAAGKTPEDAFEELKNPSTGDENWPIYKNYKMYDYTYKTMG* |
| >scaffold_39_R4 58 | MRTPSFLVLAALLVNADAASSRHLANPNVQNLAHVNDNWLDTKRFLRVHVEGEERGISAPSVKMLQGW VERGLISDEAVKLISLGDKADDLLSASLLNAWFSYVVKFNKENPSDKMNMIKTLTARFGDEALSTMIETAKRS SKTSAMAELQAKQLANWAALKKNPDDIFALLHLNSAKSLLFDQPSVNSWLKYMDDFTLKSSSVEFSAISTLR KYYNDETAEEMIIVALKSSKTSEAAKRVETELLRTWLNSMKSSTDVRRLLNLSTAPQSELSIWRNYVALLNKV DPKFKTEMLQAVVKKGLITDDTFRLLTLGNAADDLLNGSLLSAWATYIKVFNQNPTEQLNLIIVLTARFGD EAVSTMLEAAKVPYHTIANRVQTEQMKLWLDAGKLPDHIFVALKLNVTVKLFDQPLNTWVVYLDEFN KANPNSKTTLFSTLQTRYSEATLAKMLLVAKQPNLES LAVRIQGEQLQFWIKANRRRPGDIFKMLRLNILGSD LMRNQLFTAWVQYTDDEFKLNPGTKLTTLATLRKYYSDETLVTLFLKASQSPNTAKMGKRMESEMLREWFS AGTKPTHVPVGFALLNLGRTGTVFESPLYNVWNTNYVGFMMKAGAFRGDTITLLRGIYGDES LAKVLIAGK VQSTKNVANTLEKELFALWKAARVHPTQIHKLLRVENVSRSNPIYKFGDYVLAYTRA* |
| >scaffold_129_R 17 | MRLSFVLLAAAVSSLVATGHALPHEQPTVSKVTATNPDQSVTEGQTSEKRFLRSAGIGRQMLKSDSFKKTK FDEWLADNLSFTVFIDKLGKKEKYSLSYNEAALHKASGQYP* |
| >scaffold_73_R2 12 | MRLLYLAGVAILSFAADATGAKVLVSDDSNHNGAQAASAVVSTTRLLRARSVIDEERAGGISASADKLAKL FKSSKVTAVAQQGENCGKCLLSHESREHSLEYGI* |
| >scaffold_46_F1 215 | MRLSQALVIAAATLLASDTVAVATSNAQAKISKMVQSTPSQRLLRGNKYPIYEEDESEDSVDLEERGFATPD EEEEERSPLSISQVKKLEGYARRWHTTEKVALGRSSISEDKVQALLALRNAYISGVKSEKNAAKMLILRAN S* |
| >scaffold_15_R1 282 | MRLSQVLVITVASLLFASDTVAVATSNAQANLAISKMAPSQRLLRSNKYPIKEEEDSEDSIDVQERDFTAH DEEDLEERSPLSDAVVKKLNTIANGWGTSYSSVAMGRSSISQAKASALLALRDAYVSGDRGAKAAAKMAILR |

| | |
|------------------------|--|
| | ANKK* |
| >scaffold_77_F1 29 | MRLSYIFAVVVLVAILQSSGTAHPVYEDSKQIVSAGAGDATKVDSGRLLRGVEEKMEAEDEERLFKNLGTYLKKI PQKIKDSWEVKKAKEQLERSRNRKWIREQNAVPTS* |
| >scaffold_244_R 1 | MRLTFVLLAVAASVLRTRTDATPATYGEVSTSTIESPNRSALKNMAETLKERRLRRPRCPSTIYRKKSAQTFSETR CSGPACPQSLSRGPSR* |
| >scaffold_5_R34 48 | MMRWRLFILHALMATLTLPSSCWTVQILTRGPRTRKLRCLSTLPRRDNTEVFDCLYEGNATLQRRRIATVTW RRKKLVMLKL* |
| >scaffold_107_F 161 | MRVLSLVTLFTFVSAQLAAVNADRTGIAAADLNTALLQRVLVGVEKRTLRYPDPERDSEGDADSDDEDEERI ITIPGIEKLDDAVSKVTKTDDMVSKAAKASTKWKALIQKNINELAETGKLVKTLKANVLYKDVELEKMSLQGLR QLDDIEQLRKVDIENKVKGTTPDGMRRKMEHTANMKLPPAQFLESHIGRDAQLVGESGQRLLSAAVVSK GDDVHGKVLIISSNPKKGFLLPKGGWDDGEAIEKAVLREVIEEGGVKSQLI |
| >scaffold_10_F2 271 | MRLSVILLVAAFVAALDPATAANDANTVVAATNVHESIAIARFLARDEDDYPVKGERDEDDSDNDKEKDE ERMFSFFQEKAAALTEKATALSFAFKLVSQSGDDLVTAVSGLSKSEFVALFNQQAAMAKMVPGFPRGMSL DEFETVVRSAGLSDDLLENALMVGYGKYLALHMPK* |
| >scaffold_73_F3 95 | MRFGFLLAAIALFGLCESTVLNTQQKRNLRLSTTNDNEERAIGNISKVDDVADDASKNFAFMINMFKEWDE LSQAQIARMFKQTPDEFAAMWIMYKVYQKLGADDLIKTLKAKAK* |
| >scaffold_6_F17 3 | MRLFIKTLVALAVAFLATSTEAAKAVQTGGNVDDVQSSHIVPENKRLRSEDEGNLLEDDEENLLEDDEEE RKGGLNLFSTAKMEKMLGNDWYKYQVARRWKRDGHTWEHLPQHVPADLVRYFKGFRERHG* |
| >scaffold_52_R9 85 | MRFALLVAAVSLIASGDALSTQADATSRHLRSHHTNTYDAEEEERGLDKSIVKSLPEQFKNMYKPSNM ENVLESWRTGLQSVDDAVMYMKSGLMDFDAISHFVDAYRKHINKKGLPY* |
| >scaffold_78_F7 7 | MHFISFIAVALPFNVLAEEELAWELDCSVPLGFGPGLRLGVDFDRAVTFEFAIGFSFELDFLEWPRCWRRLVLL GHLSSNLRCWRVSSLRCVLPGRRLRRSFTSAEIAICIGEPGYTS* |
| >scaffold_26_F3 00 | MEAQGLFACLLWIRSVFPAGRATVFGDSTIVINQALMLNRCQAPSLRPWVGAIKALGTGRPAFYLQHVRRRA YNTASDALCNWMMDTLPASDVILRVPSGLVRHRTRRLRALPR* |
| >scaffold_431_R 1 | MKIQLLHLHVLFLVQSLDVLHVLRYGYYHAILREVHPVPLGLLPDANNHSLQRPLREILVTRISINRTGRPENHYL AQRRLRLQL* |
| >scaffold_6_F13 9 | MPLPVAVLAELEWLDTVSLYHVCTWSTAIERQQVRPNQFVRSVGLRSLRWCRRLHCKLRFHIAPVRRQT SVAIPYPLSCQLLSCSCQCFAPGIYCCLLRYSREFVRIPVHEDHFSLSIVLYHRHTAEIAFNHASSYCWLLDHV KRGVLMHPCKTKEILVTMREFEAVKEIRKVECEVG* |
| >scaffold_30_R7 9 | MRLKSLKVLALSACANRVTLQFSPAGTCASAVSALKHYGRAPARVPFVGHPLRHCKCFESRQRRPQQRKLR KTRLKWRMSSDINLE* |
| >scaffold_11_F1 229 | MKIYTVGAMLFVAATVSYALESRQLRSQQQTGARIVAGDGEETAGDDWNVLLSDRKRGAQAGTLVRD SGKDEERLVADSRRTQTYPPVQ* |
| >scaffold_10_R2 236 | MALGTVVVAHTLTFTHASSACLHFGCPAVKRTAVEDFDSRYSGASTSPPRVLRALRRKRAKREISYGVWRL HQMTAHTRGMRWSTGFCGSPPLLVPHLVFLATYVIRCSSAPKPSKR* |
| >scaffold_29_F6 61 | MRLNCFVVAALVASTNAAASSIQNHVSQTTPDIQAATQTDVVKRSLRYVKDEEDSIDTKDEERVQGI NVKELNKILSAKRIEIPANIEALSKAQRDLVHTFKGQDLTKKMFATKLGMRDVEDVANRHYPPFEKWEHLF RPGKKEKVPPEMLIGNHEYYY* |
| >scaffold_46_R9 61 | MRLAHLFVLAATFLVSSGALAATNTDSVIKTDVYKGSKPATHRALRKYKADFEDDDDDDDDEEERGIKDI PLERLQSLGRKVGVSADDVLSDAAGVMRTMSESIKQWTKGLNKLKQLYKKAKSPRISDY* |
| >scaffold_46_R7 83 | MRSFVLLVAVVALLFRIDAASGSSMKLRSGQLIPSTETRETEQIGRVLRGEAESDPESEERGITDLAKLLGK SKTMFDNNALATIVRLKSQFATADDAVQAFMTRNINPDVAVYKWLKLYKANNQVVKKTGDAPEYLLWIRY AKEYKDAHPKWVSKLPKGA* |
| >scaffold_29_F6 55 | MRLAYVVVVAVVALFASQDAVSAEADVQLGNLNVNSESTTRFLRVESTIEAEDEERALPSSLTKISEKLPVG DKVKPLTNKVAPLTKLTDKTMIAVRLKPIADFTGKLPKPIADKLMPIKPFKAAKLSAPVVQKIEKFKTFAEK IKNYKVGHTLSERYTMAKFENWFKQNKSPDDVKTLLKVGEGATVNTKNYDLSVQYNAFFQWATRDKEM KAKKAAAAATA* |
| >scaffold_710_F 2 | MRLTYITIVATAAILACCDGKSAVGDSPKPLAVPQSHVQTGNANRFLRVHDDEEERIGTQFITSLSDFASAA DKLTRSKSLNSISKVDDIAPGRAGKTLGEVLKIDDVAYLKAAGKWSKNLDDVEDAKLVSELSKLDHKEALRVLD EKNAAAFKSIEDMGYSPDGMFKMKELGKDYFDPFLKHYTRYWMAKHPTWTSSL* |
| >scaffold_2_F16 79 | MRISTLSLLLATVAVVLSCLASKGEDPSQSIANVDNIRGAGRFLRAQAQVDEERVNLKLSPESEKWLGLAG KSAAELEKAAQAPKNVNPEKLLAMMKANAEHFMMNLAKSGVTPKIMRKEYNIKRMKRVLSKDALMRNAN |

| | |
|--------------------|--|
| | YQEYERFKEFWNKLPKKLRFLAKAASLH* |
| >scaffold_225_R6 | MVFSVFFSAILLSVSASFNYNVNLETPRVQYRGWPTTWFRSPIAKILWERYLRNLADSIDIPLNPDIRGWHMS NPPLEVCNRYLLIYPFVEQRIVHSLGQFCSAGDLLRFV* |
| >scaffold_237_R14 | MQRRMSTPFFLMITTIGDAQGEVWVMPADNSPLIKSDNFSRNCGATRRTWAIGVALPTSMSQTAPRP AGGRAFPFPIRRLRFQMICLKRSICSGVLPSSASSIWISSILTSSVVALTSGSMAI* |
| >scaffold_614_F1 | MAFSSFLKFLAMATLLMVQVNAEPSQQTRNLRIQSDDLPEQARRLGNWKALIINVKNNPGVSDGVVNL LKKADFVTGTSSKIKIEKVKDTAGKVKDKVTGK* |
| >scaffold_44_R818 | MRFSYVLLVAATALLATANAIEADPTSRSLRAHKSHEKSQDEERAFTYTFNFSLWDDLFNLSLPEQFQRMKE PWYLRRIFRSWRSGMGTSDEAVAFMKSQGLSQKAIQFEDAFLKYRAHKLAKGK* |
| >scaffold_29_F663 | MRFLNILLIIAAVLTAAVSASEDAKPIQIKGANNPNEVSAHRLRAHKSDELDAEERVIPDVYHLSKLEKVAIYP KIVGWRLRKFVNAAMADLGMAREVGTKNMAIRWYQKIVKNGGFKHTPKGKYITQRK* |
| >scaffold_111_R171 | MDSTASTAVLIAMTALVDAQRLRSEASTATDSTVSTADFSTLSSGSGFTPGNVEAQSNGFALDKEGSTGGN RPSGSMQGGAGRGFGGSGHNGMQNVGSGEGLAHGSGSFGGHHGGFGRHGGHGRDEGSFSGGDFAGSF GGHDFGGSGFGGNFDRSNGEMPSIGSGEVPTSGSGEAPTAATTTSSSTSSVTASSA* |
| >scaffold_443_F6 | MRLSVILLVVAAFVAALSPTAAASEANTVAVPNVHESIATGRFLRARPEDDYPVNDERDEDDSDDDKEKDE ERMFSFFQEKKTALKAFKLVSESDDDLVMAVSALSKEEFVALFNQQAAMAKMVPGFRPGMSLDEFETV VGRAGLRDMEDALMVGYGKYPAYVM* |
| >scaffold_29_R1159 | MFPRLYAIALIVVALISTCGGTRMDEKQPLALLTTRSSRHLRSIDDRPRDTKTEERGNIDIMISDAAKKVKKAT WWKVKFAVWKHLRDKDPHDVGRILGMRNMGREALQHKNDLAAFWNFYGGKPLKYD* |
| >scaffold_52_R3 | MRICFALLAVTALTAVVSGSVNLRASQKIQSGDVAQYKTSGRELRGDLNTDEAAEERVNFGFLKPDPLKSF AKQQMKYAYNERIFNDLLKFDPPDALYTTLNLSKMKNL* |
| >scaffold_57_R893 | MAFSSFLKILAMITLITFQVNAETSQRARNLRIQDDDFPQEQARRLGGNWKAILINVKNNPGVSDGVVNLIK KVDFVSGTSKKIKIEKVKDAKDKVKDATEKVKDKLTGN* |
| >scaffold_592_F19 | MRLNLVLLAAVIALVSRCTAVSAAASSTNQINLSNLNQLNMQNHAGDGNRFLRSKIVNDDIDSDDDID |
| >scaffold_73_R384 | MRLSYIVLVVAVSFAYCEGASSVSDSERLAVPQSRVQAEQNSNRFLRVHQDDEERAGVEKILTQSASKLKRT VSNVDEGLISKLWSIRKDNDAFHFIETVMRLDPDGMMLLRMRDQGNIDLKLLKHFTKYWAEKYPDWVSKL P* |
| >scaffold_18_R458 | MRLSLVLLAAVIALVSRCTAVSAAASSNQINLSESNQVLDTMQDHAGDVNRFLRSANMVDDSDVDDIDSGD AIDDEERGKTLTFAWWNALGKTADDIYKYNLKLPLVRDAYKYNQVVGELKNNKNYQKWAAYSAYLKDRGL N* |
| >scaffold_16_F1314 | MSLLLSWRSLSLCLLSLNVSVLCLHGFIVDFTRCLRLLWFFGHCCFSGRHLRFLSCLNRSFLVHVLNVNF NLRFLNCSLHFFHRLNFSFRVNLVRFSGLLDLRFLVNCIRLHPFRKWESEGFQPVVKVFKHRLPFRIVEKRVEISV EHLKLLVTGRQSVVKDLRQFQTRDLVVRHHHEHRQFA* |
| >scaffold_63_F287 | MKLCHIVALTVALLVACVDPASASKSKLATQGDNIPTKRLLRTTEDEERGISVSLGLEKISNAFTSGKTEKFN LVKAGQSADDSFANLGLGHVNLFEKGGEVRTKMVAKFFTSDFKAWSKYTHKMNQKQDADSAMLGLTRA YGEKNAAVIILLGKDSLSSHTVAKKLETAQFTKWYTVDKIPNADELVKKVLDVPRSKLRKYPREMSIWDNYSK FVGKYVLNPRPGPPVRK* |
| >scaffold_25_F1242 | MRLSFLLPVAMIAIYCATCNANVASNQKLSMLQARLNDEAGGTRLLRVHHEEEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSCTRGRKYDRVYNGYMYRDIYV* |
| >scaffold_12_F708 | MRALKLLLVTVLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK STSGATNWLKDKKLAALKLRALPMNLKSVQKTMKDGDIPDRVFTLLNLQKSNRKNINGFRTGEYNLWKE LTVEWTKTYPNWVSNISKKPK* |
| >scaffold_25_R673 | MRLSFLLPVAMAAIYCATCNATADSDQNKVSMVQSLDARLNGQADGTRFLRAHHEEEEESDREERGLTDL FKTEKAAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSCTRGRKYDRVYNGYAIHLDY* |
| >scaffold_74_F614 | MAGRLCVMAITITLVAASVTPSQSLLTERQENDTSGRFLRVASTLKEENDEERAAAPVEAVTSLIKNKNVA NAAVEAVEKKYPLVIPGVTDLLIKARMKRWFWRKTLPTKVFKLGLRGHSEDRLKNHPFRKFYAAYLEKWKD AQAHNLVDWKGKAPALPGTITTTGNTVLKT* |
| >scaffold_1_F5431 | MKTATAFATVLALIVATNGAHVSPNTPTRLRGLRPTADTPSVEDDKEDRKHDDHHVKKVKKIAIPVVPVPEVPO YIPVPSVSPSTVVASSNNA |
| >scaffold_37_F302 | MALKIATLLSCLSLWLLFAETFYWKETLSLEVSFAFRPQTSLACVDSTPSNRRSTYRRWLRVNVPWQRLRTQS QCETVQADAQSRFSILGFCMEPTAIWSG* |

| | |
|------------------------|---|
| | RARPTERNFWLGSHPYRYHMKPQHFFMNRQMDRFPVNLPLILPRARQRQHRYFVGV* |
| >scaffold_29_F1 478 | MRCILYIALALAAAFARSDAVEAFTNADLSSKISHDFAANELVNESSQKRFLRVDPKSVSDTEERMKISSLKKIIKE LDVKDAKAKKLAIAIKDLAKARKDAQKLATNVVSKAQRTTRTKFSSLEYDNFMRMFKSNLAPEKAKSTGKIKT PEQFTRYEDFYETAKAADILTRMAHA* |
| >scaffold_93_F1 27 | MRLHCIVVLAVIAFATNGNEVSAGKSRVAITTTGALDTPTRLLRQTDEERAFGLNLLPGSKISSFMTDKK LSKYLKSNEQFDDVFIKLKDAGDKLFENPKFLAWAQYVDDFNQKHQTQNSMLPTLVRQFGDDLSIMLE KAKQADKTYGVALRLQGEQMKLWRREGLTTDMLFKIYKLLDDGATNLLNPGIKIWMRYADELFPGDSTLLF KKLQKTYSDAALSILINGKTVASSEYGEVGDGFAASSLAEGPCASGESFPVAVTQQGGRCVW* |
| >scaffold_13_F2 760 | MLVERDLLATEASQTDLLLLLLLLLGEPRRVCNFHWCSFTLLRCRLLRRCTCITSTRLLLVLTSLRWFLRLLW SL* |
| >scaffold_8_F32 13 | MCLQSIFCGAAVLCGEWLVIKRAFAVKCRLRSHSLPPAHAALSRGVLPRHLKCLLRYSSVSTKWHLYRVVQRR RWRNLRHQSRILFLVRLPLCQDKSPRDNHQFLVQVWDSIPLRWFGSRSQTTPL* |
| >scaffold_18_R4 28 | MIHICTFILLLSGERDFGVNLNKPFPNNHPLDLPFFSGNHADLLIVCLHKIIVSGYEKLSVYNCFLTIICNISPYC KKNLMVSAVRLRLFLKFAQTRYLFDNEANHHLVFFLETFDNIQYQYEGNQVYAMIQNKNVFYQLNDL QLPPIRASTEGKKEATEKTEEEKATDGEFVPTVEWLSAWKKKLPSTSLRLLQYLIPQLEDACKKAGGSLDEDA MLYFLRTTMMVGLLPVPHPIVIRKYQINQFTHLWFTFTWGVIFLRNQVLPFDGGAITLFTISVL* |
| >scaffold_8_R13 70 | MRLSFVLLAVAVSSLVATGHALPHEQPTVSKVTAANLDQSVTEGQTSEKRFRLSAAGIGRQMLRSESFKKT KFGEWLKDNLSSFTIFVDKLGKKEKYRRLYNEAALRKASGQYP* |
| >scaffold_2_F23 3 | MMGSVSAFVVWLKSLNLPPTTCTCSCPCDVQTCYRRELLLLLPVRACLRRLRLLRLLVPRRLEWDLRFLR ADLDLDS* |
| >scaffold_4_R23 6 | MRLTFLPVPTMAAIYCATCNATAVSDQGKTPTVHSLDARLNDTGDRRSLRAFKEKEEFDTEERGFLEKAAV KKVQGGNYERP* |
| >scaffold_29_R9 03 | MRLASILLVSALLCGSVFSTEGKQAHTVKNTDGRLLRSDDYDEERGFNWPWLKIKQLDETTAIKQLATDF ADLKSINREALDVFHLMREGGMSPKKATYISNLFKAYMENPRLYH* |
| >scaffold_50_R4 62 | MRFSYFLLLSAAILSNSNATATVSGEGHVMSTADAPARALETNNGKRSRYAADEDDKYDQKNGKYDDD EEEERNFTAQAQLAKWTEKAESWVSKNRTPAYIKDKLTGMNGLMTAENRKKYELFTAAYGRANPHALDRL* |
| >scaffold_25_F4 87 | MRPILLHAEFWTLWLVLAWKACCLREPSGLAFTTTYWSMTKTSYRLTWITSFASTRLSLLRMTTTRRRLRLL HLLQQTPRMYPYLYLKRMRVKRALYRWLQLTGCASTDASARFS* |
| >scaffold_7_F31 0 | MRSLLALAHHLILCLLPHPAYFHTLMSASRPLDTSMGLGFWLLPSSTSVRSKIQSVWAGDGGTRPLRSILIADS TPPPPEAATLPGDE* |
| >scaffold_50_R8 12 | MRLTYILAVAVVATLHSSATAFSSVKDSNAADSALAEGGRMLRGEYGNDAIADDDAGKKKVSYGDDALDY DDLDDDDLDYFEEERTLGDVLKLNLPVRAVKSAEKTEHAAKVKEALKDAADYQAMIARAKEMVNKD* |
| >scaffold_30_R3 24 | MGVWVLLLLDVALVAATSSVRTALFSLPVELCSFVFLTDAGRGLRPEASLVLAGDAVRGPWSKFCDFGAK LDSTVRGGNGPL* |
| >scaffold_111_F 98 | MKVFSILASAAVLIVMTVLVDAQRRLRSEASTATDSTVSTADFWTLSSGSGFTPGNVEAQSNFGALDKEGST GGNRPSGVSQGGAGRGFGSGHNGMQNVGSGEGLAHGSGSFGGGGGFRHGGHGRGEGSFGGDF GSFGGHDIGGGNFGSGNGEMPSIGSVAVPTTSGGGAPVAATTTSTSSVAASSA* |
| >scaffold_25_R8 04 | MRLTQVLVIAVASLLFARDTVIATSNQAKISKMEQSTPSQRLLRSNKYPIEEEEDEFEDTMDFEEDYTTTDE DLEERSPLSAAVAKLDDIASGWGTSWARVAMGQSSISEDKIKALLALRDAYISGSKKAKQAARMAVLRANN S* |
| >scaffold_81_F2 85 | MRLNCIVLIAAVASLSTAAATSATTDIKPVRPAINLQPPPLVVGRLLRTVQDEERGFTLPGAGKLADLFESTALK LTQSARINTWLAKGTSTDDAFIKLELNTAGSRIFENPKLLTWAVYVTKVEKKNPEEILARLSKQFTEGSLAKMI ASAKLDSKTEGLATILQAQRQVWVDAGKSSDEVFKLLQLDEAGTKLFKNQQFSTWTSFVDAFNRYKPEEA VSIFSKLAKTYDDFTLWKMLEAAKVPKTEIIASKLQAQQIDAWLDAGKSTDEVFNLLKLQRTGDKLFKNSQF LTWVSVEKFNKDPDQAIAIFSCLAGIYDQVTLSSMLEAAKHVPSTKRIASYLQGGQNNQHWLADGKSTDDI FKLLKLNTPSLENLIGPRLDAWTSFMRAFNMANEGKETTIALTLTHYKDRGLAQLLQEGTKFASTKIAEELQ TAQFARWLQLGKTEEDIFALLKLLTPTTDPEAIVFYRYKLFMDAHM* |
| >scaffold_50_R1 069 | MIVSFKTLFVSAVAVALAVSVEGYSGAVESTYETKRMLRLEAQDVAEEEMADDSECGSLEMAEDDSECGSL EMAEDDSECGSLEMAEESGTGDKGGDNGGNTWTQAPSTGDKGGDNGGNTWTQPPSTGDNGR* |
| >scaffold_29_F7 68 | MRLVSLFLVIIATLVATGNTLAVEHTEGKITSSDTLLQPTATFNPDVQTRRLRTPESPKDGEERRLVNLDIAIK DAVHEVRKLTWKWLQFAVWKEIGKPKQLIQEWGMKYHM* |
| >scaffold_10_R1 | MRLSVILLVVAFFVVALSPTAAASEANTVVAAPKLHESTATARFLRVRPEDDYPVKDERDGEDSDDDDKGNDE |

| | |
|------------------------|---|
| 133 | ERMFSFLEKKTALSFAFKKLVSESGDDLVMVAVSSLSKAEFVALFNQGGQSAMAKLVPGRPGMSLDDFVTVVKS AGLRRDMEDALMVGYGKYLAHLMD* |
| >scaffold_50_R9 34 | MHLLYLAVGVAISFAAGATDARVLASDDSNQYRAHAANANIVSTTRLLRARSVIDEERAGGISASASDKLAKL FKSSKVTDEQLQQWLSKGGAAESVFNRMDLGNTRITKLFENTQFLRWLQYADDLSASGKGTSAISILSTQYG DDTLKYMIESAKKTTNTKALGTRLQTEQLKHVVITIGKDPDEVFKLYGLNHVGSILSESQFSAWTKYVDDLN AKNEGAFVSIPTLRKHFSDDDLFHIALAAKRVDKTKMGTKLEDAFVQFWIHRKETPDNVLVGLGKSAETL LENPLLKILTKYTDAYNVKYAAKTTVIETLRTFGDDKVARMLQAGRTDSTKCKIAKQFQADQLEMWLNLSG QSVDDVYKLLKPSRRDLLVDFGNQKLFDTWLTFMNAISIKHPEKTSIFTTLAPTFRDKPMMQILEAANKFP SMEKAATKLQLEKAQSIFSTGISPYEAFKLAALDDVGDVLSLSPFKKWMLYVEDFNKKNPANKESWFLPLRG NYQGDRLDEIHKAMKDPSTVKLAKSVERERMKEWLEKWEYSPSTAFRKLHLKTAGEKVFNSPNFELWVKYL DDWNKAYPSNKMIDGFRDNYNDLNLVRLFTKAEKTPSSKKLASQLKDALADKWVAEKKTLDYVKSWSY HVPSSDDMLDRFTKLNNA* |
| >scaffold_2_F36 47 | MRLFWTLLVGLATSGVQPIAALSKVADTSETSTLDHGQSTIIEGTARHLRLAPHEDTSITDVLVERGLKVHDE ERGNFAAAAGWTANKAFNGLGKIVGHDNSNKLKTKVAEFFSILRKKGKTAGDIFAHARKQTDPSQRAKFET VATMYQQYLTKVL* |
| >scaffold_644_F 7 | MAGRLCVVMAITITLVAATVTTPSQSLTERQESDTSGRFLRVSTLKEENDEERAVAPVEAVTSLIKNKNVA NAAVEAVEKKYPLVIPGVTDLIIKARMQRWFWKRTLPTKVFKLGLRGHSEDRLKNHPFRKFAEYLEKWKD AQAHNLVDWGLKATAALPETITGNTVLTK* |
| >scaffold_104_R 67 | MRPNLVVLAAVATLVSRACTVVSATSTNQNINSLDQALDIVENHAGNGNRFLRSANIVDDSDDDIDSEDEE RGGKTWAQKFAKWNQRGESANDVYMRFGLEPLVRKAYKYGGIQGLQKNEYRKAAYSAYLKDKGLN* |
| >scaffold_36_R1 210 | MRFFMVAVAFAFASSCETASAEAPNAFLIDDNTNAVFSRYLRSTQTDSEERIAQVLKSEDRTVANVMIK YQSLYRAKITPKQAKLMLGVSDNMVFEFTKMPRIQRFYTGYYSYFTTMEKRKKWKKELKNQVMV* |
| >scaffold_379_F 2 | MNIAVLLCFFAFPSGLFFSFRRLRAASLAPFLS DIRWHV LGGPSLQHFTPKQQQLHALPLQAAAYLTSSAAFC QLHRRALISLRL* |
| >scaffold_20_R1 237 | MLGFLLLVMTVGGSTTEILGGWSSAAITPDTRAVLVQALSTTNVCVDSIISVRSQVVAGTNYEFRINGCRGLR GGNCVRASCVAPRTFVVNVFDQPWTRTRTRVMSVSEI* |
| >scaffold_24_F4 23 | MRLSFLLLAATFALLSSGTVASPTTNDPEISSPNTVLSSETTEGRHLREHKNSIDDTEERGFNPAKFDRMLNERS YRQTRFGNWVDKPYTDMDVYKLLRVESNPNYRRIFHYQNYLQEFAPHLITP* |
| >scaffold_2_R38 00 | MKFVSTLVFMTVALQSSCSTAQIDFVRSYDHHAI DFSTSSKPTQPSVADVNTVEANQPLRSGNRRRLDGSAA GIVIVGRNGGGFEPLRNGETPD* |
| >scaffold_11_F1 235 | MKAKLLLVLSFLLAVTSVAERHLRVADDANPLKPYEALNAVIEKFKNAKAGKPERRRLTGEKAFDKFVKAIA VLKEKD HARRVEAN* |
| >scaffold_55_R4 | MRLFIKTLVALAAVLLATSTEA AKAVQTGGNEDVVQSSHLSRESKRVLRSEYDEEKLEDDDEEELLENDDEERK NGANLFTDAKMKMLASDWYRYQVARRWKERKYTWQNLPPNVDPNLVAFYKGRFRERHG* |
| >scaffold_55_R6 23 | MFFLVKQLSAVNHMIRFSLASFLGLFKGTATKMESSSTKDRILRLIPILEHKVLMFVGRALFKEHRPFGMH LVHGMHPECFDKNEYEFFCGEVVELERGS GGHSTLPEWASPERKEAFTQFVETLPRLTQLCKFESHDMWIR WSKSMECEQNFHPKMDKSGSAGGLSPFQKLLVVQALRPDRLQSAIIQFICGVMQLKSLTPPSLDFKVGITEE ATNTTPVLLTTAGADPSKELEEVATSVVVGKGYFEVAMGGGQQEKALNLLKSTAEHGEWLCLQNLHLVVA WLPVLEKEFSALNASHKFRWLWLTTEPHDAFPLVLEQLKITFESPPGMKKNLQRTYAAWNPAFIAKGN SAR AQLLFLAFFHALLQERRTYIPQGWTKFYEFSGDFRAGSNVMEACQTS GAAIDWQTLHGLMENAIYGGRI DNPYDLRVLRCNLTEYFSQDLLSGHKSLTRGVKLPQSTQHADFLDIIDRFDPIDAPAMFGLPDNIERSMQRSL SGQVIAQLKALSSNEAEATTFDREKWRAQLGPLLETWSKLTTGFQLEGTSLSSSTGKNLQAMTPADAFVALE NEYALDLVQQVNSILQALKKVIYGTGLLTPAIQTVAKALLKGVPT EWAA* |
| >scaffold_325_F 4 | MLSVLSLALMLAVVTGAFETS RDVKNVGVTPDSRRLGFTADQEERKFGGPATGTDHSNSHAWENFKAWF VRTFFWRKREQTRRLRNYSSM* |
| >scaffold_50_F4 3 | MGLFYLVCAILAFVARDATGDRVIVSDGFNQHRENAARASVVSTTRLLRTRKSVIDEERVGGIPVSATDKVAK FLKSSKVTDKLQEWLRKGTAEVSYRNMNLKTSQLFENPQFLRWLQYADDLSASGKGTSAISILSNKYGDEK LYQMIRWAKEKLNQDGLMRLQTEQLEHVVVKVGDPEVFKLYLNHAGGGILSSSQFNWAKYVDDLS AKNEGATVSIPTLRKYSDNLIKIALAAKEVDETETMGMKLEDA LMQFWIHRKESPDNVLDLGLKSTGT LLESPLTILT KYEAYNAKYP SMKTTMIETLRTFGDDKVAKVLLAGRTESTTKKIAKQFQADQLDMWLNLSG QSVDDVYKLLNLPSSRRDLLGDFGGEKLFDTWLTFMNAVSIKTP EKTSTIFSKLATS FEDRPMQILEAANKFP SMEKAATKLQLEKAQSIFSTGVSPYAFR MVALDNVGDVLSLSPFKKWMLYVEDFNKKNPGKEDSWFLSL |

| | |
|------------------------|---|
| | RVNYQGNRLDRVIDKAMKDPSTMKLAKFVEKENMKEWLVRWEHPPSVAFRELHLNKAGEKVFSA PKFEL WVKYLDLDDWNQAYPSKKT MIDGFVDNYHTLDLIPILAAAEKVPSTKKLASQLKDALVDKWVAEKKT LAYVKS WLVNGVPSSDDMLKRFATKLSA* |
| >scaffold_11_R1 430 | MRSYHFLLLATAALLSSCNATAAVLCEGQVMTSADAAVPRALGTSNSKRSLRYDYDEEEDETDKHHQKND KYDEDEEEERTWSAAQIDKWTAKADEWVDLGKTPAYIKAKLTAFNGVMSDKNRKYELFLAKWGRANPDEF GRR* |
| >scaffold_22_F2 08 | MALCSCICTGIACFPVSSHFAVYFHLSPHSLLSLRRCHTDTAERLLRRVLLIARRTQEQRRAEWLQVATQR AERSGSRTIQHGHTGAGWTCLLLHRWVVISLFPRLHSHSTGHQLVSL* |
| >scaffold_29_F1 480 | MRSVYVVILAVLALLCNDALAATSKTDLAEELTKNERNRSLRGEVIADEDSYAIDEERVQTSGISKVAKELK KANLKKVSAALEKAKSNPSKLAVNAKMKKAIKAEKILAKLSSMENDNFLRFLKANFTPQRAKSTGKIKNPEQF AKYEEFYKLAKLIGLK* |
| >scaffold_3_R39 1 | MLPTELVLFAFVVGRLQAGYAPPPATTSEWPLLFSPGLGNVWHTFAFLRINRFLRIVHLRPLSDQLQRFLLYD RRLKRLTPGICYLVRALDFLLGTHWLSCLFYGVSYLAYDDGEMSWLTPDMLAFGDGVRDLADIRKVP LLQS YLRTYHFSIGAITVCYGDIIIPMNAQETEVMAVIFISVALFMSLGGFYKYFDMELGRRAEYEERVAQVGHFL KFHRFSDTWRQMQVYFALSWSRESRGRERELLSGLPPSVRQDLAQHVHASLLKNVALFTRCDPTFARAIIA ALQHEFFVRNDVIIQRGDMERSLYIVESGIVLISAVRKRQVHAQAGGAEQASNDPVGADDNRSPGVSVN EWVRASVGSALRLRKAKEKQANTPRQRQKSLVALMSPTGTITDINNQTREEKIYKGFDFYGERSLLLG TPRNATCMALCVTSLFVLTSARFEAILDEFHERSNSVSAWVMTRTPSLPPDERPIDN* |
| >scaffold_7_R68 1 | MKACVYLPSSLFLPCPSTSAEPTDPEDNSTPHLDALERLLDRERARQLVNRLLRRPDQEIDLHDSIRWH VRYGPERIPSPQPSPPTDFIETYGDEYAVSDDNSSDSGTESEQSC* |
| >scaffold_3_F35 36 | MALVFFRILLVCLQIGLDEEAIEDHPTRKTRRQLQAGPHEASSEERVEETHEPEPMARHLRQFPTQHFDIV QSIFGRIGQQIRRAGIDS* |
| >scaffold_61_F2 09 | MRLFLLLVAIATFLIASEAFSTTGDSNQIYNVDSVPGPRQRFLRAHAEVLDLEKEMKMMKMMKRKVTKEDE AKTLKITDQIDDIINKHAPGMHEFMQTPKYQRYSNYMNFLNDMAKKPEYAAALVEEIKAKSRAQVALKTRFKP TTSQNRWQIIFASLKGKRIGK* |
| >scaffold_23_R4 90 | MQRRMSTPFFLMITTIGDAQGEVWVMI PADNSPLIKSDNFSRNCGATRRTRWATGVALPTSMSQTAPRA AGGRAFPPIRRCLRFQMICLNRSICSGVLPSASSIWISSILTSSVVALTSGSTAI* |
| >scaffold_25_R1 22 | MYRILLTTFALLCGFSNAVNSDARLLRVDPTEEEKGGFTTYVKDSLTKWRINSKIKSWVKNKKTDEYVLKKG LSTLTGKDLVNAPKYSQFQDFKVG MWLKEATPTTTVFNTLGLNKVEGAVENADDFGTYVKYVITLGEKADN YPLSQWPRLFGGGSLEQLLEKRKILRLQKRNVL EINFMLG* |
| >scaffold_640_R 2 | MRICLCYLLSLTAVVSGSSVNLRASQKIQSLNVVQARMSGRELREDLNTDEATEERVNFGFLKADRLKSAF QMDETHIRTKSMNHKRN* |
| >scaffold_38_F1 173 | MRSYCVLLLVYAVMSTAGTESTAMDSTQ PSTQYDIVLHVRTLRTGEYTVSEERAAGASAVESLGKATTSVD DVETWLKRGDTANKVFKALKLQKAGDNLLDNPQLSTLMRYLRFNEANPTKKTSLIATVTKHYNHGLTKIIE AGLASTSKKTVATAKHLQTEQIHYWMAHGRSPESVFGVLIQYTKDALRRSVLLKDKLLPFTWISSDLFEKPG LKTWIRYLDEFNVRNPKQKTTLSILSKRYNDKTLANMLIAANKAESTSSIAKRIQSEQLTQWLSKGGKTS DLFTMFQQLQKAGATLFQNLFP AWIKYADDFRVIHRDTQLETMTTLMKHFDLAKMIMTAYDVPSTKSLAS R LHAELLRGWQWRQETPDDLYILLKLC* |
| >scaffold_25_F1 656 | MRLSLLLVAALVANSDAAPQSTSSLTKFSTDVAPVRSR LGASKTKTEVDEDDSFDP EEEERGISMYGKLDK VDDILKQLNLADDTVLK KLIAGKGGVEQLITTDKGVKTLVITSKKTGKKVFTFNIDIENIRKIESTPAIKKQISQ WKKLQLSPLAVSRDLRKKGIPRTTDNVMWEAFRLYSATSGRKFNSLRPTGL* |
| >scaffold_77_R2 49 | MWMTQYHLVVAPALSILTSARDEMGTQEQTAMLKEYLDMAS YTVDDLKDKPSRCRRLDGKTVQDHLGP EAYQDVARALRVLQVWEKVYPKASIEDELAFDLRLPQLNSTKS* |
| >scaffold_1_R40 7 | MRTATAFATVLALIVATNGAHVSPNTPALRGLRPTADTPSVEDDKEDRKH HHHVKKVKKIAIPVVPVEVPQ YIPVPSVPSTMVASSNNAVIGPSTNVAGPGAAPGAPGAPGAVTPAP TTLNGRPAATPAATSTTRS RPTPA PTNFAGARPSGAQLPAAPVQAGAPGMAGFPNSVGIGDNTNNNAFGAGIGAAGGFPM TGNLAGGFGG NPM SFGGANGMGSGGMFGGAGPAMGRLNGNDFGGNFGQSMGFGMQGGNGLGGFGGQGMMA NFGQTGNNGFAGGQALTQGGFGGNSFGGGFSRQRRR* |
| >scaffold_43_F1 201 | MRWLCIKQQVWLQSHCLMVVSPFLVCCPCLSDSNVFTSGW TMMLLGSQMWKSSLQNLEWLAAISSACL QMRPRLLRMQMMLFALVSTSLLS* |
| >scaffold_6_R23 12 | MLLRIVIAVAFVTISLVATGSEASSITVVDVADRSTQVKRSLRLRNLESVDEDRGPIAGLEKVD DILTKEKAS RKAGKVP SGLKNLLLRNIDEFAEHSKLAKKLSGLKLYKDAGLEKMSLSTLRQLDDIEVKRVSDIKNGITGNKDT |

| | |
|------------------------|---|
| | GGMRRKMDHVVDVAPAKYLTSIHGRGDQLYGADGSRLSSAVVSRPAEQGGGKVLIISSSKPEKGDWLLP KGGWDKGEDIAALREVMEEGVVRPVLFFLGLSYQYSN* |
| >scaffold_405_F 4 | MRLAQILVIAAASLLFAGETVAVATSNQVKISKVAQSSPSQRLLRSNKYPITEEEDESEDPVDFEERGEEDLEE RSPSSAIVSKLDDIASRWGTTYARVAMGQSTISQNKIDALLSLRDAYVSGSKAKAAKIAVLRAND* |
| >scaffold_9_F15 5 | MPTLLVVIIVSTAMAILDMTRTLRIRISICWYTRSIEFLLSLPLRWCDSSRLINNSILIHVHRDRFIRCHHGRMV WVILFLFSYRTVVSHWRSSGRRVSRVVFIFTMRSIHR* |
| >scaffold_42_R1 91 | MSKVLLLLVLSVFALVSCDALSAPVGSKLSLKTDELNAQPIDAKRMLRAQEPTNAADEERGMTELANKFK AWAAAIAIKTWTNSKLVQSMNNKLASLTQKGRVQIEKLLKQDNVNVNVLVYQNKVKPDELFLALKLDPKLL IADAPAAWANNPGLSMFYQYATYYAKMTTKA* |
| >scaffold_73_R2 70 | MRLLYLAGVAILAFIAGDATGAKVLPDDSDHNRAQAASASVSTTRLLRTRSVIDEERAGGISASASDKLAKL FKSSKVTDEQLQQWLNKGKTAESVFYRMNLENTLYTRVFKSPQFPRWLQYADDLSASGKGSAPISVLSTKYG DEKLYQMIGWAKKESSTKALGTRLQTEQLEHWVKVKGDPPEEVFKLYDLNYAGWRFLSNSQFSAWTKYVDD LNAKNEGAFVSIPTLRKYFSDDDLKIALAAKRSRGDTEAMGKLEDAFVQFVWHRKDTDPDNLVELGLKQST KTLLESPLLSLLTKYTEAYNVRFATKKTIVIELVRAFDETVARMMLLAGREKSTTKKIAKQFQADQLEMWLN GQSVDDVYKLLNLPSSRDHLGDFGGEKLFDTWLTFMNAVSIKTEPKTSAIFTTLAPTFKNRPMMQILEAANK FSSMEKAATKLQLEKAQSIFSTGVSPYKAFKMVALDNGDVSLSPLFNKWMMLYVEEFNKNKNGKEESWFLP LRGNYQGHGLDRVIDKAMKDPSTVKLAKLVQENMKEWLVRWKYSPSMALRELHLNKAGEKVFSAKPFEL WVKYLLDDWNQAYPSKKT MIDGFRGNHDLVPLMLAAAEKVPSTKLASELKALVDKWVAEKKTLAYV KSWLKGISSDDMLERFTAKLNSV* |
| >scaffold_300_F 13 | MTGSTPDHWLCFVISSSLCATTGICALHCCVSHSLTFQLDFQRLRLQEQQSMGVTTAPLKSPMGSPAVPY EPVQTMWEARPRKYKSILFEV |
| >scaffold_39_R1 000 | MMLVLLAICETIHVAAGTSLLFESDFAFGVDSTTSACAIIDRPFDPDKHLLYSNLTNLIALARCTVLERLEVS CKDMDQNRNKFSSRLRSYLLHVPVALYSTERTSYHQGRNHIHYHVGLI* |
| >scaffold_7_F14 94 | MSSKAGLISCALWLLLLVSTLGVDAHKRLLRSEVSHDNLEDRVLDNWKLSLLVKVGSRTKADKLYAKHVVG KITGNFFDTNAFAAWFVAVQMAAYAKTPAKAKVDMVSSLTARYGDQALAKMLATTEDDKFIREMKAIQLDN WQKDKRTVGSVYKLLKLDKEQDELLQSPLIATWIAIYATKLDNEDPLGAVFSTLKTQYNGKDFATMLLNKDT DDSFVVAEKLETLLMKSQREDKSVVDVYKLLNLDNEGDLFFQHPLIDTLIRYATVVDKDSFSGVFSLLQAR YNEEKMTDMFMTMRDWWPRNILDQLEDLLKTWQRQEKTMDDVFLLKLEQQGDSLFSKLLSTWVSY VAKVETNPYNVVFSLKSTYGEETLSMIIQARDMPTADYVGTGRRIRESSVQGLGEGQVHCP* |
| >scaffold_17_F9 40 | MQSVTSLACVNGAFLSSVAVIAGSDRSLRVLVDVGAGGGGRTMRVVRDAHSRAAHTVALPRPTCYTSHPSNF YDLLSSAPDSTTHLWDIRADNCVMRFCEHVNRVHTLGVAFSPCMRYVATGSEDRAAYIIDIRTGRRVLK GHTDVVTSVAFSPLHPQLATAACDGTFRFYSSTRSD* |
| >scaffold_21_F4 92 | MSSRLNLLFTGSFLLMARKEIVRIFHFLLLVNCSAIHATVSSSSGKSITATRHRRQAKPSVSRARVDG* |
| >scaffold_2_R24 30 | MGLPLMSSVFPFSLAAGLGGSHATTTWNLADFVSRLTSELPISLYGVMAGDATHLHHVLAHIGFALFSGV DVPRVTRPSIRTWSQDLVGE LRLLRSHALPTEVLEQVTGSNERRSALGNELLRVVEPFVSELVDFLVRATSAS RAAAFGTNSATFLRTMTQIVRQLRVYARGESTESEDSDERLKRLLRGLLVWLGMMENMARFVIDSLVCW AEGDNSSIRGRTRQREENSTDSIPAIRQRE* |
| >scaffold_30_R1 833 | MSDSPAKISAVCTALLLLAPFQWQSTYIPLPSGLLDFLHSPVPLVFGCHSLSETAEWSDVCFYDIDRDRIAVP AVTRHLGPSSIPNGVELCRLLRKARERFCSLRPTSKPWYELSEEQDTIITLTMQEAEIFLRDLGFDISSQDLAASI SGKSSVLLLLREHEVYSRIR* |
| >scaffold_40_F1 027 | MVVKKVGRHLHLSFLELGDCAFDFHCRLLTSRVHSIASSASLRPSRWLQSSRSSTLWTCSRLTRSSGSTRTR TCMTNCNRCWRTIRRRSRTL CVRRSRYHCRRLRDSAVTLV* |
| >scaffold_5_F25 81 | MRFHVLVIAAFVLSIDTFSTVSATNTVPSRGLRRIIDEERAGGAI SVTSEKATLKFSSKATDKQLQKWLQK RKPAEDVFYRMNLAKTGTGIFDNPLFIKVVQYADDLSATTSGKGSIASTLTAQYGDSDLKMLNVAKQDSK SKELASRLQSDQLEHWVTIGKDPSEVFKLYDLNHVGGSLRNPNQYNSWTKYVDDLNAKHGGEVSMIPTLRK YNYDEDLFAIVGAASVDALKSAGVKLENAFVQYWINDKQTPVKVLAELQLGATPKTLESPLFSLAKYTDVY NVKFPQSKTTMIETFTQAFGIEKVAKMVA AAKETEGKAKKIATELEAAQMOMWIRSSKSVDEVYNLLKLPK TLVIDLSSPLFSTWIAVMKILSIKNGDEMILQIKTLSMQFADRPMMLQLQAMEKFPNIGSTATSLQLRKAD DIFATGVTPFRAFKMALDVTGDSVLSPPVFTKWMSYVDDFNKRNPTKEESWVFSRSTYEGDFMDKLIETA RKSPKTVKIANTVESERMKDWLTRQKAPEHVHFHLKLNKGGEKAFSSPNFQLWAKYLLDFNLQYPGEKTTM IDSIRANYRDIELMPLNEAAKIPSTEKLANKLQNALRDKWVDEKVTVTQLKGLFGHMPSSNDWIQKYAEKL |

| | |
|--------------------|--|
| | NKLS* |
| >scaffold_39_F70 | MRLHCFLLAVATTLAVLNNGITTEASSLRKVPASAPIDSINAVQPETRRRLRSAETVYRHEDSYKRRPFIEEKLHKALTNPKKTKRLYARWYKSGFTKKQVAKGLDQSENRELDVYENLAKGYAKYVDARRTQQQSV* |
| >scaffold_48_F1199 | MDTSGACVLAVLLFVDPNTDTPQKIVLNVGDCRAIIREAPESSTGTLKKGKAASGMTFALSEDHCAANTKERMRALRSGAYIQNNRIAGVLEPFRTIGDIDVKGPDMDWVIPTPEIHQSELLVGRSILVIATDGVWTVLNNRRAMAHAVKELNGGRSSAESAAQAIKEAREFGSSDDITVVVVSV* |
| >scaffold_77_R178 | MLLCVFTVIAAIAVSDTTATNAIDDDGVLQTTSTRYLLGFNQEERGRVGGPATDGGTDKVSDDWWPNFKAWFKRTFYFWKKKETRRRLRDQADY* |
| >scaffold_15_R3220 | MNNLGRSSSVFFTAFFLFFESPCSAFSSINAASSASLPLAVTASCPNLENITRKLRFNAKMYSPLLERTPSTAAKITADLDWITLQQ* |
| >scaffold_18_F2149 | MRPNLVVLAALVSRCTAVSAASTNQINLSESNQALDTLQNHAGDGNRFLRSANIVDDRVDIDSDDAIDNDDEERGEKNWPKFAEWHADGKTAEDIYKYSALSHVMRYAYRTGSIQWLKKNYRKYWAAYSAYRKDRGFD* |
| >scaffold_9_R771 | MRGSSQSSQQLCMVLLIRILIVRQLRSLAHQEALLPRAHETSFLEHPARVRHRIVAPHSPQAVGYFGYEPLEWMTVLWEIRSV* |
| >scaffold_42_R407 | MRIYFVLLAVTALIAAVSGSTANLRASQTIQSGDVVQDNTSGRELRGDLIADEATEERMNFGFLKPDPLKSF AKQQMKYVYNERIFNDLLKFFEDPDALYTTLNLISKIKNRSNVNGVKTSRYNLYENFLISYLDKYPNWRSTLK* |
| >scaffold_5_F2028 | MRLSYTLIIIAVAVSSGNAVATTNGRTTELSAMASPNAVASVDTAVGGEKSLRYHNNEGLEDESDEALLEE EERKYTNMFSTTKLDEMNGTKMMSRFRKWKARGYNTYNLPAVTQKDKYTWIRQKYRDFLYHN* |
| >scaffold_73_F95 | MRCFYLAVILAFVCGDGTGTTVLASDFSNQHRDHVSSITISITRFLRTADASTGDEDRVGGVSISGTDKIAKLF KSSKVTDEQLQRWLSKGPVAFVYHGLGEDSLYEVI* |
| >scaffold_42_F41 | MANKHDACQKSLFLFLFLSLQLPLSAQSPSCPTESPLPPSAPDWQAAPPFWPLFSIYDVVTSSSPPAHWTSRTLRSATQPQP* |
| >scaffold_5_R2132 | MRLSYVIPAIAVTFASSGNALAAADGSNTGLSAITSPNVVASIDTAVGGEKSLRYHTNEDLEDDSDDEGLDDAEERERRGNMFMSTTKLDEMMDGTQLMSRFKWKQELKYNMYNLPDILASKYDELKMYRRFLYNN* |
| >scaffold_44_F578 | MRFSYVLLAAAAALLATANAIEADPRTRSLRAHKSHEKSQDEERAFTYTFNFSLWDDLNSLPEQFQMRKEPWYLRIRFRSWSRGMGTSDEAVAYMRSQGLSQKIDQFEDAYIKYRAHKLAKGK* |
| >scaffold_6_R3203 | MRLAIKTLVALAAAILATSTEAAKAVQTGGDVHVQSSHILPGENKRLLRSEHDEGKLEDEGEDDEEERKYG |
| >scaffold_11_F1232 | MKMTALLTVLTVMLAFVSADNSPRQLRSTLTTSIQDTDGNRLLRSEGLISNLLGGLNSLLNNQQSTNPPTTSTDPTTSEQNPDEIDLLRSRRVS* |
| >scaffold_31_F1104 | MRWYRVLLIAVVSFVISAQESESHIRTRSLRGNVAVNYSFKDVEDGNADERAGEKEVIKLTSSKTMKLSRSVTKAANSFSEKIPALPIKAKLQVWSNTGKSVQFVRQELGLTDLAEAAKKTQSFKYDDFVTSQLPIWAKKDLTPDEVIEQLGMKGLPAAWFKADPNFKYDNYLKVVPYWSKNNVEGDVVKMLNLTLSGAARREAVNFQYYDDFLVSQLRVWVDKLDLPGVTVMAKLDDLKLTGKEILHPNYEYKYFVKNRKLKAWATEGSLDDVAVRLGMDLHGQVQLKAHPNFKFLEKYQAKAFQYQEGWLKQGVTTFDMWNDLQVYRVPMSILRRSNTYNTYKNYVNVVDNYIIRMKKGVPDLKPLGLTSTKDATPHELREKTLIWTSAKRPEWYVKFALGLDGLGENALKEAANYQFYAYYLEAVKFA* |
| >scaffold_12_F482 | MRAHFLLLAAAIFLATSVDASANQAELSKVASPQSIETANIAPKRFLRTNKYELDEQEERIGALDDVIVKAGKTKMSNAELKKLVTPKVLENALSDRNMKLFKDLYAGKVSLLKRFKAVMKNNPNKEEVLKSYRWFERNYHISQAKARTAV* |
| >scaffold_30_R1705 | MVSAALASSGSLSGSLVAEPLLLVVLVSRSDSVALDVVDLPDDLDPDPPELLDVVPLVVALSSSPPLVTATMIPTAAPMTTRIPTTMRTLRAATVIF* |
| >scaffold_5_F1718 | MQLLRNLVFLALFAGVVAAVQEEQPQKTHLRGLLDKIVSSDPTPAPVRVLVASWEDGKVGGEARKLRPIAEKIVGGETVVMFNDRRLLSTADDLESVNFVDRSNPNCVSVCGELTKWYLSGSEQCFDLRIAPKSADQIENGLIRGVSAAYSGLRPVAKNLYLKFDETNFIDGSRCEYEILSGSKTVREEEIASGSQAAMA* |
| >scaffold_25_F1308 | MRLSFLAAAMATVYCATCNATVDSQNKVSMVQSRLNGQADGTRFLRTHHENEQESDREERDLTDVFEFKKAAVKKLAKEVMADNRRRAKDVFLWKEKGYTLDELNTFLKSAYQHVVYNQYMIFRQYV* |
| >scaffold_53_F91 | MRFFLVAALAAFAFASSCEAAVAEAPNAVQIDDNINAGFSRSLRSTDSERAIQLLSEDRAVASVKVYQSLYKAKITPNQAKVILGISDDMVELTKTTRSLRFRYTYGYSYTLMEKRRKRKKELENQVKW* |
| >scaffold_11_R969 | MILSTWLRLSAFLAAIACCTTQLGTPRPSMWINVTCSNTFGRSVSRKIMPDTRMYVRTGPSGCCDAGQH KWLIDRILRAHTRRPVSRSTKESIYSRKHSTRPSEFDDGGFHSQDQRMMAAPRNLVSDGSPVRTAVASAPSPAP |

| | |
|------------------------|--|
| | SAHIRSGPDTRLLPLPACRTGRMSSTWPWTWSCCRRGAPH* |
| >scaffold_98_R3 65 | MRLSFLVLAATFALLSCDTVASPMTKDECNPSPNQVLSSETEGRRLRVHKSSIDDVEERGFNPEKFNRLMNE QSYRRKRFPNWVSKKYTD RDVYNLLRVDSNPYKRIFNYQTYLENFAPRLISS* |
| >scaffold_66_F2 74 | MRWTSILLVAAAALTGVLDASTNVTANSVGMESAFSPVIHDHRSRLRVDDDDTVDDDEDESVDVNEERT WFSNAKQALADKLALTAVASNFAGKSTDEMGEVLKLSRDQINTIFDKGEDSIRQILPGFKSGMDSKKFDDLI KALPQEQQGVLLSAYGKYLFNGLL* |
| >scaffold_5_F17 02 | MTLLAFFLEFGFCVLAMAAKPTFFSCCTVGPSALLLKP KRELVC TRSNATSSSPSSSELLSSSESLPSSPSRRA LRFCSRSSRVISSRDVTNLRVAVPFSRFLRPLPPAAPGFPPGLPLSLIIVF* |
| >scaffold_10_F1 990 | MRVSVILLVAAFAVALNPTAAASDANTVVAVPNVHESIATGRFLRARPVDDYPVKDERDEDDSDDEKDEE RMFSEFFQEKATALSFAFKLVSQSGDDLVEAVSGLSKGEFQALFNQGAHMAKMVPGFYPGMSLGEFGTVV RAAGLSDDMENALMVGYGKYLALHMD* |
| >scaffold_26_F3 52 | MERVGGLSCIAVAGATVALVEALLSGLEQVRVGHHSREVVQARKAIAYVLGGARRVEAQHSSASVRSHGL RPRLKAEGLANCAARSLRLARSLLRGRNCEPRRERGAERHCKAAGWDQ* |
| >scaffold_96_F2 12 | MRLSHFFVVVAAAFATGAIADSEPNHRNLRKHHDTLVDEERGIPKILDQNIPIERLNSLLKKMDQRKKAELA KLIKLYESSPKFGRTKPSNS* |
| >scaffold_92_F1 4 | MRFSQVLVVAASLLFASETA AVATSNQAKISKVQSQSSPQRLLRSNKYPIKEEEDSESDVDEERGFATPD EEEEERSPLSDATVGKLNKIAKGWGTTYGK VAMGQSHISEAKAKALLALRDAYISGDKSAKAAARMAILNA NHR* |
| >scaffold_853_R 1 | MQRMMSTPFFLMITTIGE AQGEVWVMI PADNSPLIKSDNFSRNCGATRTRRWATGVALPTSMSQTAPRA AGGRAFPPIRCLRFQMICLNRSICSGVLPASSIWISSILTSSVVALTSGSMAI* |
| >scaffold_42_R6 7 | MRFCFITLVATTILASSGNISAATVLRNRSVTTSIDAVQPIEAAQTNSNRSLRRRNNYKDEEVAEEERSGIESLA AQLNKKLLPTIRQVAHLDLGRAALTQQMSLSFDERQAIQALLQLSKKDRKAVLMLIK* |
| >scaffold_7_R19 13 | MRLLLWTLVILVILSSCDAASV NENKSLQRKLYTKVASHALAADDGFEHDKRALRGASNGVTEARAATVST KFGSRLMAFFRSIKDKYLAWELKILVPGFEKMAKKGTTYQVREDFRTRLNWSGLWGTPSGFKRYAKLYRT WLEKNHYSQLAV* |
| >scaffold_8_R16 99 | MRVLGAIFSALFLAAGIDVGSLLPNSGVAFVNPVPHSLAVETVESKRLLRSYEADSEDEERAGGQLGLVDKFAA KVMQKLYKNPSDVFKRLKLDVNLENNKVFEGWL VYVNKFRQVKGVENFPDQALFNVIHQSQYYQRDLVP LFQSLTHVQGMKDLARTMQFKLFEAATPATR TLMNKAWLEGFDTSDDVFHILKLDGVDHSDKLIQWLK FSDMYKKLPTSQSTSWLDELNLVLKTKKPNQQETKFGLLFQALKEEKG METIAGKMESQLFERWMKMDSM TPDKVGGMLGGSATTNWKRIEFKLEFTDDRYIFLKAYTEAYAANRGANVLKSVEKLAENKPVAAALERAIAK* |
| >scaffold_87_R4 30 | MRLSQVLVIAAATFLFASDTVVIATSNQANISKIEQSIPSRLLRSNKYAVKEEEDSESDVDFEERGFATPDEE DLEERSPLSAAIVEKLD DIASRWGTTWGRVAMGHSSISDDKIKALIAMRDAFISGKSDRDVARAMILMANK S* |
| >scaffold_15_R1 164 | MRLTSAFVAAASIFVCFQEA SAVSDSKGPESRVQTANSNDRFLRIDQDEERAGPQPSKSLLEKFTSKVTATK LKRKAVADLTKLDDVVYLKATPGYNVPLFQRIEKMGFNP DGMLLKMRERKIDQTLKHYTNYWKGYPT WTSNPSV* |
| >scaffold_17_F1 758 | MPVFDALLTLILRAIGCDRVCANQKRRMADIVKRMEDGAFRTGKAWRILRKLNKSEMKNDAERWSTFLEL SNVLDPRAFNSRRGRALMNEAEQVISVQQLELAAVRGPVIVDKPEGENVADELWDIAEQIVEFIERKAAPVA EPQDTTNDPTFSFDVVLVTDLEDYLKVEARTAPSGVNGMSDRETHIIDEGIELSVTEQLSRMCSSLKEEYVIS SDILEKGFAELSAIERLDTVSIQVSFLLGMQMRKSCIIDHCESAKSEFERRIKRVFNQLAGKGQDMDHAASLI VLSAFCPGQVIRECIRGARTGVLHHDLSLKVLSQSSPLLEWREPSDGSGLTLELELQMTVLDISRNOSSFDRE SHNVVSFLLSLV GIDNSTSSIQSSSLMTVSKLITVCINPVWCCPDQSVEMQLNLLTLIQQLFHFFVATSADIDT DTLQISFNLAFRALSSANSNDTKV GALIREKVL LLLKSIMELMPDPVSTIRLEQFDSPLNAPLWTLISLFSDDL NEELCKGLEDVTA VEAYMQTSSVDEADPLPLPISIAIQALLWGLLWDSILSENVSVESTKAETWRLLDIIASFE FCGSESPDETIKGSSLIESAVAEMMLECGNVLF RALLCNIIPFLLEYELTENFQEEKLIPQW AIDKVS ENPEIKE QIPCRVSSHVIMRYVAKCWCLSGVANRQLNAQSDMLVLEALTHVVT AHDDQAITASKESL SGLTFCIQWLFCF LVSAARELHLDLSTWSTVRTQLELSFLRLNQL EHLKAISEAEAKFSQVFVAAWLGYPDDQFVQVFKYISTK RSN* |
| >scaffold_22_R9 74 | MILTTNLATWIFLTWAATSSMNL SKCQCTCCLNKCENPRPLRGHLLSSWVSGKQSGWPYQANLLHVHLLH PRQIRTVGSRK* |
| >scaffold_65_F3 81 | MRIPCSLLVVIALLSTTTNAISTEADAGRRSLRSMKAKVQDNQVEEERGGTGTSSLKAFVHDFDFNLLDNIFL PADFKRMTKEPEFLRHMMASWQMGFMSVDDIVLYMTRLNMSEKAINQFKLAYTAYLEYLKAVAKAEKAA |

| | |
|------------------------|--|
| | KLAKASLN* |
| >scaffold_25_F1 318 | MRLSFLAAAMATVYCATCNATADSDQNKVSMVQSRLNGQADGTRFLRTHHENEQESDREERDLTDFVET KKA AVK KLAKEVMADNRRRAKDVFLWKEKGYTLDELNTFLKSAKYQHVVYNQYMIFRGYV* |
| >scaffold_184_R 6 | MRLSYFVLVAAACIFACGEEVAAVKDSQGI AVSHTDSL RGDNRFLRGRQDVADEESDDEKNTNVEYEERAL PKLDVAAEKLMTRAAAKLTKSKLSNLAKMDD EAYHRAVVTNNFFTLERIEKMGYNPDKMFLKMKEMGKN APLTPYQKVLL EYYSMFWKIKYPSWVSEVPRMF* |
| >scaffold_1_F40 0 | MFVLFNLLSCTRLELSLFLPRPVAAAAIAAACADGG LGVTRHSPSCRLELSSVTASFVLLHQLRILRST* |
| >scaffold_6_R20 44 | MRVLLVALAVFASSVNTGVAGIQTKRSLRQYDFKSLTVADSKEEERNFVDKVDIARVTD DFVTKVKIPTN MNAAVEKVTTMADDIAAVAKTVAKTYPEGLSKGTLAQIKEVEQLRLKDIATYTKKTGDGMRRKISPFPGMKI APKYLESHVGRNMQLYGGDDGSRMLSSAVVSRSAKDG GVDLLISSSNPQKNDWLLPKGGWDKGEDIQS AALREVVEEGVCFILYM* |
| >scaffold_60_R8 42 | MFSLPTFVMFLVVPILVMAEDWSTEGISHSLPNARSLRQERVVDNSDVSDKAQRISGHVATVLGHSAPMD EVGQVVGMLLGV* |
| >scaffold_8_F32 65 | MPQTPRCLLLKVL FADVHCAALYDLLGHSNGSDLKYL MILGHPADLQLSGIQFVRLLRGQSPVALPVAHQL ST* |
| >scaffold_10_R1 425 | MRLSVILLVAAFAVALNPTAAASDANTVVAAPNVHES IATGRFLRARPVDDYPVKDERDEDDSDDEKDEER MFSFFQEKATALS AFKLVQS GDDLVEAVSGLSKGEFQALFNQ GKAHMAKMVPGFYPGMSLGEFGTVVR AAGLSDDMENALMVGYGKYLAHLMD* |
| >scaffold_81_F2 33 | MRFHALVLLSTALLATDAFKLTADVTAPQSRLLRKSEPKTNEERVFEGLTSTKLQYNLLWSKLQLGDDLSAV LKSPDVAKIAKFNLRAPGSQVSMIQRITAKYGGD DVARTLVSIEDASDNPILLTMVWQLREDQIANWLKN SETVPGVVSKLKLGTDESIFRSRALDVLEDFIKKYN TARNGDESLLKLT TIYGESELVKMIAKTRVLGRPPYSN HPASIEKANNIESQLIQKWKSNLPDFRVMNKLNFDD DVSMALSAGKVGVLKYSDSKTSAFRRLSAKYGEA EVAIAFAKAEGFLPEATALYRMQMNGWLSKGD TAGRVFSILKMKDTRDFVYKLDAL ETVYKFLKTKNPDDV TDVFKVLKKGFGAGENKLALAIVRPFEITGYHKS LFDWVWARDLPLSVAVKVKMSEADVAAAKYSDEL KPIIKQYTKFFKETVDLPEMPAVRVGRS* |
| >scaffold_6_R31 91 | MRLFIKNLVALAVLLATSTEATKIQTSGKVDVVQSS QILTGENKRLRSEHDEGELLGDDEEGLEDDEEER KYGANLFSAEKMKMLGDTDTEKKEEKSCQQ* |
| >scaffold_67_R4 06 | MWLLVTLVFGSIFTYADNSGSDSDNSIDDDY VESTYYYSSNAGLFGMLVAANTIMHLRCLRTAQLLVAE LVNAGVDRLQPPVVMLATTNPTS AV* |
| >scaffold_636_R 1 | MQRRMSTPFLLMITTIGDAQGEVEWVMIPADNS PLIKSDSFSRNCGAARRTRWATGVALPTSMSQTAPRA AGGRAFPPIRRCLRFQMICLNRSICSGVLP SASSIWMSSILTSSVVALTSGSMAI* |
| >scaffold_12_F1 261 | MGVVDWILCFTVCEEVGAACPCLTPSIVGRVLLHGKTKKIVSKPPTSNSQTMFCSFILDIWTTYTTMSKPIQ SDSDLDALRKLTRQLRVAKLVTDRLKQTD RKNATQAVMRKWLP LSTTVLKMVTRVLPSPIAAQVKRAERL CTISSEQLEQSPQHAQVFRSLQSCQTS EDAPLVIYICKVISVEANVLSDYHQSGLAATDEVYVGVGRVYSGVLR EGQPVYVMDPKFQGVSGDMDVDTIDPSTVKHVARID SGLIKTYMMMGRDLHLKDRVPAGNIVIGVGLQE HVLKTATLSSTLACPSLTKMPYQAKPIVRVAVEPED PRNFGALEAGLQRLYRSDPTVEHVQETGEHVIVALG ELHLERICDKLKERFAKAVQVSEPLVGFRESIVDGT ISSFQENIVFKELLNPDVTKDDSEKEDINASA EVQDT KVALGTTDPDGLTLKLRALPLETAKLLEESALL KRIAVSKKANDDMKDKNTVVELEAAASEDVS VFKKKLEK SLQSSSESSFLKALPLDQIWS CGPRRVGNMLINSIPTYRATGCLFPSDSVAETHDSEKDEKIRKLENSIVTGFQ MASSAGPLCDEPVWGVAFIIEDVVFHDEKSEEDK SEEKAEMSKYGPLSGQVISIMRTTCLMSFVKQPVRLV EAVYECTVQCQAEQLGKLYSVISKRRGDIYSEELSDG TALFTVKAHLPVVESEFGFATDLLIQTSGAASN PQLIFS HWSIEMDPFFQPQTEEEREDYGERVYEHYVRRYIEAVRKRKGLSRDEKVVVHAEKQRTLKR* |
| >scaffold_81_F9 0 | MRFPFLLLLLVTFLLVNGKADSQSTELRLLRDGNSPREEERALPPP KVS LFADWLKATGLKVSDKVRARYWL WRKQSAEDVFRLLKLDGGLEKLLGSRKFNTWTSFVNIYNKKNPNEKVTMDGILSKTYGDLELAKALEVAMGS MVKSERKMGTTLSLQREGWQAAGKTADDV FILLKLDKVGADLFTTPQLNSWYKYVSMQADSKSLMASV LRNHYSDETL SKIFREAKPEIKRMRIIRVQLET AVAKSKPKQLSPEEYFKMLKLDGDGVDKFLASNLD TWIKYV ARYNTKNPGQDVSTIKILT KFYGDEELAKVLEAGK MATTEKVATELQSAQFNQWLSLYSTDDVFRLLKLDN DLDSLITNPNLITWITYLQG FNANNPGKGTMMIKTFTK FYGEIPLAKMLESSLVPKTEKVATQFQAQFKQW LRDGKKPAEIWKALKMEKATWMQNPDAQVWYKYKDWYKLNKPQ* |
| >scaffold_6_R31 37 | MPLLILFIASILATTIYVNTKLLNILPGNSLGVSTY NHSHLDWKMDTAAFETSVGT SIHPTSSFYRRGKTFDRSA SRERAVMVCLHDAMLNMGSLLLRELRLCLGNHELIQVYHCGNELSDRSVELL FSLDNRVELVDVCSDSLSSRGII |

| | |
|------------------------|---|
| | SKEMATKFRSWWIKPLAMYHTDVRHVMLLDVDDVILKDPVAVVRTLDGYKNTGTTFFYDRVISNKRFLTGND TGEMYVHKLRLTFNYTRFNVSSEGFNPSQHMLNTFAFNGKSIHEMDSSMVLIDKQRAGKVVMDILLWFITEE RFRFTYSWGDKETFWLAFEMARVPYFFSPWGVSVVDSMPNEDLKNHPDTLCGSILQFMPVKGMNDTAEV LYMNGKALIDPYPQIGIFIRKAKQNNLNFNTIPTQMTTPQKRRQINTKAYPGKKFNTECLVGMGAVPLPHTFS PNLLRRRVHFLGLAMGVLGSLDHCETY* |
| >scaffold_2_R11 14 | MPVCYFMFLCVSYFASSSCQIKQHVEAFIGDLADAFVPEGGVLVQRRRNLTLQGLVARQVHTHELEVPAH VARYLRHEEVSRLDADLDAREGAMYDVRREHETELHELREQTREPHHEEQQLDVALWPLQ* |
| >scaffold_5_F94 7 | MRTTFLWLLALVLCVCAAEPKTPEPTSSANARDNDPVVQEIRGLRNSGMKLNDAKDFKGAIEKLRERAITLL HNRVFGEGRHAIIDPSEISQDAALYAQILNDYGSVLIRAKQYDEAIEVLEDSVTMVEKIYGDSSHPSLGLSLRSL ADAYMAKEEYKLAIKKYKTLRKHVKKGLEETHEAYIEASLRIAEGYKKGKLNKKNLKVLRDAVKAQGTEINGLT TGIAELYMELSTAHTAVGEIDDALRAETAASAIFLQRDGEETLSYAFSLNALAGVKMRQKKVDEAVKLEHAH KIAVKIYGENDPITQASAKTLKEVKEYRLDMHAQKDEL* |
| >scaffold_48_R1 407 | MRALCFIFSVLLLLLEIDASSVFNSGAAVTHPHLELTQVTTPTLRLLRSTHEEENDERAGASVLDGIVAKAMQLI NKNPEDVFKLKLANTNLQNNAVFEQWLQYVYKFRAAKGEDKFNHRLFNLLRKSQNHPPDLLVPLFQSLT HVQGMEDLARTMQLKLFESGYQSTRNLMNKAWLQGLDTPDDVFHILLEKNALESQQRLPQWLKFAEMY KTQNKISSWENELSLLLKTPHEKETQFGLLFQSLKKTGEMETIAAKMEAQLFTRWIKTDTMTDPKVGCVLAS PTNTNWKRIFEPLVTEPRHVLESYTVAYAASRGGKVLKSVEKLFANNQPVAALERAIV* |
| >scaffold_59_F7 11 | MRLTYIMIVATAAILACCDGASAVSDSKRLTVPLDRVQTGNANRFLRVHHEEEEEERTGPQFIKLSLEK FAKAAETGNFAKTVEKLRKSLNGISKVDDVVVPGRAGKTLTEVLKIDDVAYLKAAGKWPKNLDDIQEAKLV REISTLDDQATLKLITKENAAEFKIEDMGFNPDMKLRMKQLSKNDKDYFDLVLLKHYYQYMAKHPTWA SSL* |
| >scaffold_29_F1 645 | MRVFSILLVAAATLVASASAESESKQRDLVSPAAPQWRTIAENEVPTKRNLRKKEIEEERAITAISLFDVVKA KGLQALPYPELANLDSKLRQYKLLVNNLDRKQIVEITGQVPHYLTHGDKARRIVQYNKWWRKEPVPDP WVLKNYPAFFKGYEEFFNNRFRTRGYKYA* |
| >scaffold_2_R17 01 | MRFSHFVFIAAAILLSSENAIADTPVEGQALMTETDAETPVRALSSNNDKRFRLRSYKEEEAYLTEDKYDEEKKR EES* |
| >scaffold_29_R9 98 | MRLLYIAVVAILASTNVYPAAADA EVSSFTTERPHWDRSLTAATEGDNDGKRFRLRKRKTTTNETIRERSKQ GSENTVSYCTACLHYP* |
| >scaffold_9_F34 50 | MRFSHVLLLATAALISSCNATAIVPGEGQVTTSDAVVPVRALETRNGKRSRYYDAEEEDETKYDQKNGK YDDDEEERVMTAAQIAKW TAKAGEWVEQGKTPAYIKDKLTA FNQVM SDKNKEYRFLATWGRAHPNEL GRRYLRPEAESSPRFRKLIVFISNHLLDVLFCVTR* |
| >scaffold_6_F30 32 | MSAFLVAFRCCHVVFAFWELRSVEFALALGLNRARGSRWLRRLVAHLLSLYEAVLDGPGSASTSVAFMPIV* |
| >scaffold_32_F2 41 | MSRAGSPVALVPPKTVLIVVNLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCEPVFV AKSEEALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWVVFHNVDVYNSAELQDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSYCSVNRTVARRVVEHYHGDGLVWLHDFHFLMLPSYL LRLRLTALVTMYLHVFPSSIEFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVDPVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDRPNDYHRTRDNVLFTEEINRRYAPPGVVVVYFEERKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDMDV GALAKACGMSYIEKHNRFYMNKVFVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFDFRVMRFESGFVCLDDELVKKCANTSRRFLIFDYGGTSSANILDEEGARFSHRPG CVEQSAENADSSDRSGDCTGRSAARYVDGKVRKPISETRASLRTLCDVPCNIVFVTSNTQRGALEDQFDSI PNLNLIAENGLFIRMGGQTRWECVCEEDKRSFGWKDDVKRVMEAYAARTNGSFLVENAASLLYDYNSSDQ EYGEIQSLELCAQLRKIVEVCIKVILSSR* |
| >scaffold_399_R 2 | MRLTCILLVAAASLVGVLDA SAATTGNTVVANAAMVISPLAPESQGRRLRLVYDDEDDSADEKDEEEDSA DKVDEERGWLSDKMALTS LASKFVGKSTDEMGEVIKLT PAQIDTIFDKGEDSIQKMLPGFYSGMDFQKFD DLIRALPQEQQAVMLSAYTKYLHQNGRFS* |
| >scaffold_7_F17 75 | MRLVNVWSMLGLVLVLMVTEAFETTKGAKNGVTPDARRLLGFAADQEERRFGGPATGTDHSNSHAW ENFKAWFKETFFFWRKWEQKRRRLRS* |
| >scaffold_18_F2 220 | MQRRMSTPFFLMITTIGDAQGEVWVMI PADNSPLIKSDNFSRNCRATRRTRWATGVALPTSMSQTAPRA AGGRAFPSPIRRCLRFQMICLNRSICSGVLPSASSIWCISILTSSVVALTSGSMAI* |

| | |
|------------------------|---|
| >scaffold_50_F2 79 | MRFLVFLALAIVALVADGEAKDSRLLRARSDEERGGGITVPVADKVAKLFQFSKLTDEQLQQWLSAGKTAES VFYRMGLDNTFVTKVFKSPQFIRWLQYADDLSASGKGTSAISILSTKYGDEQLYKMGWAKKESSTEALGLRL QSEQLDHWVVKVGDPEVFKLYDLNYAGSRILSTSQFSAWTKYVDDLNAKNEGAFVSIPTLRKHFSDDDEL HIVLAAKRSGETEAMGMKLEDAFVQFWIHRKETPGNVLVGLGLKSMETLLESPLLSILTKEYEVYVNVYAAK KTTVIETLRTFGDDKVARMLLAGRTESTTTKIAKQFQTDQLEMWLLSSGQSVDDVYKLLKLPYRDLVADIGTN KFLGTWLTFMNAVSLKNPEKMSAIFTRLEPTFGNRPMMQILEAANKLPSMEKAATKLQLEKAQRIFSTGESP YQLFKLVALDDVGESVLSPLFKKWMLYVEDFNKNPSDEVSWFVALRAGYQDDKLGPIIDKAMKDPKTMK LAKLVEKERMKEWLEKWKRPPSMAFRELHLNKAGEKVFSAKPFELWVKYLDLWVQAYPSKTTMIDAFVD NYHMSTLVPMLAAAEKVPITKKLASQLKDALVDKWIAEKESLAYVKS WLKVPSSDDMLERFTKKLNSA* |
| >scaffold_7_R19 03 | MSVLCLALMLAMVTGAFETSRDVKNGVVTSSRLLGFTADQEERRFGGPATGTDHNSHTWENFKAWF KKTFFFWRKREQTRRLRN* |
| >scaffold_24_F3 57 | MRLSFLLLVAAVALLSSGTVVSAMTNDPEISTPNQVLSSETTEGRKLRMHNIASDDTEERGFNAQKFEQLMD DGTYSKRFRANWVTKRYTDTDIYNKLQISSNPYKRIKLNKYQTYIEHFAPGLISP* |
| >scaffold_17_F3 59 | MAANSCVAVLVLCGLPGAGKTLVKQLVATACSISSRLHERISFDDLYEQHVTAEGKPGFEFDPKWKMCQ QDMLKRVSNRLKEQNDPVHRNECNQLVLLVDDNFQYRSLRKRFFHLTAKRRRPSGFTLCTTTGMPLMTRSE LWFWCALCGCSFRYLSGTECWSKQTRTSTQ* |
| >scaffold_8_R31 93 | MRLSFLLPVAMAAIYCATCNATADSDQNKMSTVHSLDARLNDEAGGRRFLRVNQEEDVDAEERGFYFEKA AVKKMAKAIMADPSKADEVYNTWAEKGYTSLKMSEYLKAKKYDQVYNGYALHLDI* |
| >scaffold_83_F3 73 | MHPPSLVLTVLSPAALKSSTLHVALELQDTPVSSHASEPFSFPRCRRPLRVASSTCAKAGADPLLAPNGH LK* |
| >scaffold_25_F1 083 | MRLSFLLPVAMVAIYCATCNANVDSNQNKASMLQARLNDEAGGTRLLRVHHESDTEERGFLEKAAVKKMA KAIMADPNKADEVYKWKWADKGYTLTQMSNFLKSKTAGKYDRVYNGYVIHLDY* |
| >scaffold_22_F1 823 | MRLTFLFAAMLAASFSTSDAASVDQTTGVRSLRRYQAEDEERGVTSVSTSIDDVLNNLIKQKSSLSKLISQK LDNMLTKNFIKLLRSNKSYTDEVFTKATLNKMLTSEKFAEKKFVEWYALGLTDKILQLRNLNGVGEHFGLTHSQ YVTFINRIHGVA* |
| >scaffold_157_F 14 | MRVASIALLAVVTALASVTDSSAATTGTVLAKVVSNEAAPSVEHEHATRFLRKHKDHADTEREERNGISLQ GLKSTFEKVADLPFDRAWHHLQMLNLSWDKREALLKHLRLSAKDREAVLKIT* |
| >scaffold_7_F23 10 | MISRFTLITAGVMASSTLIHASPLQYDPYTPVNISTPLTSSHPAYGAQTEGCIPIVPEDPNQAKAESMIIQAD IYRKLRSMEDTTNSDIQDLETYFGTKMEVNFQTLKQQYSSGHAPATPWASSYWPWFQDSINYVWKTGEP SA SEKYATAYGLNVTEFKDKISERSGVSRRRSTRCTADSDCKDGSVCGKRDGISSGYCIPGWFGICHAWAPAAI LEPEPQCDVTKNNVTFHVM DIKGLVTSIYDGAIEKTVFTGARFSGLDSPANKDQYGRFTDAARRDLGPGYFH IAITNVMAKQNRSFVVDVTAGSEVWNQPVRSFNVQSMDLVDTRVASMQYFGVPSYPFNDK MVRLAYVKT TFSWVSESYKDGPLVSSGRIDRYTESKDYELLELDADYNIIGGEVWVQSKDEHPDFWLPTAKPSASAVTST GLKYADVQELLNLSQSC* |
| >scaffold_39_R8 34 | MPVLPRLSLAGSSSCGAQVLSLLALLDASSAQYASLPSFDELFPYLYLLLHALVKLEDTKVSEVNAVISKLH NRLETCWNARRPLRLQTFAPTILPTFAPQFDENYTVRKDKTAPKDTAQLKQLQRQVKRARKGAARELRRDA EFIHREKQKEEEARLSAKEEKQKEIRRWLEEQNATFNQQVRKGGHMLKGGGSARGPAPRARTPRK* |
| >scaffold_104_R 20 | MRLTSILAAVVVTLHTSATAFPAVKDTAAIENGAVADIVDSSVTGGRMLRVNYNDDDDLDLDDKKKTRN SEDPLDNYDEEERVGIVEALKKLNVPVTAANKS AKKLQSTLRRSRK* |
| >scaffold_61_R5 21 | MKVTKVVVALAALCVLWTSPTDSEDISNILEVSASRHLRQTSAEFAAKPQETGKKRDSTNPLQRRDQALVS AHRVYDPVSGLACSLVGEVACVQSERDESFCRETGYRQELDCPRPNDPKDEALLTKPEDERETRFKACSPA DSARPGVAVVKFELLMAAVLAASVLLRRERRNHMSSFDLRKDPRQRTGLLGGNSDKSSD* |
| >scaffold_90_R9 4 | MANLYKLFVTLVLVSWGSKVSVCCLTSLRFLRTQNLKTRYLAKSTSCFQC GSGHHLPLRHYAVSVSQLASLN LRLACC* |
| >scaffold_25_R3 32 | MYRVLLLVTFALLCGFSNAANSGRLLRVVDSAEERGGFTSYLMDSF TKWRINSKINSWVNKQKTDEYVLAK LGLSTLTGKELVKAAYPQFQDFKVGVLKEATPTTSVSTLGLDKVEGAVEKADDFGTYYKVMALGEKAD DYPITRWRELFGGGSPEQLKLRQLLFLAKRNAIDIRIMLG* |
| >scaffold_18_F2 130 | MRPNLVVLA AVIALVSRCTAVSADSSTNLNQALDTLQNHAGDGNRFLRSANIVDDRVDIDSDDAIDNDDE ERGEKLGRLNLPNGMRAERPQTSTKGS HSSLTYAKRTNMAKSDGSITTSTIASGPITWPS* |
| >scaffold_74_F6 12 | MVFIAPKLTAEQALVIISCLLAARGNRKRFLRTMHFLIASHASVFETHLLGSTQATASTTPLTTRRHGHASILR YLSYDC* |
| >scaffold_37_F1 | MGLLQKVKTMMIPTVALSLTATRIGVPSVLAARAVPTQTHQPLLRRAKLIRELRQGINVSRLLVAKSPRIRL |

| | |
|------------------------|--|
| 084 | QGHLRAVVRYRTRRMIL* |
| >scaffold_11_F1 319 | MRIGYVTLTATTAILASFGNVSGDSDLTQVVKVASLDAVLPNTNRLRGRKNNEEMEERNGFEALAAKL DKSVLPAILKVANLDLGRAALTQLRIPFEQLAIQALLRLSKDRKAVLLLIK* |
| >scaffold_11_R2 801 | MKWCRASSASLALLACSLAGLQDQAADRWTVHTVQRTRSAEELVHEGTSRWPGDAGAVASVTCPSDLLL ALQCREADGQRELRRDDQDLRRLTRMCMCCADHRRIKSGGLFALYMKASVRLTANCDRYELDAPLVSTLTC RSVDKDIHLLPDKEPL* |
| >scaffold_11_F2 657 | MQRMMSTPFFLMITTIGDAQGEVEVVMIPADNSPLIKSDNFSRNCGATRRTRWATGVALPTSMSQTAPRA AGGRAFPFPIRRCLRFQMICLNRSICSGVLPASSIWMSSILTSSVVALTSGSMAI* |
| >scaffold_50_R1 378 | MGLFYLVCAILAFMAGDSTGDRVIVSDGFNQHRENAARASVSTTRLLRTKSVIDEERVGGIPVSATDKLAK FLKPSKVTDKQLQEWLRNGKTAESVFYRMLNLPSTKYLFEDLQFTRWLKYADDLSASGKGASAVISLSAKY GDEILYLMIDRAMQEQSALGIRLQADQLAHWVKVRKDPDEVFKLYDLNYAGRGILSNQFNAWTKYVDD LSAKNEGAFVSIIPTRKYYSDDNLIKIALAAKEVDETEAMGMKLEDAFVQFVIHRKETPDNVLDLGLKSTK TLLKNPLNLTKEYTEAYNVNYPMSRMTTVIETLRTFSDEVMAKTFLAGRTEYTTKIAKQFQTDQLEMWLSGG QSVDDVYKLLSLPPRNSLDFGNQKLFDTWLTFMNAVSIKNPKDTSIAFTTLAPTFNDRPMMQILEAAKFP SMEKAATKLQLEKASIFSTGVSPYTAFRMVALDDVGESVSSPLFKKWMMLRPDRYHLMASHLCCRPAAQ HFLKRPRRRDVCGGNSCVLPRYVVRGRGVAGVALLCRKPNEQLLLLSTTDC* |
| >scaffold_17_F2 587 | MIVYTGTLVTTLVAFVATVCLETEHTACVLSNDLGISSSTAQVLGLSLWGSVHLEQVEQSPFSRQLHGHVVA LDQLEDGRYKGHQVFARLLRGALSRLNHAQRATDHVLLVLRHTLHQYWD* |
| >scaffold_25_F1 282 | MRLSFLPAAMAAYCATCNATSDQNKMSMVQSLDARLNGQADGTRFLRAHHESEESDREERGFTDLFK NEKAAVKKMAKAIMADPSKADEVYNTWAACKYTLTQLSNFLKSKTAGKYDRVYNGYALHLDY* |
| >scaffold_68_R3 1 | MRLNLAVLAAVIALVSRCTAVSAASSTNQJNLSNLNQAINTMPNHAGDGNRLRTAKIVDDSEDDIDSDDDI DRDEEERGGKTWAEKFAKWHARGESADDVYQRFALPEVVRQAYKYGQIGRLDDNEYRKYWAAYSA |
| >scaffold_11_F1 227 | MKLSMLVLALVCISQLGGSSANEATDIMRRQLRVGKAVASLFENQHQSTRELEENIMQDEDNKPNEVQAEF TKFRMRRLRSYVELIE* |
| >scaffold_117_F 62 | MLRSVGGAHFDVHIVSLLFLLHGFSTHTLELLVANVQSQGIETSECEVDDQNGNTEPNVARALIDREALEL SFALSISVSVTRELHQHIGT* |
| >scaffold_53_F9 79 | MRLSVILLAVFALSSSVSATRNHPGETATANTAMQGVDMRTDTNQMRFLRTEADDDEERLAGKNMNFN AEKIEKALQDTSYAKTLFRRWKRYEVEHGAADFCLKIFNIGKDDKVFGLYKSYVSWLEKHHPLGAETGGGPNL FSKAKLKDAMKDPKYENTMFGRWKRQGFESDAAYNKLLAFNLASDADVYKIYKYVTWLNIIHPLAKTRKT TAKDFLFNVDRIRAKKDSEFAETLFKWKTSGLDEKPVYKLWDMGLKTDELYKLYKNYVKWLDIHYPLP AKAT* |
| >scaffold_22_F2 097 | MGGLRRPALLWIPHFLVADANCNAAVEIDFKSGRILRDIDYAVISDEFARVAISRSTFAVVSVPKVFNRNDEE GTGIVDRVRLMLHPTDLSSSRFQDVKLSFTLVRDVTNYCREAEVRLRQAREIDGQLPCLTMHSFQLNIRE NFSLNLHISEGHRVFRWPPPTAEVLLTHVEIPLDNLQESVTYPAFMEIPIHAVVGKTPAQKRTAVNSADTK TPNTEQLFQRDWAVVPIRLDSKAELDVAPPSPSIERTSTYLVDLKTANEGDPTRMTVEQKEFTPYFYV EMAFSPTFWCRYDQTWVFDKTKKVLDMYQVVHRGFDTKVMISTSAYAGCVRVARCSIDCLENVHLF FCHHPYISKAIICLVKISQPTLLKWMMPRAADCITYLKSFKLIIPS* |
| >scaffold_13_R9 5 | MRLFWTLLVALVALSSCDAASVDENKALQRKLYTKVASQALAADNRHGKRALGESSKIAPSSVTEARAAT VSTGFSGKIMAFRAIKEKYLKWEQKILAPSFKEKAEGTTYSEVLANYRTRLNWSGLWGTPSGFKRYARLYE TWLKNPNKYSHLAV* |
| >scaffold_73_R3 02 | MSSRGHFFSTAFPILAACKFQVSSSPFPHVALPFGSFLMSIANTFHAPGSLQYLVASLVSAGVMSSGTATSG KSGSDASMRWLSNMMRMLRRLRASLTITSRTSR* |
| >scaffold_25_F1 407 | MRLSFLSVAMAVIYCATCNATVDSQNKVSMVQSLDARLNGQADGTRLLRTHHENEQESDREERGLTDL FKTEKAAVKKMAKAIMADPSKADEVYMKWEAKKYTLTQMSNFLKSKTKGKYDQVYNGYVIHLDH* |
| >scaffold_60_F3 66 | MLIHATTICHVVWLVLQLANGFATFVTMEMKTLTITRLLRRATAKETRNTCTWIASAGITIWMRTERTRRR CVPSPIQMAWTCASARRPT* |
| >scaffold_5_R26 54 | MRPYFLLLAVLLAISNQLRPADQSIGPLKTRRSLRLAKKSDTSNEKEDERLGLSSLKTEIKKVPLKVKLAWWQ HFVGGPSEYVKKLGANHPLYLYRHTREGLEMWSWLAEAIRHTKSGKGLMAWSHCMNLDQIARQIKK PEGTEPFRVYKRYAKEFDGHRLLSSDITYFIDESASVAGRYARAHWAESKVDKEYVLEFLGLLYMKPIYVKHNPY YQYCSASSVLVAMVIKVFVEVHGRVEA* |
| >scaffold_13_F1 424 | MLGLPLAILELLHCHVVLAHMSGLTSQNILVSNFTFCKKQKLHVCSRLRLPDTNSRGGRIIPRTARAQVCAG ADQEQGEADPEPTGADPSTYQQAETLGCGEGLTF* |

| | |
|------------------------|--|
| >scaffold_7_F13 78 | MSVLCLALMLAMATGAFESTGDAKNGVVTSSRLLGFTADQEERKFGGPATGTDVNSHAWENFKAWF KKTFFFWRKREQTRRLRI* |
| >scaffold_24_R3 31 | MVLFSFCLVYIILCCYAEKIRPLTLGQDRRLRCAVIAFPVHEKQRSEDEVDITIPKVFLAKGSSVEMLWHFCAIS ADRRTLFLQRGSLRLRLISRPFLECAWGIPTFLDVVPAAPARWRRRTSRAQNRRAIRPREVAVRLDQGG QIGFKEKIKPSPVARTRAKGNDEKECVLHDIRDKFHQLLHVLR* |
| >scaffold_11_F9 7 | MPTSRPCSWLHPAALCPVVFPMMLPLLDLDCREFSNGPFNLAYVIAVGSSEPTPIPTTARMCRSGQTPFCP KFRLGCPFFIFSLNCRGRHPHHRALFHRCLRHFLHYPRRHCPRRWQCPRSVRTS* |
| >scaffold_6_R29 10 | MVSFFLPFFLFLAFVFFTFSCSSPHHDNEPPSRTDRSTPRSSGQPDLHPLRRLRGKSGSGTARRCILPGQTPSD ASKDVPLRPPPHAS* |
| >scaffold_25_R1 205 | MRITYILAVTVAATLHSSVTAIPSVKSSKATENGAVPAVIDSTHTGTGRMLRWVNKYEGDLDKYEGDLDDN DDLDDDDLEEERGFSDTLKKNPLKLVKKGTKLTAEQAAKVQALKDAADYQKMIENANKLIRSD* |
| >scaffold_12_F7 09 | MRASKLLMLTVFLLASLDATSGYNKLGQNTTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLK KSTSGATNWLKDKKLAALKLRALPMNLDSVQKTMKDGIDPDRVFTLLNLQKKS |
| >scaffold_12_F2 959 | MRLISIVLAVATIFACCGAPTVDSDKSIQVPHHVQTENNANRLLRVHQDEEERTGAQLVKSLEKFSKKA AEKLTRSKSFNDFVKLDDVAYKENFHSATQALYQRIEKMGNPDGMLKTMKRKGDVLDLLAEYTRYWMK KYPTWTKNQ* |
| >scaffold_5_F36 83 | MFCGVLIVALVQSLFFNFLDLSPEKRVRYLIELEWWEKATRQNAAKLLQTAWRSGNLRRTDIDGQCHLFS LMRTARRLRIDKPAIELSVEDQVAEMEATILAEVDRMEAQKVKVLRQIQAQATQLATLKHKLQMKKKK* |
| >scaffold_495_R 1 | MRLSLLVAAAVLVANSDAAPQSTSLTKFSTDVAPVRSRSGASKTKTEVDEDDSDPDEEEERGIAMYGLDK VDDILKQLNLADDSVIKLLAKGKGGVEQLITTDKGVKTLVITSKKTGKKVFTFNIDIENIRKIESTPAIK |
| >scaffold_536_R 3 | MVLMLVSAVCMCSCLLYCAAQRLVNPQQLPTGVMVCRHLRWNHWSRRDALTALQASDTLQDATYAKGA SACEVTKLCELVSSCRRRGDAGSRLCPA* |
| >scaffold_5_F17 32 | MALGVAMAITLGLLSLVSNNTAVGTENAVGPVTRHLRSLLDQKVSEHGAALDQQPLAASMEATSRRSVPQ NPGGPLGGGRTMEAIAPRYEKKAAPETIAEAVAKAEAEIEEALIVALGAENDLQAEIAITEASILSLATNSIT EGLGGEEIVVGEIEDSDSDSSDQEEHTTRTKAKKHKDW* |
| >scaffold_1_F44 48 | MNCSQQSRLLFSLAFCQHSRLIFALVFSQQSRLQCVYDDRILPDEHLRAIRRDVSPSLYSRDLRRPSNAP* |
| >scaffold_53_F9 2 | MRLGYFLLATIVGFLACDNATASVSESTSKLTAREEHPIHGRIGDFTAGHDNKRALRSEDEDDGADDSDEE RDLILSTIHRPKYWRWFKAGMTPYAVQVGLTGVRRLLWPKFKRREYKGYVVFYTEQCHKPEYHDFCKKHA DP* |
| >scaffold_12_R4 2 | MRLTSIVLVAAVSIFVCCQALADSDSKSISVPQHVVHTENANRLLRVHQDEEERAGPQLVKSLEKFSKKA AEKLTRSKSFNDFKLLDDVAYKENFHSATQALYQRIEKMGNPDGMLKTMKRKGDVLDLLAEYTRYWMR KYPTWTKNQ* |
| >scaffold_3_R28 40 | MMALLLVGAPWLLPLMLPLPRFDLAPDVTLTPRERALRFSSLRILPSLISSLANLFTLLRLPMTAPPRGFLPLR * |
| >scaffold_68_R5 30 | MKTIIASLLTAVAVNAANGDVNALSIASTISDAKQVSIRALRAAHGSHEHSSGMGSMEDSHDTTSSSTHESTV AGDDDDDDHSDHDTSSKSSMAGTAGSAGPSNSTESTQAPDTSSAASITVAAGSIFLAAAAAFL* |
| >scaffold_24_F1 366 | MRLLLVVALTLAAFLAAADVSALNDAPSKRLLRSTVRVDEEEERGMWETLSSKVTKLIKPNQVAIKAMDDPKI AEVAGTSLTSLKSVNPKKFDSVDGLFSSKAFNNLENYVLRNLNKQDINKQTSVAKVFSTGLGDKQAFHLFFTA TQSSDSAVEKSGRFFRDQLLTQWATEGKTWTEVSKSVKGLPATYYPRENKYFDILFNLAHDTQKRAARLA RLEKARMAANTAA* |
| >scaffold_77_R3 94 | MRLTYIALVIAATLHASGTAISTDKSVKIPAIADDAGRMLRIVKEKPVSDKEAEREERFANPIKKGAFLLKKK WDKQTLKEAIKRDENRRKWIREQGFEPD* |
| >scaffold_5_R20 89 | MRLSFVLPVAVIATFASSGNAVATADGRNTGLSAITPPNVVASIDTAVGGEKRSRHYHNNKDLEDDSDDEG LEDAEEERRGNMFSATKLEMDLGDGKQVMSRFKKWKAFGYNTYNLPDAIQARKYDELKMYRKFLYYN* |
| >scaffold_12_R9 67 | MRLQFAVLFALSIVITAANGFSETTAQQFNELSTGEKQHDEKRILRTEKVEDEDEEGTEGEERVQVSPVSWII DLFTPKTAEQIAEAAKAEAVKFYTKLANSPSFRAPERFNWKIDGMQVESVLVHLKLWGLDGEKFKAIATKYT EFLASGKLS* |
| >scaffold_535_F 2 | MRLTCILLVAAASLVGVLDASAATTGNTVVANAAMVISPLAPESQGRSLRLVYDDEDDSADEKDEEEESA DKVDEERGWLSDKMALTSVSKFVKGSTDEMGEVIKLTTPAQIDTIFDKGEDSIQKMLPGFYSGMDFQKFD DLIRALPQEQQAVMLSAYTKYLHQNGRFS* |
| >scaffold_29_F1 | MRVFSIFLLVAAATLVASASTESESKQRLDNSPAAPQWRTITENEVPTKRNLKRRKKIEERATTAISLFDDAVK |

| | |
|------------------------|---|
| 837 | AKGWQVLPYAELANLDATVRTQYLKLLVNNLERKQIVELTGQVPRYVLKHGDSKATRLVQYNKWIFNHFKE AVDPAWVLKNYPAFFKGYDKFYQNRFRGYKYA* |
| >scaffold_11_F1 676 | MRLPHFLLVATATFLSSYNSTATVSTEGQMMPADAAVPRALEANDGKRFLRYKKEDMYDENDGDDDDV DDDDKYDEEVEERGVMATQVAKWTDKANKWVRLKETLVTIKEKLTNMKGVISAKNREKYNLFTAVYGRA NPHVFERL* |
| >scaffold_31_F1 184 | MRFGLFLALLVATFVACSTVANAESVALTLDNEVRRRLRNQQNIAKAAGDFISKSSESATLTKAINIAKTANG DEAAARRAVMLAAGAKEGAKLSDETMVKLSAMIAESAKKNPKSWPRLKFKVKITLGIQVGLAIYGAYKLLF DKGSSTAAATTTTTTSSGAA* |
| >scaffold_40_R1 072 | MAFSSFLKFLAMATLLMVQVNAEPSQQRNLRISDLPQEQARRLGNWALVINVKNPVGSDGVMK LLKADLITGTSSKIKIEKVKDAAEKVKNSAGVKDKVTGN* |
| >scaffold_62_R1 2 | MRVTQTLFAIAVVLFAATDNLTTASKATTSSKVQKDGLYAEIYNVEDGVVTKMRVPLGDVEQYSDTDFEQL KSTLTTEERAAVHLPFGVDSFFKGLEKVFVGFVKGRRLRIEEA* |
| >scaffold_4_R32 84 | MVKCAFVVGLVMALASGENNVVDVGDVGGNGVKTEAVVRVNAVESLRFVFLDAIINRSVVLEHSRETRD LRVLSIEREGVVEVEVDAATVSSCNATTACPPENVNDEIDENSVPPLSQQTDRLVLDGENVSTLSFQD VIDSAAGSVAVLPLFKPIGSAHVLSGFARVEILSPKLEFPAIPEREQVVMAWIEEKARLKAEREAREIANN KELQERLEKERLLAEALAKKEEELLAKLDREYERTRMTPHNLAAGKRRDGFWEFVEFEKTKGPIGLNWDL NTRDKAVVSHLEPELPAQQLNVIAPRDQIISLNGVDTSKMGPPQEVVEVYLSAIPKRMVFLVQMSAERAAK SAEKNPVKRVVMNWTAFDAPEVLRGWEVRLHLASWSVPPQINETNASLPLQFELPTPITGCSFVPVQSS NETAGVVYLAYRGACTLVEKAKNARTANGSALLIVNANGEGRFTPSGTVVVERVDVPTLYESTFVFIRCIW NLICGVCTAG* |
| >scaffold_211_R 6 | MRLSQVLVIAVASFVFASDTVATSNQAKISKTVQSSQSQRLLRSNHYPVKEEEDSESDVDFEERGFTTPDEE DLEERSPLSAAIVEKLDDIASRWGTSWAAMGQSSISEDKIKALLALRDAYLSGNKNAKAAAKLAILRANWT RSQQKW* |
| >scaffold_80_F4 73 | MSFSLFCFTQLLWSCNFQTVKAVFEAYAQTYVRIPPEGTVVFSTALPTCTWKSTDSVFSFITRILRF* |
| >scaffold_62_R7 23 | MIGCKSLISRWPTFFLTLAADSIWSDFFVACNPQLTDQARHLNSRERKRYLREYPGIARHFHRRFKAFFFL QHLLWKRSSSWGNCRLLLAR* |
| >scaffold_1_F54 50 | MKTATTFATVLALIVATNAAQVSPHTPALRGLRLTADTPSVEDDKEDRKHDDHHVKKVKKIAIPVVPVVEVPQF IPVVPVSPSTVVASSNNAVGPSTNVAGPGAAAPGAPGAVTPAPTTLNRPAAATPAATSTRSRPTPA PTNFAGARPSGAQLPAAPVQVGAPGMTGFPNSAGVDNTNNAFAGIGAGGFPMTGNLAGGFGFR NPMNGFGGANGMGGGIFGGAGAGPAMGGNGFGGNGLGGFGGQAAMNFGQAGNNGFEGGQGGF GSGFNRRERHRRR* |
| >scaffold_26_R1 820 | MAAIKISWQPGCFPPVALSYLVVWSELCAANRRMVWIRRSRFRGKTRLRSPYLTEHKLCMFHVFMHFFYL N* |
| >scaffold_52_R8 | MSALRAQIYLFVVARAAVDGQVQVCLVAVQQDLVRARAIRLATTNHVHQRLISTRPPQSRHQARRRESAH TLQQVQGLFRALRVRHGVQKVLERHHNHERRAANRRFALLRVVYFYGILVQNDGRILLDR* |
| >scaffold_33_F2 96 | MGVWFCAIILLIMASMSIPDPYGDPAHMSIDEEDWTDQIDSINADAPNSAGKYVPMMLASFGLMCE LQQMPWWSSTHNENRSNNVAFAKPPFTPSKRRSLRSELLSRSS* |
| >scaffold_16_F1 464 | MRFSQLVIAAVSLLFANDTVAVATSNHAAISKTVQSSQSQRLLRSNKYLVDEEEDQSESDVDFEERGGGKS HPLTNRKLRSLKQSLVGEELLIRELLWVSQASRPRLMRCLPCETRTCFATGLLSRQQR* |
| >scaffold_2_R38 99 | MRLQSIVLIVATLVATTQATTKLNAPSSETVDISPQRFLRKHHTQNDEERGAASLIEKAKRVFPKITDKTLQR WANKKSPKQALTRLKLDIPGRTSLRSLSSASGPLSWPSATRTPPKRWSRLYWRNTAMRP* |
| >scaffold_28_F9 19 | MDTCLSVLLSLVVSACSSQALPEVLNASVEMASVTVSLYGVGTPSLISRWLRAIAEFYDYPISQLLDDHFCGL WHRFISASHRPVPQEEENSTGWMKNSQLAPLPNGHSLQQFPLCILLGEEPMNTQETHFAKMLDKIVPIG VLHSFISGDMSENGDRFKFVDEILSCFPVDDMNAERKIDFGAQLTDLFAFSFMLLVYDPDLKQLAQQM VDIAEERAHGNTLQLSHLGHIAASKMARFTVWNIQGCDEDLVTQSDLWRDALKMMKEKYSAFDWKLLNIA DLLGEFVYLLRTEHIDPRAVCAVECFKIFVEETRDVAESSLVQLLQSICFQSIKRLAVRNRNRIGRVLVLLVRE NCEYFMKSTDKFGKYLGFVVQEISDILSKCNASSQTRNSVSGALSVSADDQAELEWVIFAVCNELGSGLGKH ALDIDMVDGISSSLDKLNALISSRKVVSSETSQSEERKKNLSQSAMGTRNQAIQILMFIQREQSRGAQFY NPHPFSGTASAISETYVNAATQSPAGSLADVRLSAVTTCIDMVRDSSQSTMKLYGKLAQTLVYSSSGTFTGT SRYNCEAASLANTLGQLGALHASEYVSPAEEGELSRLYWRHFHREGALREIKTTFGLVMHENVLYLSSL LFEGTRFGHVDPTVVEETLKTQLTVLNLEEGLAALARSKDNELKAFKPFESSSPSNWSSSYSSSGGDWGVKP |

| | |
|------------------------|---|
| | KRTFRQFLKHWTSAEELGFEAWVRSCLAARESSDPVLKACALSAMRVDMAVFLFPYALERILRLDNRDV DNPDEEKSSGVSEPSRIMKAANQGIRFVLGTGSDALSHHVLASLDGTQRESEFSQPPEAVQLVVHSINFLR ETEKAQFVETNGRGPQTVSTIKGKGSRRSSTYTAASIGRQHLNDLAYGCLVDVDFLAVAKAAVRVKMPYSA MQYVEMWLEKKQGGKITSLSLDRDGMVDTVRDILVEAYSFDSDDDGIYGVNDGRTVKSQLVKYNREGLH ARALPLYDVSQFSSQQLVSTDELTLNTPRRLVEGILTSLSLQSLGYNHLLTGYLQSLQSGDVAGNKSQTIAIQA EHKYKLAWKSMQWEAVLSGLSASGEHSSHQMFQGLRAIAHGNFTRLQGITTAKAEQVLRVSLHSFE STKDSYSALVRLQAIHEIEELANHIRNSVPPEPLVFSATTTGSGPFLPSLGGTLAAPRQETLTVLPLEQWHQR RDQIKNDFDKAESLLALEEVLVQVAKPSDNARVVTKLYDLASLSRKAGRIAIAYRALQKLEHLDERGSLGIYER MQCQIQKAKLLWKQEQEARSIAWTGKSVSSELTGYLRDTSISATEVTSLQLLLVKLTFTGKWIAFQRSESSQVI LEDFFQKATEIMSNMDPEAVSERSRDAAKAHFALAEFMAGMYQQVSTRVTSQEWLTGKMMVVQARHDEL QELQSMEQNMQENRAHIFALNKEVIYDMNERSKVEASVDQFLIGAICSYGKGLTSLQAEIDMVFRVLSL WFNNQHKPDINRVVIEVIDMVPSYKVVPLSYQIISRISASGTFQALRKLVMKLEQHPHHTLIQIALKNS GDVEGKALQFRTNVGDAAKAEKAVYLTLMKTEQRELLQSLDSIANAYVQLALFDTSEYHGKKKKIPLSTVK IFETNSGRSGGTTFDQCLRARARRGDSVVLPAVLTSQIAPQPDMMNYSNVVRMYSFEPQFSITDSGIHRPKIYC YGSDEPEYKQLVKGQDDTRQDLVIEQVFETMNQFLMEEKATRKRKRLRTRVPLSPIAGVLEWVENTMP WGSYLVSRTSKRLSAHERYHPHEWKHTECRQYLNAPDKLPAFLEIEANFTPVFHHFFLEKFPDAAVWYQRR LSYVQSAAVTSIVGYILGIDRHSQNILIHEKTGELVHIDFGVVFQGMALYTPETVPFRLTRDMVDGGMGISG VDGVFSRCCVTLQLLRKKSASVVILEVFVHDPYRWLTSPLKALRIQEGQGHGKPTRSRSSSSGSAEYDG TGSMQDTQPADEMHAEPGSTDAAARALIRVKQKLEGYEDPNGSALSIEGQVQQLINAAQDPLNLCKLFPG WAPWL* |
| >scaffold_29_R3 08 | MTSSSCRLLPLWSGVSSVYSDAEIPLRRPFLVFLHRGFWYSRCQVPVSSLCFRIRTFLEVPWPPARVTRHR VHRSSGRLRFTLLSRRVCCSYRARRSLRSACRLRAFVRSYRWYSIGRLFSTQVLVFTVLRMDHPHLGDVLLRL LHCRRLCGALDCGLRSLFSILSDRPIAPISHATIRYIPDAGEFIPNSLWVRENCVGVSRCCRRIHRIPYANT WYYHGVSGRPNRYRIRHLRSYPLLPDVLHPLSYIIGVVHLRRLVYRLVKCLRLALSH* |
| >scaffold_21_F2 030 | MQRRMSTPFFLMITTIGDAQGEVEWVMIPADNSPLIKSDSFSRNCGATRRTRWATGVALPTSMSQTAPRA AGGRAFPFPIRCLRFQMICLNRSICTGVLPASSIWISSILTSSVVALTSGSMAI* |
| >scaffold_494_F 2 | MRLLYLAGVAIFAADAKVLVSDSVNNRAQAASASVVSTTRLLRTRSVIDEERAGGISASASDKLAKLFKSS KVTDEQLQKWLNNGKTAESVFYRMNLENTLYTRVFESQPFRWLQYADDLSASGKGASISVLSTKYGDDT LYKMIGWAKKESSTKALGIRLQTEQLEHWVKIGKDPDEVFKLYKLDYAGNRILGNPQFSAWTKYVDDLNAK NKGAFVSIPTLRKYISDDDLFKIALAAKRSGETKAMGTKLEDAIVQFWNRAGGPVGNPT* |
| >scaffold_100_R 322 | MRICFVLLLTVAALVTAVSGSSVNLNRDSQRIDVVQDKTSSRELRGDLNTEATEERASDFVKKLVAKLKGDPL ESFAKRQTKYIFSDDIFDEMLKKFPDPDALYTTKLHVIKNRSNKYGVTTARHKLHQNFLLTSYIDKFPNWRSKL N* |