Lawrence Berkeley National Laboratory

LBL Publications

Title

Genome Sequencing and Mapping Reveal Loss of Heterozygosity as a Mechanism for Rapid Adaptation in the Vegetable Pathogen Phytophthora capsici

Permalink https://escholarship.org/uc/item/5rw79183

Authors

Lamour, Kurt H. Mudge, Joann Gobena, Daniel <u>et al.</u>

Publication Date

2012-10-01

Genome Sequencing and Mapping Reveal Loss of Heterozygosity as a Mechanism for Rapid Adaptation in the Vegetable Pathogen Physophthora Capsici

Kurt H. Lamour,¹ Joann Mudge,² Daniel Gobena,¹ Oscar P.Hurtado-Gonzales,³ Jeremy Schmutz,^{4,5} Alan Kuo,⁴ Neil A.Miller,⁶ Brandon J. Rice,² Sylvain Raffaele,⁷ Liliana M. Cano,⁷Arvind K. Bharti,² Ryan

S. Donahoo,⁸ Sabra Finley,¹ EdgarHuitema,^{9,10} Jon Hulvey,¹¹ Darren Platt,⁴ Asaf Salamov,⁴ AlonSavidor,¹² Rahul Sharma,^{13,14,15} Remco Stam,^{9,10} DylanStorey,¹ Marco Thines,^{13,14,15} Joe Win,⁷ Brian J. Haas,¹⁶Darrell L. Dinwiddie,^{6,17} Jerry Jenkins,^{4,5} James R. Knight,¹⁸Jason P. Affourtit,¹⁸ Cliff S. Han,¹⁹ Olga Chertkov,¹⁹ Erika A.Lindquist,⁴ Chris Detter,¹⁹ Igor V. Grigoriev,⁵ SophienKamoun,⁷ and Stephen F. Kingsmore^{6,17}

¹University of Tennessee, Knoxville 37996, U.S.A.

²National Center for Genome Resources, Santa Fe, NM 87505, U.S.A.

³Pioneer Hi-Bred International, Johnston, IA 50131, U.S.A.

⁴United States Department of Energy Joint Genome Institute, Walnut Creek, CA 94598, U.S.A.

⁵Hudson Alpha Institute of Biotechnology, Huntsville, AL 35806, U.S.A.

⁶Children's Mercy Hospital, Kansas City, MO 64108, U.S.A.

⁷The Sainsbury Laboratory, Norwich NR4 7UH, U.K.

⁸University of Florida, IFAS-SWFREC, Immokalee 34142, U.S.A.

⁹Division of Plant Science, University of Dundee, Invergowrie, Dundee DD2 5DA, U.K.

¹⁰Plant Pathology Program, James Hutton Institute, Invergowrie, Dundee DD2 5DA, U.K.

¹¹University of Massachusetts-Amherst, Department of Plant, Soil, and Insect Sciences, Amherst 01003, U.S.A.

¹²Department of Molecular Biology and Ecology of Plants, Tel Aviv University, Tel Aviv 69978. Israel

¹³Biodiversity and Climate Research Centre, D-60325 Frankfurt am Main, Germany

¹⁴Senckenberg Gesellschaft für Naturforschung, D-60325 Frankfurt am Main, Germany

¹⁵Goethe University, Department of Biological Sciences, Institute of Ecology, Evolution and Diversity, D-60323, Frankfurt am Main, Germany

¹⁶Broad Institute, Cambridge, MA 02142, U.S.A.

¹⁷School of Medicine, University of Missouri-Kansas City, Kansas City, MO 64108, U.S.A. ¹⁸Roche Applied Science, Branford, Connecticut 06405, U.S.A.

¹⁹Los Alamos National Laboratory, Department of Energy Joint Genome Institute, Los Alamos, NM 87545, U.S.A.

October 2012

The work conducted by the U.S. Department of Energy Joint Genome Institute is supported by the Office of Science of the U.S. Department of Energy under Contract No. DE-AC02-05CH11231

DISCLAIMER

This document was prepared as an account of work sponsored by the United States Government. While this document is believed to contain correct information, neither the United States Government nor any agency thereof, nor The Regents of the University of California, nor any of their employees, makes any warranty, express or implied, or assumes any legal responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by its trade name, trademark, manufacturer, or otherwise, does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government or any agency thereof, or The Regents of the University of California. The views and opinions of authors expressed herein do not necessarily state or reflect those of the United States Government or any agency thereof or The Regents of the University of California.

Title: Genome sequencing and mapping reveal loss of heterozygosity as a mechanism for rapid adaptation in the vegetable pathogen *Phytophthora capsici*.

Authors: Kurt H. Lamour¹†*, Joann Mudge²†, Daniel Gobena¹, Oscar P. Hurtado-Gonzales³, Jeremy Schmutz^{4,5}, Alan Kuo⁴, Neil A. Miller⁶, Brandon J. Rice², Sylvain Raffaele⁷, Liliana M. Cano⁷, Arvind K. Bharti², Ryan S. Donahoo⁸, Sabra Finley¹, Edgar Huitema^{9,10}, Jon Hulvey¹¹, Darren Platt⁴, Asaf Salamov⁴, Alon Savidor¹², Rahul Sharma¹³⁻¹⁵, Remco Stam^{9,10}, Dylan Storey¹, Marco Thines¹³⁻¹⁵, Joe Win⁷, Brian J. Haas¹⁶, Darrell L. Dinwiddie^{6,17}, Jerry Jenkins^{4,5}, James R. Knight¹⁸, Jason P. Affourtit¹⁸, Cliff S. Han¹⁹, Olga Chertkov¹⁹, Erika A. Lindquist⁴, Chris Detter¹⁹, Igor V. Grigoriev⁴, Sophien Kamoun⁷, Stephen F. Kingsmore^{6,17}

Affiliations:

¹University of Tennessee, Department of Entomology and Plant Pathology, Knoxville, TN 37996, USA.

²National Center for Genome Resources, Santa Fe, NM 87505, USA.

³Pioneer Hi-Bred International, Johnston, IA 50131, USA.

⁴US Department of Energy Joint Genome Institute, Walnut Creek, CA 94598, USA.

⁵Hudson Alpha Institute of Biotechnology, Huntsville, AL 35806, USA.

⁶Children's Mercy Hospital, Kansas City, MO 64108, USA.

⁷The Sainsbury Laboratory, Norwich NR4 7UH, UK.

⁸University of Florida, IFAS-SWFREC, Immokalee, FL 34142, USA.

⁹Division of Plant Science, University of Dundee, Invergowrie, Dundee DD2 5DA, UK.

¹⁰Plant Pathology Program, James Hutton Institute, Invergowrie, Dundee DD2 5DA, UK.

¹¹University of Massachusetts-Amherst, Dept. of Plant, Soil, and Insect Sciences, Amherst, MA 01003, USA.

¹²Department of Molecular Biology and Ecology of Plants, Tel Aviv University, Tel Aviv 69978, Israel.

¹³Biodiversity and Climate Research Centre, D-60325 Frankfurt am Main, Germany.

¹⁴Senckenberg Gesellschaft für Naturforschung, D-60325 Frankfurt am Main, Germany.

¹⁵Goethe University, Department of Biological Sciences, Institute of Ecology, Evolution and Diversity, D-60323, Frankfurt am Main, Germany.

¹⁶Broad Institute, Cambridge, MA 02142, USA.

¹⁷School of Medicine, University of Missouri-Kansas City, Kansas City, MO 64108, USA.

¹⁸Roche Applied Science, Branford, Connecticut 06405, USA.

¹⁹Los Alamos National Laboratory, Department of Energy's Joint Genome Institute, Los Alamos, NM 87545, USA.

[†] These authors contributed equally to this work.

Running title: Adaptation in Phytophthora capsici.

*Corresponding author: Kurt Lamour, klamour@utk.edu.

ABSTRACT: The oomycete vegetable pathogen *Phytophthora capsici* has shown remarkable adaptation to fungicides and new hosts. Like other members of this destructive genus, *P. capsici* has an explosive epidemiology, rapidly producing massive numbers of asexual spores on infected hosts. In addition, *P. capsici* can remain dormant for years as sexually-recombined oospores, making it difficult to produce crops at infested sites, and allowing outcrossing populations to maintain significant genetic variation. Genome sequencing, development of a high-density genetic map, and integrative genomic/genetic characterization of *P. capsici* field isolates and intercross progeny revealed significant mitotic loss of heterozygosity (LOH) and higher levels of SNVs than those reported for humans, plants, and *P. infestans*. LOH was detected in clonally propagated field isolates and sexual progeny, cumulatively affecting >30% of the genome. LOH altered genotypes for more than 11,000 single nucleotide variant (SNV) sites and showed a strong association with changes in mating type and pathogenicity. Overall, it appears that LOH may provide a rapid mechanism for fixing alleles and may be an important component of adaptability for *P. capsici*.

INTRODUCTION

Phytophthora capsici is a virulent, hemibiotrophic pathogen of vegetable crops, inflicting significant losses worldwide (Leonian 1922; Erwin and Ribeiro 1996; Gevens et al. 2008; Lamour and Kamoun 2009; Meitz et al. 2010; Lamour et al. 2011). It's main hosts are peppers and cucurbits and during the last ninety years *P. capsici* has spread geographically as agricultural cultivation of these hosts has intensified; dynamically adapting to fungicides and new hosts (Leonian 1922; Erwin and Ribeiro 1996; Lamour and Hausbeck 2001; Hausbeck and Lamour 2004; Gevens et al. 2008; Hurtado-Gonzales et al. 2008; Lamour and Kamoun 2009; Meitz et al. 2010; Lamour et al. 2011; Gobena et al. 2012). Like other members of this destructive genus (e.g. the late blight pathogen *P. infestans* and the sudden oak death

pathogen *P. ramorum*), *P. capsici* has an explosive epidemiology, rapidly producing massive numbers of asexual spores on infected hosts (Lamour and Hausbeck 2001; Hausbeck and Lamour 2004; Lamour and Kamoun 2009) (Fig. 1B-G). In addition, *P. capsici* often features meiosis between two mating types, producing thick-walled oospores that remain dormant for years, making it very difficult to produce crops at infested sites, and allowing outcrossing populations to maintain significant genetic variation (Fig. 1A). Members of this genus are notoriously plastic in field and laboratory scenarios and we examined the molecular basis for rapid adaptation in *P. capsici* through genome sequencing, development of a high-density genetic map, and integrative genomic/genetic characterization of *P. capsici* field isolates and intercross progeny. Development of these tools and subsequent analyses revealed significant mitotic loss of heterozygosity (LOH) in clonally propagated field isolates as well as multiple sexual progeny. Follow-up tests revealed a strong association of LOH with changes in mating type and loss of pathogenicity.

RESULTS

Genome sequencing, assembly and annotation

Assembling the nuclear genome of a diploid organism with frequent nucleotide variants can be difficult and a partially inbred line of *P. capsici* (LT1534) was developed for sequencing (Supplementary Fig. 1) (Hurtado-Gonzales and Lamour 2009; Gobena et al. 2012). The 64 Mbp reference genome was assembled using Arachne from 30X paired and singleton 454 FLX and Titanium genomic DNA reads, 5X Sanger paired reads, 56,448 Sanger ESTs, and 1,260 full-length cDNA sequences. This resulted in 917 scaffolds and an N50 of 706 kbp (Supplementary Table 1). The size of the *P. capsici* genome is intermediate among the sequenced oomycetes, larger than *Albugo* and *Pythium ultimum* (up to 45 Mbp) (Levesque et al. 2010; Kemen et al. 2011), but more compact than *P. ramorum*, *P. sojae*, *Hyaloperonospora arabidopsis* and *P. infestans* at 65, 95, 100, and 240 Mbp, respectively (Tyler et al. 2006; Haas et al. 2009; Baxter et al. 2010). The *P. capsici* draft genome had fewer scaffolds and/or larger scaffold N50, which may reflect hybrid assembly or fewer repeat sequences in *P. capsici* (19%) than *P. infestans*, *P. sojae* or *P. ramorum* (74%, 39%, and 28%, respectively) (Tyler et al. 2006; Haas et al. 2009). Like other Phytophthora species, the majority of *P. capsici* repeats were retrotransposons (84%) with long terminal repeats, of which 57% were Gypsy elements. The mitochondrial genome (80,148 bp) was assembled in 4 scaffolds.

Gene model prediction using a training set of 16.4 million sequencing-by-synthesis mRNAsequences from nine life stages predicted 19,805 genes supported by cDNA sequences (50%) or previously identified proteins (75%) (Supplementary Tables 2 and 3). After excluding 2,682 genes with homology to transposable elements, *P. capsici* had a similar gene content to *P. infestans*, *P. sojae* and *P. ramorum* (17,797, 16,988 and 14,451, respectively) (Tyler et al. 2006; Haas et al. 2009) and had slightly more core eukaryotic genes identified compared to these other species (Supplementary Fig. 2). The nonrepetitive gene density in *P. capsici* (268/Mbp) was higher than other Phytophthora species (74/Mbp in *P. infestans*, 179/Mb in *P. sojae* and 222/Mb in *P. ramorum*) (Tyler et al. 2006; Haas et al. 2009).

Comparison to other Phytophthora species

Genome comparisons revealed almost perfect scaffold-level synteny between the gene models of *P. capsici, P. ramorum* and *P. sojae*, with very little duplication (Fig. 2A, B). There was also extensive synteny with *P. infestans* genes (Supplementary Fig. 3). The genome organization of *P. capsici* was similar to other Phytophthora species, with ²/₃ of the genes located in gene-rich blocks that have a conserved order and few repeats (median intergenic distance 351 bp), separated by relatively gene-poor regions without order conservation and more repetitive sequences (median intergenic distance ~3kb; Fig. 2C) (Haas et al. 2009). Like *P. infestans*, the 365 predicted *P. capsici* genes without a homolog resided predominantly in gene-poor genomic regions (Fig. 2C) (Haas et al. 2009; Raffaele et al. 2010).

Seventy eight percent of the predicted *P. capsici* genes clustered into 2,483 candidate multigene families. Among these families, there are cytoplasmic effector genes of the RxLR type including

homologs of known avirulence genes in *P. infestans* (Whisson et al. 2007; Dou et al. 2008; Boutemy et al. 2011). *P. capsici* had 357 RxLR effector genes, which is a similar to the numbers reported for *P. sojae* and *P. ramorum*, whereas *P. infestans* had > 500 (Tyler et al. 2006; Haas et al. 2009). *Phytophthora capsici* also had 29 putative full length and 70 pseudogenes for Crinkling and Necrosis (CRN) effectors, an ancient class of intracellular effector proteins that share a highly conserved N-terminal domain, required for translocation, and diverse C-terminal effector domains (Haas et al. 2009). While markedly fewer than in *P. infestans*, the majority of predicted CRN effector domains were conserved between species.

Single nucleotide variation (SNV) and genetic mapping

Nucleotide diversity was assessed in seven *P. capsici* field isolates (including parental isolates TN1 and TN2) and an isolate of the sister species *P. tropicalis* by restriction-site associated DNA (RAD) sequencing (Supplementary Table 4) (Baird et al. 2008). Between 5 and 9 million sequencing-by-synthesis reads, flanked by a *Sgr*AI restriction site, were generated per isolate (Table S5). The reads were aligned to the *P. capsici* reference genome and covered ~2.3 Mbp at a depth of \geq 30-fold (Kim et al. 2009; Baranzini et al. 2010; Bell et al. 2011). The SNV density among *P. capsici* isolates was 23.4/kb (range 16.9 - 25.9/kb) and between *P. capsici* and the closely related species *P. tropicalis* was 51.7/kb (Supplementary Tables 6 and 7; Supplementary Figs. 4 and 5). This is much higher than the SNV diversity observed among *P. infestans* strains (0.65/Kb average) and between *P. infestans* T30-4 and species in its clade (7.67/Kb average) (Raffaele et al. 2010). The nucleotide diversity (π) was 0.012 (range 0.009 - 0.016), ~16-fold higher than *H. sapiens* and ~4-fold higher than obtained by RAD in natural stickleback populations and most outbred plant species. The threespined stickleback (*Gasterosteus aculeatus* L.) is an outcrossing species of fish studied to better understand adaptive evolution (Hohenlohe et al. 2010).

In addition, the inheritance of SNV's was examined in 65 putative F₁ intercross progeny by *Pst*Ibased RAD (Supplementary Figs. 1 and 6; Supplementary Table 8). Of these 65 isolates, two were found to be clonally derived from the TN2 parent and three additional isolates were clonal to progeny isolates. The remaining 60 true F1 progeny were genotyped at bi-allelic sites exhibiting either a di- or tri-modal distribution where the peaks corresponded to homozygous reference alleles, heterozygotes, and homozygous alternate alleles (Supplementary Fig. 7) (Kim et al. 2009). Genotypes were imputed at 500,352 nucleotides with >10 uniquely aligned reads in >55 progeny. Homozygosity was inferred when >90% reads displayed one allele, and heterozygosity when 20-80% of reads displayed one allele. We have previously shown such genotype imputation to be >99% accurate (Kim et al. 2009; Baranzini et al. 2010; Bell et al. 2011). Forty-five percent (9,254) of SNVs showing Mendelian inheritance were in coding regions, and 3,382 (37%) of these were non-synonymous (Supplementary Fig. 8a, Supplementary Table 10). The distribution of mutations in *Phytophthora* clade 1c species is in a similar range with 35% of SNV's in coding regions, 41% of which are non-synonymous (Raffaele et al. 2010). This was surprising since there is typically extensive purifying selection on genes. A parsimonious explanation of these findings is that rates of mutation relative to generation times are much greater in P. capsici than higher eukaryotes.

A total of 20,568 SNVs were either polymorphic in the parents and had Mendelian segregation $(\chi^2 \ge 0.05)$, or were homozygous for opposite alleles in the parents and heterozygous in progeny that did not exhibit loss of heterozygosity (LOH) (marker spacing 3,111 bp; Supplementary Table 9). A genetic map was developed using markers on the 108 scaffolds that had sufficient SNVs for independent genetic linkage analysis in the F₁ progeny. Of these, 92 scaffolds had SNV's that were inherited as a single co-segregating block, while markers on 15 scaffolds broke into 2 blocks and markers on the largest scaffold (Scaffold 1) broke into three blocks (Supplementary Table 11; Supplementary Fig. 9). Scaffold-level linkage analysis, a genetic map of *P. capsici* was constructed with 1,136 SNVs that had segregation patterns <95% similar.

The genetic map was 1,654 centiMorgans (cM) in length and comprised eighteen linkage groups accounting for 84% (54Mbp) of the reference genome and containing ~90% of the predicted genes (Fig. 3; Supplementary Table 12). The order for the majority of markers was the same on the genetic and physical maps. Genetic linkage analysis localized the mating type locus to a ~300kb critical region on scaffold block 4.1 in LG10.

Loss of heterozygosity

Integrated genomic and genetic analysis allowed assessment of chromosomal variation among the 65 putative F_1 intercross progeny. Twenty-three F_1 isolates and two clonal isolates contained regions with LOH. LOH was identified by homozygosity at all markers in ≥ 2 adjacent (linked) RAD sequences that were incompatible with parental genotypes (Supplementary Fig. 10; Supplementary Table 13). The minimum length of LOH tracts varied from 299 bp to >1 Mbp. LOH occurred in 14 of the 18 LGs and in 59 scaffolds or scaffold blocks. Fifty-four percent (11,048) of SNVs exhibited LOH in at least one isolate and the LOH was bidirectional. The frequency and distribution of LOH tracts varied among isolates and 36.5% (19.7 Mbp) of the genome was affected by LOH in at least one isolate. Two isolates (TN37 and TN47) had LOH exceeding 10% of the genome (Supplementary Table 14). Interestingly, LOH showed bias toward gene-rich regions (Supplementary Fig. 8B, C). Given the requirements of informative parental genotypes, at least two adjacent RAD tags to be affected, and that LOH was sought only in the mapped genome, the true extent of LOH is likely to be considerably higher.

Sub-chromosomal mitotic LOH has several molecular mechanisms including deletions (resulting in hemizygosity) and translocations (resulting in homozygosity) associated with homologous and non-homologous recombination or gene conversion (Chamnanpunt et al. 2001; Cvitanich et al. 2006). Of note, mutagenesis-induced LOH in *P. capsici* was not associated with sub-chromosomal deletions (Hulvey et al. 2010). In only one of 12 isolates was LOH due to hemizygosity as assessed by batched, normalized sequence coverage plots (Bell et al. 2011) (Fig. 4A-C). Forty-five percent (4,967) of the

marker nucleotides exhibiting LOH were exonic, 17% (1,911) were in coding regions and 3.5% (391) were predicted to be non-synonymous, affecting 264 genes. Phenotypic consequences of genome diversity were sought among F₁ progeny. Five F₁ isolates switched mating type from A2 to A1 between assessments in 2005 and RAD sequencing in 2010, while one switched from A1 to A2 (Fig. 5A). Isolates were in storage for most of this interval and were sub-cultured < 10 times. Four of the isolates that switched from A2 to A1 had >800 Kbp tracts of LOH encompassing a region that appears to contain the mating type locus (or loci) on LG10 (Fig. 5A). TN25, which changed from A1 to A2, did not exhibit LOH at the mating type locus but did have >1.6 Mbp LOH in LG16. However, A1 mating type isolates TN37 and TN70 had similar LOH on LG16, but did not change mating type. Thus, LOH at the mating type locus was associated with mating type switch in four of six isolates.

LOH was also associated with a loss of pathogenicity to pepper and cucumber fruit. Between tests in 2005 and 2011, progeny isolates TN37 and TN47 became unable to infect wounded or un-wounded fruit (Fig. 5B). In 2011, host cells consistently elicited a strong hypersensitive response (cell necrosis) at point of inoculation, suggesting loss of a factor necessary for infection, such as an Avh protein. This is characteristic of Phytophthora Avh proteins, which evolve rapidly in a presumed host-pathogen arms race (Tyler et al. 2006; Whisson et al. 2007; Jiang et al. 2008; Haas et al. 2009). These isolates had similar growth rates *in vitro* in 2005 and 2011 even after substantial LOH (Supplementary Table 14). Indeed, the TN37 and TN47 genomes each had > 10% LOH and share an LOH tract on LG1 (17_31,860 - 17_870,195 and 105_51,382 - 105_63734; Supplementary Tables 13-14). A plausible explanation is that LOH changed the expression or expressed sequence of one or more *P. capsici* effectors in the avirulent isolates.

DISCUSSION

In summary, the genome sequence of *P. capsici* was typical for the genus in organization and gene content. A prototypical, dense genetic map indicated the recombination rate of *P. capsici* to be 38.7 kbp/cM, an order of magnitude less than yeast, but 30-fold higher than *H. sapiens* (Li and Freudenberg

2009). SNV density and diversity among *P. capsici* genomes was much higher than in other eukaryotes and coding and non-synonymous variants were remarkably common. This suggests that germline mutation rates may be much higher in *P. capsici* than in *H. sapiens*, *Saccharomyces cerevisiae*, *Drosophila melanogaster* and *P. infestans*, or that purifying selection was less (Baranzini et al. 2010). The overall SNV diversity for the seven *P. capsici* field isolates was similar, but markedly different from the *P. tropicalis* isolate. The isolates from Peru and Argentina appear to be part of long-lived, widely dispersed clonal lineages where, unlike the situation for the North America isolates, sexual recombination may be rare (Hurtado-Gonzales et al. 2008; Gobena et al. 2012). Interestingly, the isolates from Peru and Argentina share the most polymorphic loci and may be derived from a common, invasive population or widely dispersed clonal lineage. The high level of SNV diversity in *P. capsici* has practical implications for research. The SNV's can be used to finely map segregating phenotypes in larger populations of progeny, conduct sensitive analyses of population structure and overall dynamics, and to track the fate of numerous heterozygous loci during asexual growth (e.g. tracking LOH).

Phytophthora capsici populations are often characterized by high levels of diversity and rapid adaptation (Leonian 1922; Erwin and Ribeiro 1996; Lamour and Hausbeck 2001; Hausbeck and Lamour 2004; Gevens et al. 2008; Hurtado-Gonzales et al. 2008; Lamour and Kamoun 2009; Meitz et al. 2010; Lamour et al. 2011; Gobena et al. 2012). In addition to elevated genetic diversity, *P. capsici* genomes also had a remarkably high frequency of mitotic LOH occurring more commonly in gene rich areas. Mitotic genomic instability was also recently described in *P. ramorum* (Vercauteren et al. 2011), suggesting LOH may be a general characteristic of oomycetes. Here we show loss of pathogenicity and mating type switches are likely phenotypic consequences of mitotic LOH. For the latter, the evidence for causality of LOH was strong: four of five progeny that switched from the A2 to the A1 mating type had tracts of LOH spanning the region where mating type maps in this cross. This was reminiscent of the mating type locus in the fungus *Candida albicans*, where a repressor suppresses switching to the matingcompetent form (Morschhauser 2010). Loss of mating type heterozygosity relieves this repression,

Page 10 of 90

ensuring that only mating type homozygous cells are mating-competent. An ability to switch mating type has important consequences, particularly in northern latitudes where survival of the winter or fallow months requires thick-walled, sexually-produced oospores (Lamour et al. 2011). It also provides a mechanism to introgress mutations efficiently into a population (Supplementary Figure 10). The fungicide mefenoxam, for example, is strongly inhibitory to *P. capsici* and has been widely used in some areas. Mefenoxam resistance is inherited in *P. capsici* as a single incompletely dominant trait (Lamour and Hausbeck 2000; Lamour and Hausbeck 2001). Mating type switches in a rare, partially resistant field isolate will double the chances for outcrossing. In addition to aiding introgression of a novel resistance allele by sexual reproduction, LOH could directly catalyze the conversion to full resistance. Using the SNV markers presented here, it should be possible to detect instances of LOH within naturally occurring isolates.

The dramatic expansion of repeated sequences observed in several pathogen lineages of oomycetes (*P. infestans*, *H. arabidopsidis*) and fungi (*Puccinia graminis* f. sp. tritici, *Melampsora larci-populina*, *Blumeria graminis*) was proposed to contribute to fast adaptation (Haas et al. 2009; Spanu et al. 2010). A clear correlation was indeed observed between the rate of gene evolution and the amount of flanking repeats (Raffaele et al. 2010). By contrast, the *P. capsici* genome is rather compact, with a relatively low repeat content, and yet shows remarkable plasticity. This plasticity may be due in part to LOH which represents an alternative mechanism for rapidly generating diversity, preferentially affecting gene dense regions. Previous studies suggest a role for LOH in adaptive evolution (Rosenberg 2011). It has, for example, been associated with oncogenesis and embryonic stem cell adaptation in *H. sapiens* (Narva et al. 2010), and fluconazole resistance in *C. albicans* (Forche et al. 2011). Here we extend these observations, suggesting that the co-occurrence of frequent mitotic LOH and high nucleotide diversity in genes may provide a general mechanism for rapid adaptation in *P. capsici*. In addition, LOH may contribute to the difficulties encountered when studying oomycetes in the laboratory (e.g. loss of pathogenicity and virulence) and the overall success of oomycetes as plant pathogens.

METHODS

Isolates. Three *in vitro* crosses were completed to produce a moderately inbred isolate for sequencing the reference genome (Hurtado-Gonzales and Lamour 2009). Crosses include an F1, and two recurrent backcrosses (BC1 and BC2). An isolate from the BC2 progeny was chosen for genome sequencing based on growth rate and asexual and sexual spore production (Hurtado-Gonzales and Lamour 2009). The initial cross was between an A1 isolate recovered from cucumber in Michigan in 1997 (TN1) and an A2 isolate recovered from pumpkin in Tennessee in 2004 (TN2). The backcrosses were to the TN2 parent. Crosses were made by co-culturing the parental isolates on V8-juice agar plates, incubating the plates for 8 to 12 weeks in darkness, and then separating the sexual oospores from the subtending asexual mycelium and other asexual spores (sporangia and zoospores) through mechanical shearing and an enzyme treatment to destroy the asexual material (Lamour and Hausbeck 2000). The enzymes not only destroy the asexual material; they stimulate oospore germination. Germinating oospores were harvested using a 5µl pipette while looking at the spores under a light microscope. Once harvested, the germinated spores are placed onto V8 media, sub-cultured to produce mycelium for DNA and to conduct mating type analysis, and then stored long term (3 to 5 years) in water with sterile hemp seeds.

The progeny from all of the crosses were genotyped using AFLP markers, tested for mating type, assessed for growth rate on agar media, and assayed for pathogenicity and virulence on detached pepper (variety Early Jalapeno) and cucumber fruit (various varieties) (Hurtado-Gonzales and Lamour 2009). All fruit inoculations were conducted on two fruit and the experiments repeated twice. Pathogenicity was assessed by placing a 7mm plug of mycelium at both a wounded and non-wounded site. Wounded sites are distal and non-wounded sites proximal to the stem end of the fruit. Pathogenicity to pepper fruit was assessed in 2005 and 2011. Sixty-nine progeny from the F1 were further analyzed using a focused re-sequencing strategy (outlined below). In addition, the mating type was tested at three different time intervals: within 6 months of the cross, at the time of the focused re-sequencing, and one year following the re-sequencing.

Genome sequencing and assembly. High quality genomic DNA was prepared from the isolate selected for genome sequencing using a phenol/chloroform protocol. Aliquots from the same batch of DNA were submitted to 454 Inc. (Branford, CT) and to the Joint Genome Institute (JGI, Walnut Creek, CA) for 454- and Sanger-based sequencing, respectively. A total of 2,451,737 random shotgun genomic reads, 2,022,402 paired-end reads (2.3 Kb), and 833,606 paired-end reads (18.0 Kb) were produced using the 454 Titanium chemistry. In addition, 197,306 random genomic paired-end reads from a 6.3Kb plasmid library, 146,588 paired-end reads from a 34.7Kb fosmid library and 65,000 ESTs were produced at the JGI using Sanger sequencing. Additionally, 1,260 full-length cDNA sequences were generated by Sanger sequencing and cDNA sequence assembly.

The sequence reads were assembled using a modified version of Arachne v.20071016 (Jaffe et al. 2003) with parameters maxcliq1=100, n haplotypes=2 and BINGE AND PURGE=True. The output of this assembly was passed through Rebuilder and SquashOverlaps with parameters to merge adjacent assembled alternative haplotypes, and subsequently run through another complete Arachne assembly process to finalize the assembly. Prior to assembly, redundant 454 pairs (defined as at least 98%) identical across their length) were removed and a single pair retained. An error correction step was applied to all 454 data that attempted to set bps to equal the Sanger sequence. This was accomplished by shredding the Sanger sequence to a similar size as the 454 reads and then aligning these shredded reads and the 454 simultaneously. Where the 454 read sequence had no support we changed the 454 read sequence to match the Sanger read sequence. This removed many of the 454 sequence errors (primarily indels and occasional bp substitutions) and coalesced the haplotyopes if the Sanger data was all from one haplotype. Error correction was also applied during the Arachne assembly process by screening each scaffold against bacterial proteins, organelle sequences and GenBank using Megablast against Genbank NR and blastp against a set of known microbial proteins. Twenty-eight scaffolds identified as prokaryotic contamination were removed. We classified additional scaffolds as mitochondrion (4), small repetitive (6), alternative haplotyope (17), and less than 1kb (1050). The final assembly consists of 971

scaffold sequences, with half of the genome contained in 29 scaffolds of at least 705.7 kb in length and contigs greater than 34.6kb. Scaffolds ranged in size from 1 kb to 2.1 Mb with an N50 of 705.7 kb (Supplementary Table 1). The genome size was tested using the approach outlined by Baxter with nt by nt coverage calculated by aligning back to the reference (Baxter et al. 2010). This resulted in an estimated genome size of approximately 55 Mbp. We also estimated the genome size by measuring the percentage of all reads that were captured by the assembly using a high quality 10,000 read subset (average quality \geq 35, length \geq 100 and no missing bases) and extrapolating genome size if the assembly had captured all the reads. We extrapolated up from the genome size that doesn't include gaps (56.1 Mb) and 97.44% of the reads were captured leading to an estimated genome size of approximately 57 Mbp. Both estimates likely underestimate the genome size due to over alignment of repeat copies, possibly explaining the discrepancy between our gapped genome size (64.1 Mb) and these lower estimates of total genome size. Nevertheless, both estimates are similar to the non-gapped genome size (56.1 Mb) suggesting that the assembly is missing very little of the non-repetitive genome.

In addition, total RNA was isolated from different growth conditions (rich media, starvation, and sporulating), pooled, and submitted to both 454 and the JGI for Sanger cDNA and EST sequencing. At the JGI, ESTs were produced as previously described with the following modification: 2 cDNA libraries were constructed and sequenced and the size ranges of the cDNA library inserts were 0.6k-2kb and >2kb (Jeffries et al. 2007).

Genome sequence analysis and annotation of gene families. The genome assembly of *Phytophthora capsici* LT1534 was annotated using the JGI Annotation Pipeline, which combines several gene predictors: A) cDNA-based gene models were derived from 1260 full-length cDNAs and 11,090 consensus sequences clustered from 56,448 ESTs, and then mapped to genomic sequence, B) protein-based gene models were predicted using FGENESH+ (Salamov and Solovyev 2000) and GeneWise (Birney et al. 2004) seeded by BLASTx alignments of genomic sequence against sequences from the NCBI non-redundant protein set nr, and C) *ab initio* gene models were predicted using FGENESH

trained on the set of putative full-length genes and reliable protein-based models. GeneWise models were completed using scaffold data to find start and stop codons. ESTs, EST clusters, EST contigs, and full length cDNAs were used to verify, complete, and extend the gene models. Because multiple gene models per locus were often generated, a single representative gene model for each locus was chosen based on homology and EST support, and used for further analysis. This led to a filtered set of 19,805 gene models (including 2,682 genes with homology to transposable elements) with their properties and support by different lines of evidence summarized in Supplementary Tables 2 and 3. All gene-based analyses were done on the full 19,805 gene set. A total of 56,448 ESTs are available in Genbank under the title "DOE Joint Genome Institute *Phytophthora capsici* EST project". An additional 1260 full length cDNAs are available in Genbank under the title "DOE Joint Genome Institute *Phytophthora capsici* EST project". The genome sequence is available at http://genome.jgi-psf.org/Phyca11/Phyca11.home.html.

All predicted gene models were functionally annotated by the JGI Annotation Pipeline using InterProScan (Zdobnov and Apweiler 2001) and hardware-accelerated double-affine Smith-Waterman alignments (http://www.timelogic.com/) against highly curated databases such as SwissProt (Bairoch et al. 2005), KEGG (Ogata et al. 1999), and Pfam (Bateman et al. 2004). KEGG hits were used to map EC numbers (Bairoch 2000), and InterPro, KEGG, and SwissProt hits were used to map GO terms (Ashburner et al. 2000). In addition, predicted proteins were annotated according to KOG classification (Koonin et al. 2004). Protein targeting predictions were made with signalP (Nielsen et al. 1999) and TMHMM (Krogh et al. 2001). Finally, all proteins were aligned by BLASTp to proteins in nr and to each other; after the latter analysis the alignment scores were used as a distance metric for clustering by MCL (<u>http://www.micans.org/mcl/</u>) into a first draft of 2,483 candidate multigene families. The same method was used to group genes with those of other *Phytophthora* species. We used BLASTP to identify Crinkler (CRN) coding genes from the *P. capsici* gene model set (Supplementary Table 16). For this purpose, 16 well characterized CRN proteins from *P. infestans* were searched against the *P. capsici* protein data set. Sequences with significant hits (E= <10-5) were retrieved and a non-redundant CRNlike protein set was created (Win et al. 2006). CRN-like protein sequences were manually examined for the presence of LFLAK motifs and C-terminal effector domains. To determine whether candidates are full length or pseudogenes, we aligned our CRN candidates with previously characterized CRNs. CRN proteins of at least 250 amino acid residues or more and that aligned with both the N and C-terminal domains of PiCRNs were considered full length (Haas et al. 2009). Finally, we used SignalP3.0 to identify predicted secretion signals for all CRN candidates (Supplementary Table 16). Candidate RxLR effectors were mined via the protocol outlined previously (Haas et al. 2009). In brief, all open reading frames greater than 210 nucleotides (70 AA) from the genome were generated and translated to amino acid coding sequences. Those containing an RXLR (Arg - Anything - Leu - Arg) between 30 - 60 amino acids from the start, a signal peptide score greater than .9, and a cleavage site prior to the RXLR are considered effectors (Supplementary Table 17).

The CEGMA pipeline was used to compare the completeness and continuity of six Oomycete genomes (Parra et al. 2007). CEGMA checks the completeness and continuity of genome assemblies on the basis of 248 core eukaryotic genes. Here, we compared the CEGMA analyses (without KOG 69) for the genomes of *P. sojae, P. ramorum, P. infestans, Pythium ultimum,* and *Hyaloperonospora arabidopsis* to the draft genome of *P. capsici* reported here (Supplementary Table 15).

Transposon-like coding sequences were identified by searching the annotated protein-coding sequences of *P. capsici* against databases of transposon sequences with TransposonPSI (<u>http://transposonpsi.sf.net</u>). Nucleotide sequences corresponding to the coding regions of the annotated protein coding genes were searched against the TransposonPSI collection of transposon protein sequences using BLASTX. This included ORFs derived from mobile elements discovered in other sequenced Phytophthora genomes. In addition, the *P. capsici* protein sequences were searched against a collection of PSI-BLAST profiles corresponding to common families of transposable elements, including Gypsy-retrotransposons and Piggybac-transposons. Any *P. capsici* annotated coding sequence

identified by having a BLAST E-value of at most 1e-10 was identified and reported, yielding 2,682 candidate gene annotations with transposon homology (see: Pcap.annot_transposon_mining.maxE_1e-10.txt.gz). Transposon family (PSI-BLAST) and top transposon protein BLASTX match are provided. Scaffolds, gene models clusters, and annotations thereof, may be accessed at the JGI *P. capsici* portal (http://www.jgi.doe.gov/capsici/) (Grigoriev et al. 2012).

Focused re-sequencing. Restriction site Associated DNA (RAD) sequencing was employed to identify and genotype single base substitutions in 7 field isolates (including the parents TN1 and TN2) and 69 progeny (TN3 – TN71) of the F1 cross (Baird et al. 2008). Briefly, genomic DNA was digested with the 6bp recognition restriction endonuclease *Pst*I or the 8bp recognition *Sgr*AI (Floragenex, Inc.). The digested DNA was sheared to between 200 and 500 bp and adaptors ligated which contained sequencing priming sites and an isolate-specific 5bp tag (Baird et al. 2008). Between 5 and 9 million sequencing-bysynthesis reads of length 72 nt were generated per isolate using an Illumina GAII sequencer (Supplementary Fig. 6, Supplementary Tables 5 and 8). The short reads abutting the restriction site are referred to as RAD tags. The RAD approach reduces complexity by focusing the sequencing to the regions directly adjacent (left and right) to a restriction site and the isolate-specific 5bp tags allow multiplexed sequencing. Reads were aligned to the *P. capsici* reference, covering ~2 million nucleotides at a depth of \geq 30-fold (Supplementary Tables 5 and 8) (Kim et al. 2009; Baranzini et al. 2010; Bell et al. 2011).

SNV discovery and genotyping. Alignments of the RAD sequence and subsequent variant detection were carried out using the Genomic Short-read Nucleotide Alignment Program (GSNAP (Wu and Nacu 2010)) and the Alpheus pipeline (Miller et al. 2008), as described (Kim et al. 2009; Baranzini et al. 2010; Bell et al. 2011). Reads were aligned to the *P. capsici* genome assembly (version 11) and alignments retained that matched the genome with 95% identity or higher and that had \leq 5 equally high scoring alignments. Reads with equally good hits to more than one genomic region, reflecting repeat

content or redundancy in the assembly, were not considered uniquely aligned. Four isolates were excluded from analysis due to insufficient aligned reads (Supplementary Fig. 6). Read count–based allele frequencies of bi-allelic nucleotide variants had a tri-modal distribution, with peaks corresponding to homozygous reference alleles, heterozygotes, and homozygous alternate alleles (Supplementary Fig. 7) (Kim et al. 2009).

Potential segregating SNV sites were identified in the parents and progeny using the following criteria: $\geq 10X$ unique sequence coverage, an average quality score ≥ 20 and an alternate allele frequency $\geq 20\%$ within unique reads. This pool of potential segregating sites was then analyzed across all lines and SNV genotyping criteria were developed based on a visual assessment of the distribution of alternate alleles at different levels of unique sequence coverage (Supplementary Fig. 7). SNVs were genotyped at the identified segregating SNV sites for all lines with at least 10X unique coverage where alternate allele frequency within unique reads was <10% (homozygous for the reference allele), >20% and <80% (heterozygous), or >90% (homozygous for the alternate allele). Our final pool of markers was limited to loci with 5 or fewer missing genotypes in the progeny. This approach does not enrich or bias for gene rich areas of the genome.

Nucleotide density and diversity was calculated for SNV sites in the *Sgr*AI RAD tags from the *P*. *capsici* parents and five additional unrelated isolates (two of which represent very large clonal populations in Peru and Argentina) and a *P. tropicalis* isolate. Heterozygosity and homozygosity were distinguished by allele frequency cutoffs of 15% and 85% and there was no requirement for number of missing genotypes in the isolates. SNV diversity was calculated as previously described (Begun et al. 2007).

Loss of heterozygosity and clonal isolates. Loss of Heterozygosity (LOH) was assessed using graphical genotypes constructed by ordering the SNV loci according to (i) the genetic linkage of markers in the individual scaffolds and (ii) the genetic linkage of markers from all scaffolds together

(see below). The minimum tracts of LOH were calculated for regions with continuous, uninterrupted switching of loci to either the reference or alternate allele (homozygosity) in a way that is impossible by normal meiosis or clonal reproduction (Judd and Petes 1988). To avoid minor genotyping errors, LOH was only assigned for tracts where the switched loci spanned at least two different sequencing sites. In addition, the junction of LOH versus non-LOH in an isolate with good sequence coverage was analyzed to compare the coverage on either side of the junction.

Genetic similarity of the parents and progeny was assessed in JMP Genomics by analyzing a relationship matrix for our final pool of markers using a Fast Ward Hierarchical clustering at the default settings. Alleles were assumed to be identical by descent. Putative clones were confirmed using the graphical genotypes. Genetic similarity analyses revealed three clonal lineages. One clonal lineage contained the parent isolate TN2 and the (mistaken) F1 progeny TN3 and TN50. Two other clonal lineages were derived from true F1 progeny and included isolates TN56, TN57, and TN63 in one clonal lineage and isolates TN8 and TN11 in another clonal lineage (Supplementary Fig. 11). This was not surprising since occasional sporangia survive the enzymatic treatment used to destroy asexual propagules, and premature oospore germination can produce sporangia that appear to be discrete oospore progeny. A single isolate of each lineage was retained for genetic linkage analysis. Apomixis, which occurred in other crosses stemming from these parents, is another possible explanation for the clones of the TN2 parent (Hurtado-Gonzales and Lamour 2009).

Genetic linkage analysis. Putative segregating markers were analyzed using a Chi-square test for simple Mendelian inheritance (5% significance). A total of 20,568 SNVs were either polymorphic in the parents and had Mendelian segregation ($\chi 2 \ge 0.05$), or were homozygous for opposite alleles in the parents and heterozygous in progeny that did not exhibit LOH. These were used for genetic linkage analysis (average marker spacing 3,111 bp; Supplementary Table 9).

A total of 3,141 SNVs exhibited segregation ratios <5% by χ 2-test. Some of these clustered in blocks and may reflect gene conversion, as described in other Phytophthora species (Chamnanpunt et al. 2001; Cvitanich et al. 2006). An additional ~100 loci were heterozygous in both parents and all progeny and may reflect assembly artifacts, copy number variants or obligate heterozygosity.

Mendelian markers were further analyzed using JoinMap 4.1 at the default settings for a CP type cross (heterogeneously heterozygous with phase unknown) (van Ooijen 2011). Initially, markers from individual scaffolds were analyzed to determine if they are inherited as a linked unit (expected if the scaffold is assembled correctly). If a scaffold broke into more than one piece, the pieces are referred to as scaffold blocks. Following this, markers from all the scaffolds, or scaffold blocks, were analyzed together to produce a linkage map. The LGs were further refined to include only markers with segregation patterns <95% similar and to exclude isolates showing any LOH within the LG. Markers are named with the scaffold number followed by the nucleotide position on the scaffold. Additional information was included for the markers used in JoinMap 4.1 to designate the three possible segregation scenarios: hk = heterozygous in both parents, nn = Parent 1 is homozygous and Parent 2 is heterozygous, and lm = Parent 1 is heterozygous and Parent 2 is homozygous.

The SNV genotyping error rates were assessed for both parents (TN1 and TN2) at 95 polymorphic sites in portions of 30 genes across 14 linkage groups by Sanger sequencing of PCR products. Of these 190 genotypes, all but 5 were confirmed. The 5 genotypes that differed occurred on a single gene that switched to homozygosity at all 5 sites.

ACKNOWLEDGEMENTS

Sequences, assembly and annotations are available via the JGI Genome Portal at http://jgi.doe.gov/capsici and at DDBJ/EMBL/GenBank under the accession ADVJ00000000.

This work was supported by grants to SFK from the National Science Foundation (05-512), US Department of Agriculture, and Joint Genome Institute Community Sequencing Program. The work conducted by the U.S. Department of Energy Joint Genome Institute is supported by the Office of Science of the U.S. Department of Energy under Contract No. DE-AC02-05CH11231. JM was supported in part by NIH grant RR016480. SR was supported by a Marie Curie Fellowship (contract 255104). RStam and EH were supported by the BBSRC and Royal Society of Edinburgh. LC, JW and

SK were supported by the Gatsby Charitable Foundation. RSharma and MT are supported by the LOEWE Initiative of the state of Hesse. KL was supported by a grant from the UTIA AgResearch program. We thank Paul Richardson and Eddy Rubin for JGI support in genome sequencing.

AUTHOR CONTRIBUTIONS

Manuscript Preparation: KHL, SFK, JM, SK, EH, SR, IVG, AK, RStam, AKB, MT; Crosses, phenotypic analyses, and genetic analyses: KHL, DG, OPH, ASavidor, JH, and DS; Genomic or Transcriptomic Sequencing: JK, JA, EAL, SFK, JM, DLD, CD, OC, CSH; Assembly: JS, DP; Annotation: AK, ASalamov, IVG; Alignments and variant detection: NAM, BJR, JM KHL; Analysis: KHL, JM, SFK, SK, EH, DS, SR, LMC, AK, IVG, AKB, RSD, SF, ASavidor, DG, OPH, JH, RSharma, RStam, DS, MT, JW, BJH, CD, CSH, OC.

References and Notes:

- Ashburner M., Ball C. A., Blake J. A., Botstein D., Butler H., Cherry J. M., Davis A. P., Dolinski K., Dwight S. S., Eppig J. T., Harris M. A., Hill D. P., Issel-Tarver L., Kasarskis A., Lewis S., Matese J. C., Richardson J. E., Ringwald M., Rubin G. M., and Sherlock G. 2000. Gene ontology: tool for the unification of biology. The Gene Ontology Consortium. Nat Genet 25: 25-29.
- Baird N. A., Etter P. D., Atwood T. S., Currey M. C., Shiver A. L., Lewis Z. A., Selker E. U., Cresko W. A., and Johnson E. A. 2008. Rapid SNP discovery and genetic mapping using sequenced RAD markers. PLoS One 3: e3376.
- Bairoch A. 2000. The ENZYME database in 2000. Nucleic Acids Res 28: 304-305.
- Bairoch A., Apweiler R., Wu C. H., Barker W. C., Boeckmann B., Ferro S., Gasteiger E., Huang H., Lopez R., Magrane M., Martin M. J., Natale D. A., O'Donovan C., Redaschi N., and Yeh L. S. 2005. The Universal Protein Resource (UniProt). Nucleic Acids Res 33: D154-159.
- Baranzini S. E., Mudge J., van Velkinburgh J. C., Khankhanian P., Khrebtukova I., Miller N. A., Zhang L., Farmer A. D., Bell C. J., Kim R. W., May G. D., Woodward J. E., Caillier S. J., McElroy J. P., Gomez R., Pando M. J., Clendenen L. E., Ganusova E. E., Schilkey F. D., Ramaraj T., Khan O. A., Huntley J. J., Luo S., Kwok P. Y., Wu T. D., Schroth G. P., Oksenberg J. R., Hauser S. L., and Kingsmore S. F. 2010. Genome, epigenome and RNA sequences of monozygotic twins discordant for multiple sclerosis. Nature 464: 1351-1356.
- Bateman A., Coin L., Durbin R., Finn R. D., Hollich V., Griffiths†Jones S., Khanna A., Marshall M., Moxon S., and Sonnhammer E. L. L. 2004. The Pfam protein families database. Nucleic acids research 32: D138-D141.
- Baxter L., Tripathy S., Ishaque N., Boot N., Cabral A., Kemen E., Thines M., Ah-Fong A., Anderson R., and Badejoko W. 2010. Signatures of adaptation to obligate biotrophy in the *Hyaloperonospora arabidopsidis* genome. Science 330: 1549.
- Begun D. J., Holloway A. K., Stevens K., Hillier L. W., Poh Y. P., Hahn M. W., Nista P. M., Jones C. D., Kern A. D., Dewey C. N., Pachter L., Myers E., and Langley C. H. 2007. Population genomics: whole-genome analysis of polymorphism and divergence in *Drosophila simulans*. PLoS Biol 5: e310.
- Bell C. J., Dinwiddie D. L., Miller N. A., Hateley S. L., Ganusova E. E., Mudge J., Langley R. J., Zhang L., Lee C. C., Schilkey F. D., Sheth V., Woodward J. E., Peckham H. E., Schroth G. P., Kim R. W., and Kingsmore S. F. 2011. Carrier testing for severe childhood recessive diseases by next-generation sequencing. Sci Transl Med 3: 65ra64.
- Birney E., Clamp M., and Durbin R. 2004. GeneWise and Genomewise. Genome Res 14: 988-995.
- Boutemy L. S., King S. R., Win J., Hughes R. K., Clarke T. A., Blumenschein T. M., Kamoun S., and Banfield M. J. 2011. Structures of Phytophthora RXLR effector proteins: a conserved but adaptable fold underpins functional diversity. J Biol Chem 286: 35834-35842.

- Chamnanpunt J., Shan W. X., and Tyler B. M. 2001. High frequency mitotic gene conversion in genetic hybrids of the oomycete *Phytophthora sojae*. Proc Natl Acad Sci U S A 98: 14530-14535.
- Cvitanich C., Salcido M., and Judelson H. S. 2006. Concerted evolution of a tandemly arrayed family of mating-specific genes in *Phytophthora* analyzed through inter- and intraspecific comparisons. Mol Genet Genomics 275: 169-184.
- Dou D., Kale S. D., Wang X., Chen Y., Wang Q., Jiang R. H., Arredondo F. D., Anderson R. G., Thakur P. B., McDowell J. M., Wang Y., and Tyler B. M. 2008. Conserved C-terminal motifs required for avirulence and suppression of cell death by *Phytophthora sojae* effector Avr1b. Plant Cell 20: 1118-1133.
- Erwin D. C.and Ribeiro O. K. 1996. *Phytophthora* diseases worldwide. The American Phytopathological Society, St. Paul, MN.
- Forche A., Abbey D., Pisithkul T., Weinzierl M. A., Ringstrom T., Bruck D., Petersen K., and Berman J. 2011. Stress alters rates and types of loss of heterozygosity in *Candida albicans*. MBio 2.
- Gevens A. J., Donahoo R. S., Lamour K. H., and Hausbeck M. K. 2008. Characterization of *Phytophthora capsici* causing foliar and pod blight of snap bean in Michigan. Plant Dis. 92: 201-209.
- Gobena D., Roig J., Galmarini C., Hulvey J., and Lamour K. H. 2012. Genetic diversity of *Phytophthora capsici* isolates from pepper and pumpkin in Argentina. Mycologia 104: 102-107.
- Grigoriev I. V., Nordberg H., Shabalov I., Aerts A., Cantor M., Goodstein D., Kuo A., Minovitsky S., Nikitin R., Ohm R. A., Otillar R., Poliakov A., Ratnere I., Riley R., Smirnova T., Rokhsar D., and Dubchak I. 2012. The genome portal of the department of energy joint genome institute. Nucleic Acids Res. 40: D26-32.
- Haas B. J., Kamoun S., Zody M. C., Jiang R. H., Handsaker R. E., Cano L. M., Grabherr M., Kodira C. D., Raffaele S., Torto-Alalibo T., Bozkurt T. O., Ah-Fong A. M., Alvarado L., Anderson V. L., Armstrong M. R., Avrova A., Baxter L., Beynon J., Boevink P. C., Bollmann S. R., Bos J. I., Bulone V., Cai G., Cakir C., Carrington J. C., Chawner M., Conti L., Costanzo S., Ewan R., Fahlgren N., Fischbach M. A., Fugelstad J., Gilroy E. M., Gnerre S., Green P. J., Grenville-Briggs L. J., Griffith J., Grunwald N. J., Horn K., Horner N. R., Hu C. H., Huitema E., Jeong D. H., Jones A. M., Jones J. D., Jones R. W., Karlsson E. K., Kunjeti S. G., Lamour K., Liu Z., Ma L., Maclean D., Chibucos M. C., McDonald H., McWalters J., Meijer H. J., Morgan W., Morris P. F., Munro C. A., O'Neill K., Ospina-Giraldo M., Pinzon A., Pritchard L., Ramsahoye B., Ren Q., Restrepo S., Roy S., Sadanandom A., Savidor A., Schornack S., Schwartz D. C., Schumann U. D., Schwessinger B., Seyer L., Sharpe T., Silvar C., Song J., Studholme D. J., Sykes S., Thines M., van de Vondervoort P. J., Phuntumart V., Wawra S., Weide R., Win J., Young C., Zhou S., Fry W., Meyers B. C., van West P., Ristaino J., Govers F., Birch P. R., Whisson S. C., Judelson H. S., and Nusbaum C. 2009. Genome sequence and analysis of the Irish potato famine pathogen *Phytophthora infestans*. Nature 461: 393-398.
- Hausbeck M. K.and Lamour K. H. 2004. *Phytophthora capsici* on vegetable crops: Research progress and management challenges. Plant Dis. 88: 1292-1303.
- Hohenlohe P. A., Bassham S., Etter P. D., Stiffler N., Johnson E. A., and Cresko W. A. 2010. Population genomics of parallel adaptation in threespine stickleback using sequenced RAD tags. PLoS Genet. 6: e1000862.
- Hulvey J., Young J., Finley L., and Lamour K. 2010. Loss of heterozygosity in *Phytophthora capsici* after N-ethyl-nitrosourea mutagenesis. Mycologia 102: 27-32.
- Hurtado-Gonzales O., Aragon-Caballero L., Apaza-Tapia W., Donahoo R., and Lamour K. 2008. Survival and spread of *Phytophthora capsici* in coastal Peru. Phytopathology 98: 688-694.
- Hurtado-Gonzales O. P. and Lamour K. H. 2009. Evidence for inbreeding and apomixis in close crosses of *Phytophthora capsici*. Plant Path. 58: 715-722.
- Jaffe D. B., Butler J., Gnerre S., Mauceli E., Lindblad-Toh K., Mesirov J. P., Zody M. C., and Lander E. S. 2003. Whole-genome sequence assembly for mammalian genomes: Arachne 2. Genome Res. 13: 91-96.

- Jeffries T. W., Grigoriev I. V., Grimwood J., Laplaza J. M., Aerts A., Salamov A., Schmutz J., Lindquist E., Dehal P., Shapiro H., Jin Y. S., Passoth V., and Richardson P. M. 2007. Genome sequence of the lignocellulose-bioconverting and xylose-fermenting yeast *Pichia stipitis*. Nat. Biotechnol. 25: 319-326.
- Jiang R. H., Tripathy S., Govers F., and Tyler B. M. 2008. RXLR effector reservoir in two *Phytophthora* species is dominated by a single rapidly evolving superfamily with more than 700 members. Proc. Natl. Acad. Sci. U S A 105: 4874-4879.
- Judd S. R.and Petes T. D. 1988. Physical lengths of meiotic and mitotic gene conversion tracts in *Saccharomyces cerevisiae*. Genetics 118: 401-410.
- Kemen E., Gardiner A., Schultz-Larsen T., Kemen A. C., Balmuth A. L., Robert-Seilaniantz A., Bailey K., Holub E., Studholme D. J., Maclean D., and Jones J. D. 2011. Gene gain and loss during evolution of obligate parasitism in the white rust pathogen of *Arabidopsis thaliana*. PLoS Biol. 9: e1001094.
- Kim J. I., Ju Y. S., Park H., Kim S., Lee S., Yi J. H., Mudge J., Miller N. A., Hong D., Bell C. J., Kim H. S., Chung I. S., Lee W. C., Lee J. S., Seo S. H., Yun J. Y., Woo H. N., Lee H., Suh D., Lee S., Kim H. J., Yavartanoo M., Kwak M., Zheng Y., Lee M. K., Park H., Kim J. Y., Gokcumen O., Mills R. E., Zaranek A. W., Thakuria J., Wu X., Kim R. W., Huntley J. J., Luo S., Schroth G. P., Wu T. D., Kim H., Yang K. S., Park W. Y., Kim H., Church G. M., Lee C., Kingsmore S. F., and Seo J. S. 2009. A highly annotated whole-genome sequence of a Korean individual. Nature 460: 1011-1015.
- Koonin E. V., Fedorova N. D., Jackson J. D., Jacobs A. R., Krylov D. M., Makarova K. S., Mazumder R., Mekhedov S. L., Nikolskaya A. N., Rao B. S., Rogozin I. B., Smirnov S., Sorokin A. V., Sverdlov A. V., Vasudevan S., Wolf Y. I., Yin J. J., and Natale D. A. 2004. A comprehensive evolutionary classification of proteins encoded in complete eukaryotic genomes. Genome Biol. 5: R7.
- Krogh A., Larsson B., von Heijne G., and Sonnhammer E. L. 2001. Predicting transmembrane protein topology with a hidden Markov model: application to complete genomes. J. Mol. Biol. 305: 567-580.
- Lamour K.and Kamoun S. 2009. Oomycete genetics and genomics: diversity, interactions, and research tools. Blackwell Pub, Hoboken, NJ.
- Lamour K. H.and Hausbeck M. K. 2000. Mefenoxam insensitivity and the sexual stage of *Phytophthora capsici* in Michigan cucurbit fields. Phytopathology 90: 396-400.
- Lamour K. H.and Hausbeck M. K. 2001. The dynamics of mefenoxam insensitivity in a recombining population of *Phytophthora capsici* characterized with Amplified Fragment Length Polymorphism markers. Phytopathology 91: 553-557.
- Lamour K. H., Stam R., Jupe J., and Huitema E. 2011. The oomycete broad-host-range pathogen *Phytophthora capsici*. Mol. Plant Pathol. (online at 10.1111/j.1364-3703.2011.00754.x).
- Leonian L. H. 1922. Stem and fruit blight of peppers caused by *Phytophthora capsici* sp. nov. Phytopathology 12: M08.
- Levesque C. A., Brouwer H., Cano L., Hamilton J. P., Holt C., Huitema E., Raffaele S., Robideau G. P., Thines M., and Win J. 2010. Genome sequence of the necrotrophic plant pathogen *Pythium ultimum* reveals original pathogenicity mechanisms and effector repertoire. Genome Bio. 11: R73.
- Li W.and Freudenberg J. 2009. Two-parameter characterization of chromosome-scale recombination rate. Genome Res. 19: 2300-2307.
- Meitz J. C., Linde C. C., Thompson A., Langenhoven S., and McLeod A. 2010. *Phytophthora capsici* on vegetable hosts in South Africa: distribution, host range and genetic diversity. Australasian Plant. Path. 39: 431-439.
- Miller N. A., Kingsmore S. F., Farmer A., Langley R. J., Mudge J., Crow J. A., Gonzalez A. J., Schilkey F. D., Kim R. J., van Velkinburgh J., May G. D., Black C. F., Myers M. K., Utsey J. P., Frost N. S., Sugarbaker D. J., Bueno R., Gullans S. R., Baxter S. M., Day S. W., and Retzel E. F. 2008. Management of high-throughput DNA sequencing projects: Alpheus. J. Comput. Sci. Syst. Biol. 1: 132.

- Morschhauser J. 2010. Regulation of white-opaque switching in *Candida albicans*. Med. Microbiol. Immunol. 199: 165-172.
- Narva E., Autio R., Rahkonen N., Kong L., Harrison N., Kitsberg D., Borghese L., Itskovitz-Eldor J., Rasool O., Dvorak P., Hovatta O., Otonkoski T., Tuuri T., Cui W., Brustle O., Baker D., Maltby E., Moore H. D., Benvenisty N., Andrews P. W., Yli-Harja O., and Lahesmaa R. 2010. High-resolution DNA analysis of human embryonic stem cell lines reveals culture-induced copy number changes and loss of heterozygosity. Nat. Biotechnol. 28: 371-377.
- Nielsen H., Brunak S., and von Heijne G. 1999. Machine learning approaches for the prediction of signal peptides and other protein sorting signals. Protein Eng. 12: 3-9.
- Ogata H., Goto S., Sato K., Fujibuchi W., Bono H., and Kanehisa M. 1999. KEGG: Kyoto Encyclopedia of Genes and Genomes. Nucleic Acids Res. 27: 29-34.
- Parra G., Bradnam K., and Korf I. 2007. CEGMA: a pipeline to accurately annotate core genes in eukaryotic genomes. Bioinformatics 23: 1061-1067.
- Raffaele S., Farrer R. A., Cano L. M., Studholme D. J., MacLean D., Thines M., Jiang R. H., Zody M. C., Kunjeti S. G., Donofrio N. M., Meyers B. C., Nusbaum C., and Kamoun S. 2010. Genome evolution following host jumps in the Irish potato famine pathogen lineage. Science 330: 1540-1543.
- Rosenberg S. M. 2011. Stress-induced loss of heterozygosity in *Candida*: a possible missing link in the ability to evolve. MBio 2.
- Salamov A. A.and Solovyev V. V. 2000. *Ab initio* gene finding in *Drosophila* genomic DNA. Genome Res. 10: 516-522.
- Spanu P. D., Abbott J. C., Amselem J., Burgis T. A., Soanes D. M., Stuber K., Ver Loren van Themaat E., Brown J. K., Butcher S. A., Gurr S. J., Lebrun M. H., Ridout C. J., Schulze-Lefert P., Talbot N. J., Ahmadinejad N., Ametz C., Barton G. R., Benjdia M., Bidzinski P., Bindschedler L. V., Both M., Brewer M. T., Cadle-Davidson L., Cadle-Davidson M. M., Collemare J., Cramer R., Frenkel O., Godfrey D., Harriman J., Hoede C., King B. C., Klages S., Kleemann J., Knoll D., Koti P. S., Kreplak J., Lopez-Ruiz F. J., Lu X., Maekawa T., Mahanil S., Micali C., Milgroom M. G., Montana G., Noir S., O'Connell R. J., Oberhaensli S., Parlange F., Pedersen C., Quesneville H., Reinhardt R., Rott M., Sacristan S., Schmidt S. M., Schon M., Skamnioti P., Sommer H., Stephens A., Takahara H., Thordal-Christensen H., Vigouroux M., Wessling R., Wicker T., and Panstruga R. 2010. Genome expansion and gene loss in powdery mildew fungi reveal tradeoffs in extreme parasitism. Science 330: 1543-1546.
- Tyler B. M., Tripathy S., Zhang X., Dehal P., Jiang R. H., Aerts A., Arredondo F. D., Baxter L., Bensasson D., Beynon J. L., Chapman J., Damasceno C. M., Dorrance A. E., Dou D., Dickerman A. W., Dubchak I. L., Garbelotto M., Gijzen M., Gordon S. G., Govers F., Grunwald N. J., Huang W., Ivors K. L., Jones R. W., Kamoun S., Krampis K., Lamour K. H., Lee M. K., McDonald W. H., Medina M., Meijer H. J., Nordberg E. K., Maclean D. J., Ospina-Giraldo M. D., Morris P. F., Phuntumart V., Putnam N. H., Rash S., Rose J. K., Sakihama Y., Salamov A. A., Savidor A., Scheuring C. F., Smith B. M., Sobral B. W., Terry A., Torto-Alalibo T. A., Win J., Xu Z., Zhang H., Grigoriev I. V., Rokhsar D. S., and Boore J. L. 2006. *Phytophthora* genome sequences uncover evolutionary origins and mechanisms of pathogenesis. Science 313: 1261-1266.
- van Ooijen J. 2011. Multipoint maximum likelihood mapping in a full-sib family of an outbreeding species. Genet. Res. (Camb) 93: 343-349.
- Vercauteren A., Boutet X., D'Hondt L., Bockstaele E. V., Maes M., Leus L., Chandelier A., and Heungens K. 2011. Aberrant genome size and instability of *Phytophthora ramorum* oospore progenies. Fungal Gen. Biol. 48: 537-543.
- Whisson S. C., Boevink P. C., Moleleki L., Avrova A. O., Morales J. G., Gilroy E. M., Armstrong M. R., Grouffaud S., Van West P., and Chapman S. 2007. A translocation signal for delivery of oomycete effector proteins into host plant cells. Nature 450: 115-118.

- Win J., Kanneganti T. D., Torto-Alalibo T., and Kamoun S. 2006. Computational and comparative analyses of 150 full-length cDNA sequences from the oomycete plant pathogen *Phytophthora infestans*. Fungal Genet. Biol. 43: 20-33.
- Wu T. D.and Nacu S. 2010. Fast and SNP-tolerant detection of complex variants and splicing in short reads. Bioinformatics 26: 873-881.
- Zdobnov E. M.and Apweiler R. 2001. InterProScan--an integration platform for the signature-recognition methods in InterPro. Bioinformatics 17: 847-848.

Figure 1: Life cycle. Thick walled dormant sexual oospores (**A**) are produced when plants are infected with both the A1 and A2 mating types. Asexual reproduction following infection rapidly creates very large populations of sporangia on the surface of the infected plants (**B-D**, **G**). Sporangia can detach and cause infection directly or, in the presence of free water, release 20-40 bi-flagellate swimming zoospores (**E**) that swim to plants and cause infection indirectly (**F**).

Figure 2: Synteny and genome architecture of *P. capsici*. Pairwise comparison of gene models from *P. capsici* to *P. ramorum* (**A**) and *P. sojae* (**B**). Sequences other than gene models were replaced with Ns. Maximal unique matches in all 6 frames were used as anchors for amino-acid-based alignment with the PROmer package of MUMmer. Scaffolds >450 kbp are shown (*P. capsici*: 46 Mb, *P. ramorum*: 36 Mb, *P. sojae*: 58 Mb). Scaffolds were ordered to maximize the center diagonal. There were some translocations, which possibly were assembly errors. A small region of *P. sojae* (top) had no syntenic genes in *P. capsici*. (**C**) *P. capsici* whole 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 binning. The color scale shows number of genes in bins. In addition, the 365 predicted proteins for which no homolog was found (10e-5 e-value cutoff) are indicated as dots.

Figure 3: The 1654cM *P. capsici* genetic map. The map contains 1,136 SNV markers and 18 linkage groups (LG's). Linkage groups are colored by genome scaffold.

Figure 4: Ploidy for LOH regions. (A) Normalized sequence coverage of RAD tags across 50,649 nt of a 850,537 nt LOH tract on scaffold 4 (LG10) at nt 86,143 - 936,679 encompassing the mating type locus. The 6 isolates with LOH are denoted by red dots. Normalized coverage was obtained by dividing the average coverage across the tract by the total unique coverage for each sample. Parents are in light blue and non-LOH progeny are dark blue. **(B)** Normalized coverage of tags across 14,143 nt of a

415,652 nt LOH tract on scaffold 35 (LG16) at nt 141,346 - 556,998. Five isolates with LOH are in red. TN58 had ½ of the expected coverage, indicating LOH to reflect a sub-chromosomal deletion. **(C)** Normalized coverage of tags across 22,124 nt f a 655,528 nt LOH tract on scaffold 1 (LG9) at nt 87,441 - 742,969. TN50 had LOH of this region.

Figure 5: LOH in *P. capsici.* **(A)** Graphical genotypes for LG10 in the region of scaffold 4 encompassing the mating type locus in parents (TN1 and TN2, black) and 59 F_1 progeny. R = reference allele and A = alternate allele. Red asterisks to the left of the marker names denote unambiguous LOH loci. Dots represent missing genotypes. Six progeny had LOH in this region (in red), including four that switched mating type from A2 to A1 and two that remained A1 following LOH. Group 1: progeny 10, 13, 15, 16, 26, 29, 31, 35, 37-41, 46, 51, 54, 59-61, 64-66, 70, 71, and 9. Group 2: progeny 14, 17, 19-24, 27, 30, 33, 36, 42-45, 49, 5, 55, 58, 62, 67, and 68. (B) Association of LOH with loss of pathogenicity of *P. capsici* isolates in wound inoculated jalapeno fruits between 2005 and 2011. TN1 and TN2 are the parents. TN31 was representative for all progeny except TN37 and TN47. In 2011, the TN37 and TN47 genomes had 10.4% and 11.1% LOH, respectively, and these isolates no longer infected wounded fruit.



Lesions on pepper, pumpkin and cucumber contain millions of sporangia.

Asexual reproduction rapidly creates very large spore populations on infected plants.

Figure 1



100

10

5' Flanking Intergenic Region (Kbp)

Figure 2

0.01

0.01

0.1



Figure 3_Lamour_MPMI



Figure 4



Figure 5

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 F_1 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 \geq 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*.



Phytophthora capsici LT1534

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 \geq 4 uniquely aligning reads of Q score \geq 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	с	Silent		e_gw1.2.451.1
2_1437339	2	1	0.41	0.00	0.00		1	1	3	1	с	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	1			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	1			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	с	Silent		e_gw1.2.451.1
2_1441385	1	2	0.59	0.00	0.00		1	1	1	1	с	Silent		e_gw1.2.451.1
2_1441919	3	1	0.00	0.00	0.00	Yes	3	3	3	1	с	Silent		e_gw1.2.451.1
2_1441939	1	2	0.89	0.00	0.00		1	1	1	1	с	Р	R	e_gw1.2.451.1
2_1441973	1	2	0.89	0.00	0.00		1	1	1	1	С	Silent		e_gw1.2.451.1
2_1443263	3	2	0.00	0.00	0.49		3	3	3	1	с	Silent		e_gw1.2.451.1
2_1447270	1	2	0.89	0.00	0.00		1	1	1	3	С	N	н	estExt2_fgenesh1_pg.C_PHYCAscaffold_20222
2_1452313	3	2	0.00	0.00	0.49		3	3	3	1	с	Silent		fgenesh2_kg.PHYCAscaffold_2_#_197_#_4100895:1
2_1452392	3	2	0.00	0.00	0.59		3	3	3	1	с	Silent		fgenesh2_kg.PHYCAscaffold_2_#_197_#_4100895:1
2_1452394	3	2	0.00	0.00	0.59		3	3	3	1	с	Silent		fgenesh2_kg.PHYCAscaffold_2_#_197_#_4100895:1
2_1456984	1	2	0.58	0.00	0.00		1	1	1	3				
2_1457008	1	2	0.68	0.00	0.00		1	1	1	3				
2_1482757	3	2	0.00	0.00	0.68		3	3	3	1				
2_1482805	2	2	0.00	0.69	0.00		1	1	3	1				
2_1482817	2	1	0.41	0.00	0.00		3	3	1	1				
2_1482847	1	2	0.59	0.00	0.00		1	1	1	1				
2_1482853	1	2	0.59	0.00	0.00		1	1	1	1				
2_1482862	1	2	0.59	0.00	0.00		1	1	1	1				
2_1484820	1	2	0.58	0.00	0.00		1	1	1	3				
2_1486427	3	2	0.00	0.00	0.58		3	3	3	1	С	Silent		estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1486428	3	2	0.00	0.00	0.58		3	3	3	1	С	Silent		estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1486435	2	2	0.00	0.62	0.00		1	1	3	1	С	М	к	estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1486458	3	2	0.00	0.00	0.58		3	3	3	1	С	А	т	estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1487541	2	1	0.33	0.00	0.00		3	3	1	1	I			estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1487551	2	1	0.33	0.00	0.00		3	3	1	1	I			estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1487569	2	1	0.33	0.00	0.00		3	3	1	1	I			estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1487581	2	1	0.33	0.00	0.00		3	3	1	1	I			estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1487584	2	1	0.33	0.00	0.00		3	3	1	1	I			estExt2_fgenesh1_pm.C_PHYCAscaffold_20199
2_1490736	2	2	0.01	0.71	0.00		1	1	3	1	С	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.



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 assemb	bly 97.1%

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

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%)
ab initio	2261 (11%)

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

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

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

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 5: RAD sequencing statistics.

Sample	Nt with ≥ 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

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

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	SNPs	RR·AR	AR·AR	A R · A A	A A · RR
Group	5141 5	KK,AK	AN,AN	AN.AA	
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

Linkage Group	Coding Silent	Coding Non- Silent	Intron	3'	5'	Non- Gene	Genes Per Linkage Group	Total Genes in Linkage Groups
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 10. Summary of genic and non-genic markers showing simple Mendelian inheritance.

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

Scaffold	Lowest	Highest	Total	Linkage
Block	Marker	Marker	Covered	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

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

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

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

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

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

Linkage				Tract
Group	Isolate	1st LOH	Last LOH	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

	Total LOH
Isolate	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 14. Summary of total nucleotides affected by LOH per isolate.

Organism	Source	Genome version	Reference
Phytophthora ramorum	http://genome.jgi.doe.gov/Phyra1_1/Phyra1 _1.download.ftp.html	1.0	Tyler et al. (2006)
Phytophthora sojae	http://genome.jgi- psf.org/Physo3/Physo3.home.html	3.0	Tyler et al. (2006)
Phytophthora capsici	http://genome.jgi- psf.org/PhycaF7/PhycaF7.home.html	11.0	
Phytophthora infestans	http://www.broadinstitute.org/annotation/ge nome/phytophthora_infestans/MultiDownlo ads.html	4.1	Haas et al. (2009)
Pythium ultimum	http://pythium.plantbiology.msu.edu/downl oad.html	Release 1	Levesque et al. (2010)
Hyaloperonospora arabidopsis	http://vmd.vbi.vt.edu/download/index.php	8.3.2	Baxter et al. (2010)

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

Gene Name	Amino Acid sequence
>jgil104199le_g	MVKLFCAIVGEAGSAFSVEVDETDSVDDLKTAIKAVNEDITCPPRKLQLFLAKKADGT
w1.9.417.1	WLDGAGAAAVTVDEADRVPLMLDKHGNHHKFVKMNPLLWIKNDQHFGENFRPVHV
	VVVVPDVAHAQTGLWLVTGFVKNALNTKGIRCKLYWMATLRIGYYDPARRTDKKNV
	AFWYEDTKLCFHVLFETKDAALLFETDLRIEPQTLGSPLTNQVVETRVAPVNAVSTELQ
	RVFYGDYVPDDSKSPQNSVSSISLTTSGSNLDSSTDEFRFQRIEHEKFFLPYGKAESCHLV
	SRKQSRNHKREFAKYDRDSNSRLALSRDMHGWFDGMSIEVPIVNMLPGSVKENQSIGN
	RRKVVEVFVKVLDAGCTDRVFSRLKEGSTTTNDPLMMKTFVHVEDPETFCLCMRWKH
	DDNAERWRSFWDMTPAVD*
>jgil106435le_g	MLKLFCVVVGVAGDAFPVNIEPNETVGDMKKKIKHEEMYQFPASELQLFLAKVPKEKH
w1.12.217.1	DMAWLSSRSEDVKKLKKGEKTPLIDILTEEDQELQAEDPLDDVLRGMDPPSLCQIHVLV
	MAPPQDSLRSHLTTLLSVLLCHVLTKAPTTPTDRNVDFKDDVCNFYGCYSPDESCVRC
	MLLNDAFPSELVVASHLFRCSNEDVSDVMMQITLSDIDDERNGLLLFKPLKYAFDHFQI
	SFIRDDTDVFRLKVFDPSILATPIVDLKDRKGNKVLSTEQTQLLLSRISENPCRFNTQTTF
	GDVDDSALTFTGLERPFYRCLNLQARVARVMALEKKWIDASYDFQDFWSEVSLDDKM
	EMFHRSILNS*
>jgil106457le_g	MKLFCAMVGMAVGIIEVDIDNNAYVTALRDAIATKNEEIKKSALRLKLFLAKKGNAWL
w1.12.537.1	TDTEAAGVGGDLESLGFKLMKSVRLLKNPEYFGEDFQTGEGHVHVLVVVPEENMTVG
	EPVVDVDGVNIYVTSNMTLNPPDLVAFWRAFQAIDTKIEADSVIALPEGTFILGNPKVGS
	RIYIRPCYPQLWEVCWHIIHHETPNLVILGNPGIGKTYFGYLLLLFLARLGKTVVYESRR
	TKRRFLFSRNVVIKGSQQDFDDILEQDTTYYVVDAMEPREFQARTILVTSPDRDVWYTF
	NKISCQTRYMPVWTEQEIFSCREQVYSTIPKSVVQKCFYRWGGIPRYVLQYAQFDNHQ
	ALIEKALEVVDFDWLMNAYGKLDDNNSQAHRLLHYRVNERFTCDYFGFASSFVQHEV
	YQHLHKKEKRKLLEFIGRSVGSGDLSVLRDRLAEEHDHHCARTHKKLKRF*
>jgil106668le_g	MVKLFCAIVGVEGSVFSVVIGEGQTVEELKKAIKERNDDKINVSWLGLQLFLAKKAKG
w1.12.123.1	DGDWLTEKDVQEGVYDMSDLQMLRAARAKLRLVKLSDNDVNEEQKVEGGKSVNVL
	VVLPGLTTTIKVNERENNALTDELAYYQRIGQEIQSNCQQHCGPILDKIDSIYEKKPYPM
	PFICVQGSSGMGKSQLAFALGGEGREHPRPWFYWTHGTVSDYDQRIYRNFASIGSAFDS
	VVWKDEVRKEEEDDILNCTSVLYMTKKLWTYGFIIELLRYCSRSNVGAQMVRVENQTF
	Y VI KSNLED VIEV RSRMEKNGE V LPFFILDEMPPSRTKKLSAFQLNVFRACGLVVIGMG
	TDANISNLVGKPEHSRTDPHWWMTVVSCFPPWQSIPFGDPAKEEVWQKVIELHPVVKH
	IAEHSKGLFSKCFVDAVVKFAMEEVSENEIFALADLLDAAFKAVYAELRIIKGFMFIEE
	GRDAQLMALSY HGSNKPPGI KKPV I DSDSI SEPI PKKKKLDVDVGVASMKAHFANSG
	YEGIAD VDVLQGDLKFKDISDAWSPICQFPAIEKDVLLILAVLGGKKFSDINDISSILD
	VFEEFLEKNUNKEESNALKKDFINFIKFENVVAHALFCASKKNUAKUALIEFLSULVSE
	FQDEY YQKDEFDASALLKEFDGLKEKFAEKKIPFLAPPNAKWPD YILEAGGDCNFGHFE
	SNWSLALVFCKKLEEFESEQKEDWEFPSTGIATVDCKAWEVNWISKPEAGEKLVIVVQT
Sigil10672612 g	
>jgi100/30le_g	
w1.12.192.1	VI VENELI VLOV V KOV MDANOFI I LUVADAKLKA VOLKSSEL VEV NEED VA VOKOH V H
	EDDHVVADETVSDOI SMESSEISVEDDSNDVEKVODEAEDI EGSHGKAESAHI ISASHOP
	NUTSVDEVDKDDNNDI ALSDEMUGAVDCINCDEDI VNIEVVSASDUDEI DUDEKVELO
	VSVVSHEVVELLGRI KDGSTKTEDDI VMKTEVVVODKNIECTCIOWKVNIVIDOL DEEE
	FTTNPR AT*
Sigil1086431e g	
w1 15 666 1	KGWI PDKSFAAI FI KKGEVHEDIOVI INGEEMEATKTI NVWI FEKNOMEEOI SSEOIH
w1.1J.000.1	VI VVVPFIRDGKKRYRSSI WFAFSFPI KKRRVKTDESEDEDDDVSFFRTEGI CCCODM
	AHPARKDSKI MERKAYTVIFAOI VEHVKDCEEENRTSNNPGI SSNVVVTGNPGIGKSW
1	

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

	FYLYCIFQLIRNREREDIKQLPPYELVMNYDDNFVKYDAARAEFVRLNKEDVDDLMDK
	PFVLRLVDARSTKLMGWRGVSVLFAVPDAEDLHDFEKVPGPKFIMPAWSLEELQDCNQ
	VLPDDLKLAEDELVSRFDAFGGIPRYVFSKNKTAIENKLKRAMASFSVKEILSYCKRGA
	AVKESDOSDCVLOMVPSEANFRLKFYLDFLSSDICEKIVFOAEGEDLTMLAKFAMGK*
>igil110274le_g	MVKFFCAIIGVAGSAFPVDIDASLSVGDLKNAIKGKNDDIKCTARELOLFLAKKEKGAG
w1.18.174.1	AWLTEDEAAAVSLDEGGHLOGFKMMKSSLYLKNPKHFGSNFOSDEDOVHVLVVVPD
	OAOPOTGI WI VTASVENAL NTKGIRCEL YELMASYI GYYDPVRETGDKDTAL WYED
	KTI CIHSIFKSFFNALLFDNVLODECITOTSPI DGHDVSTNVAPVSROLSFL RRIVSRHVA
	PODTESPOVSMI SISTNTSIVDVMTDEEKVORIESEEWEGDVGKAOSCHI MSRDHCRKC
	PSDPKVDNDPNNPI AI SSAMHDWVDGPMVNVPVMNISVESVSEPPVIGNPVKVNI IV
	PALNARVAKWISI II KEGEVASEDSI EMHTCVVVONDKVECVCMEWKRKEIDKOWKS
	VVDMEPAVD*
Sigil112102le g	MVKLECAIVGAMESAISVEVGEGI TVGDI KKAIATDOKEDEAASKI OLEI TETEGGGW
\sim Jg111219210_g	
w1.21.101.1	
	IAAI VENNKKKKEDDQEDE WWKAIEDDK VEIELELKEREQKEEF VKIELOUKE
	VIAK VLMY VSDUTCSKNSSVFTSTGLFKPNLCFYNSANDKVCVFKGEEQAKGEMTVPL
	IDLQKLIWKIDDAPIVFGIAAVAQDVCLVIIKESEINAPGEKKKAKVEIIEKINLKDL
	RGRLSIFLALLNLSTLFRPVVGRIRPLGIAEYKTIIRPNGVKIAFGENCVVKTYPLTMPSD
	KIISDLRDLHWQMKENAVPNVVELKSTNMRKRSVELAPVGRQLPLENVHQLLMAMRD
	ILRALVALHTIGLMHRDLRWENVLRYPDEDKWFLIDFDEGASSPAVKVDHLKAETHAP
	EILSSSTHTTMVDIWSVGYLLETSHVHDLPVALEDIKTQCLQENPSVRPTAQSLLEAVEA
	LIAN*
>jgil113302le_g	MVFASQDNDFNRCSNPFFSQLPTAEETDEWLEFPSLLPLTRRRSLYIRPSFKSIAAQALLK
w1.24.271.1	VDSNRRKYAVVTGTPGIGKSVFLYYVMWKLIKDKKRMLFVTGRPPIYFDGESVLDCYQ
	LPYAGNRNFWSPDLWCLVDATNPCKIAGLPIHHCSVLLASEPRDDYVRHFRKLVPTPQV
	FYMPIWTEEEMEKIIPLYPSAASVWRDRFETLGGIPRLVLQAVQTDPQEFMRVCFYSLEN
	CMRLVLFHSKTKTGFIDSLYTLEMPRHVHILSQEPYHEYTLAYASETAMRAVIDAKWIV
	NRAEMLHFLVMNLKSTDSLTQTTCHCIFELYAMQLLELGGTFSYRSLQAGVEQSSETLD
	EDENDIDIPMSWREIVDRVEADQNEDQLYVPKSAKDVAIDAWMPEVGGFQIALGKEQK
	IKSQAADELALLGQGGNRLFFLVFPRDFDSFTKQEPLSIEQYALLIPYPEV*
>jgil114043le_g	MVMVKIFCKIVGEAGSIFSVKIDDSESVEDFMRKIKDRCDGKIVAPWMDLQLFLAKKKD
w1.25.260.1	GVWLTKRDVLEVTSELDPLDDTLAPLNSVGLSEEDVRYRMTKEDIKAKKVPVHVLLVV
	PTEDYLRSPATILLETILPHVLTHATTTLTEDNRDFRHNLCNFYGCYTREQSLVRCMLLD
	VPLPKSLVLASHLFRRSNEYLSFRMMQISDIDDVKNGLLLFKPLKYAFDHFQISFIRDDT
	DVFRLKLFDSTIKETPLIDLVDHHGKKVLSEEQTGELLSDVNNDTCLFDVGKTFGDVDG
	CALAFTGIERPYYHCLNLQARVALMVALKKGWIDESYDFKDFWSEVSLDDKMEMFHR
	SILNSVADI*
>jgil117630le_g	MIKLFCAIVGEERNVIEVDISDVESISALKEAIKTKMPITLKYVDANTLQLFPAKMPDDK
w1.33.284.1	WLRDDSVAADDLMSGRIHDDIKAMIADSKILRPACRIRDELSAKKTGWKWSLPWSPITP
	KIKTKQIHVLVKVPDMMTGTLPLCGVAPEAQWRLWQSIVRVTSGTYSGTALVLHRSSR
	NLYLLTNLHFWLAEYEEEEIFFEHMSAGFKMNVELYLKRNPRKKRSGKQKSDGGELQQ
	TFNGSPVVIVDQLLPGHTTLTEVHSFVFKSDACWCSSVDYDYAVFKVLAPPPRIQLLGV
	EPSVSHFPTNVYVFGFHDGHEEKKFGHSYAIVPARIKYHRYKCLILSLSTVSLPESGVICT
	SRGLSIGYLAGSTIDESGNEVNQWLELGFFRRIL*
>jgil128631le_g	MVKLFCAIVGDPSWSFEVDIDENVSVSELNKAIKTKNKIKLKCVDANDLQLFLAKTDGP
w1.77.29.1	SGGKVWLDRAGAAAVETDDLHRFTRMDSTLYVRNPKHFGSGFKPDEGQVHVLVVVP
	KWAVEAETIGQNVERVGLPRTTALNDPKKYAEEIIALDDWGVNTVHQIPSIWEFMSSLS
	GCTKTGELFWRLEEKQVASLLLDGWFRESSPGSINQFEDMKSILMGSPGIGKSTLLCVM
	AFHLVLKYKKNVLVYRQLKGENCLLYLGYEDDEVVYFTVKRCKADRAVSIYEELGYR
	QGFPNVWLLLDGFRYKEIPEGLETFRMLATSQQVSLKSQESTDAYCCLLPCWSKKDLLS
	MGSLIYNFTPDEMEERFFYSGGSVREFTYATWEDIQRAMDVAVSGVEDYSKLLTTASC
	MFTDTSQVVRTFVENTNDRSHYFSSRYWEPMVDSEYAVLALSVRLKADALHRIYTWA
	KMAGHGSLAGCAFEIYLHRLAIDNRLEL
>jgil129711le_g	MMKLFCAVVGEAESVFHVDIEPGETVSDLKDAIKEINKHDPVLKNVTAMNLQLFLAKK

w1.86.23.1	GGAWLSGDDPAVLELEEGEIHPDILEMMDAKPMLDDKTLQFLLFEKNKLPQPSTNQIHV
	LVEIPAGARAIPYSNRPPKPFVSSEGATWDFQNPLDNEQLSNAIRKHYDAWKRGYYDKI
	LHPLFTCWSGPGTGKSRLLDEFPKLLKDWLFAGKSENPDMIRLLONAFTFNIAFDKETP
	HEAGSFSSAAELIGTRMLYQLQDTLKWDPFVQEKSRHSVPSDVMDKLSKILGTRHKDM
	CVILCVDGMEKLSHENGGKDCEFYKVLTVLSYLIGTSKCWVIAICSATIYSPVKNFLLSS
	POWAYEVPTAILSRPTVEGEDIFATFNGDQLIELLIDDMGGFGRALEVLHVMMRKARRK
	GSLEFMSVLTAVLAELRVLYPRIKKKMASMOEAFLAVVARRPVDKYSRFGKLSLDDVI
	STGLVRREGRFLTCPYVLYLLLDTPDSPWSKYKCYSSOETRENAKPWOTWEAFNYKLR
	ALKSVACOGKVDWRDIHRGARFGRGCYRVVIEEPRTYSLDVNRKTAKLDGFGEGNIFR
	CKYDPDOOGYFFEDAFTGVKDAESRAFHEIHOCKKIKDNLSLEDLLEEKKKAAGPHDL
	FLLYCTSEVEGDIESLENCAVVDRTCWEKYYGPFAARALYVSTVSPPDINTSAIOSLOLV
	NGIGPAISKRIVEKRPYSSLEEAHEKTGVSMNILSOTSCKRSKVDK*
>igil131973le_g	MKLFYVIVGVAGSPFPVDIEPSETVGDLTKAIKKENKCRFKHVDAYDLOLYLAKKDKG
w1.125.9.1	NGAWLTEDDVATVRDDAVFOTYKLMKPTLFLNNTEICGESINOCDVHVLVKAPMRLPT
	IHOL HYRHFTVGSVDIRTKKSMTFDPPPL VRFWRAL OGDRTEFKADAVLTL PEGTFLLG
	NPMLGSRIVIRHCYPRLWOVCLKMINDEAMNTPHLVILGNPGIGKTYFGYVILWLLARS
	GNTVVYESRVCHRRFLESODMVVOGSKKDFIEILEOTTTYYVVDGVEPRYYSAKTILLT
	SPOREVWYEENKDDCRSCYMPVWSRDEVI TCREI MYSDIPESVVODCERRWGGIPRYV
	I HYATAGGROOWI EKAMENITI DSI MDACGDI YENPSEESI HI I HYRVTKEENTDYE
	DFASOYVI FEVYRRI YNHNKKKI I FFIDKSGWVGAAAVI RDHI FEVYTVAASI TDED
	GDGVAVFOFKANKLI *
Sigil20879lfgene	MVKI FCAVVGVOGSAFPVDIDASOSVGDI KDAIKTKNKIKI KNIDASDI OLFLAKPKD
sh1 ng PHYCA	GPWI RSDDSDVIRMRSGAIPFOVKKI I NFOIDPAAGIGAI FGDAKPTMFIHVI VRVPDY
scaffold 75 # 8	DSDSEVNOORKI TSEOKI RKESGATGEI PVOGDEMKI EDI TDDDIGKVI DIKAIGDIVG
searroid_75_n_0	ETGSEEVIRKEII TSDOWSGSARI I VAMIAODNPNWKNIMGEI GVINCTOI SI NNDTKV
	STTVDASOFIARNNOLOVI PDSD*
\igi 30322 gw1	MVELSI OCAIVGOIGPSEDVEIDDGKKVSKI KEMIOVKNPETIKCDAKDI PLELAKMGP
×Jg11393221gw1.	PTTTOLI KIWKKESTDGNSPESEIPEEGEMVKI EVI NNMEIOEEOMEI PSTDOIHVI VVV
05.10.1	PPDDVGSKPSADIEVAKVI KPI KMIEDKVI TEEIDI DVDEKEEOI DNI SMVOOSDDDIVM
	TOTI LEEWKKEGEEDI VVEVDMEEVVEWKVIKKI LEGEDDVVIVGSDGVGKSCEI MI
Sigil506730lfgan	MVLVALTCALVCKTCVLCVKIDDSAOVWELKKAITVEKSNDLKDVDADKLELELAKM
>jgilo00/59ligeli	ECCTWI DCACAAAVAI DEDGUDOCCVOMDDTI WIKNSKUECDNEKDCECOVUVI VV
Acceptfold 21 #	
Ascalloid_ $21_\#$	KERKNEDIDKSMAWALOVI SLITI I ITTIKITEKLEKEW AAINEMIKQKNQDONEKI OTKDIN KKEKNEDIDKSMEVSSI SWNDI EDII SVEDEDI KASAVDENVVEAI EDEI MOVEKI VG
111#_Contigo	DVVSGREARDOVEIVALIEAVCI MI CDATIL VEEEVRGRNVI VHGDEEEVI RDGRNDVS
05.1	IVE AKODDIDOGIAONIVACI EAI SDVECI EDTI CIVTNVI EWVEISDDDEKIDDMNITTI K
	VYGAVDSIOELTEIVGMIVGELANSS*
Sigil511381lfgan	
>jgiij11301iigeii	DAWI DCACAAAVEI DEUCHDOCCVOMDTI WVKNDKUECDNEODCECOVUVI VVV
Acceptfold 82 #	
Ascallolu_ $83_{\#}$	DVDAVDETCCSDTDVEEEEWIKCDTEEOOSCDVDDVVEANICDVI DNNKI CVESVEKCA
9#_Conug727	
.1	ELIAMDEIVDESVMALLTNI TNIWEELWVSNKSNNDDILATTTI TTDGEAEEVIDTI LAO
	STADADIMI DELAEDVEDELINITIWELEW VSIKSINIKLIATITETTOEALEVIKTELAQ
	VADOVTDSIDTI SVEC*
Sigil527260lastE	
vt2 franceh1 =	
mC_PHVCAsee	
ffold 180005	Υ ΕΘΆΘΕΥ ΑΙΨΙΕΛΥΔΕΙΨΙ Ι ΕΛΙΑΕΙΝΛΚΑΚΙ Υ ΠΟΕΙΨΙΟΣΙΑΟΚΑΣΤΥΡΕΥΝΙΚΕΙ Υ Υ DI ΥΡΓΙ
11010_180093	
	ΓΙΝΓΙΟΠΨΥΓΓΨΥΟΟΚΑΝΑΝΠΙΓΓΑΥΟΟΟΑΠΙΝΓΕΛΙΟΓΑΥΟΛΑΕΙΝΤΑΓΥΓΕΕΥΛΚΚΚΑΓΑΕΥΓΕΟΥΟΕ Εξνιζνόε αιερυνοία ενι ζοριεμάζο ανανιουτοςιρινές νός *
Simil54007(1) (F	
>jg115409/6lestE	
xt2_Genewise1P	EIDVMEDVSDTTDLELLEAGKATLKSVGLSDEDVGEVDEADAAAGEGPVNVLVVIPEL

lus.C_PHYCAsc	YNVPSPVMALFNVMLPHVLTQVPTTESDVVIEFKMELCKFYECYSRHRTWVRCMLLD
affold_50927	VAFPKSLVSASHLFRCNNAFMAPLTVQLWDIDDMRNGLLLFKPLKHAFDHFQLSFILDD
	TNVFRLKLFDPSIYNTRLLDLKDCDNENVLSMEEMGVLFYNTSLTRNPCEFDTQTTFGD
	VDGSALVFTGLKRPFYRCLNQQARLARVFALKKHWIDESYNFTDFWSEVSLDDKMDM
	FHRSILEN*
>jgil556686lestE	MVLVALTCALVGKTGVLGVKIDDSAQVWELKKAIKEKSLNKLKNADADELELSLAKK
xt2_Genewise1P	GAGWLSIEDLAAIQKGEDVPGFERVSLVDTEDEAYSASIRDVLKTNGMPPPQTRQIHVL
lus.C_PHYCAsc	VVVPKDENDRSAMALDVPSLPPTTIHRHPERLKRWAAINEMIRQKNQEGNEKTSTRDT
affold_940014	NKKRKNRDIDSSMPYLSLSWTDLEPILTMEDFNLEASAVPQNVVEELRDRLMQVRKLY
	GDVYSGKEAKRQVFIMPIFEAVCLMLGDATILVEEDVKGKNVHVHGRFEFVLKHGKKR
	VSIVIAKRDDIPQGIAQNMAGLEALSDVEGLERTLGIVTNYLEWVFISDDDEKIRRMNTT
	LKVYGAVPSTKELREIVGMICGLLANST*
>jgil558527lestE	MMKLYCAIVGVAGSVFAVEIGEDKTVYDLKDAIKTQNKIKKVDAGDLQLFLAKKKKK
xt2_Genewise1.	GKGMWLTEKDVQKGVNNTSDFNLLGTVGAPLKFVGLLKDDVEFEPTLKDVESMNTPV
C_PHYCAscaff	HVLVAIPQQWTISKKTDAKRLEKEENIPLEMLWQYSEMEITTFPQPDELSSLLQRPLPFQ
old_11463	LNLQKFLTPKTIFDPSGPFLVCNELSALIDGFSYSCDYRPDPMASENTWQRMYDQLLDIS
	YRLCRAHGFDVVSNRN
>jgil559084lestE	MIWLYCAIVGKAGGVFGVKINKGDQVWELKKKIKDENQATITCDANELQLFLAKKDG
xt2_Genewise1.	MWLPDEDLVAVDLENGTIHPDIDKMMNAEQQMQDNKTLQFWLFEENEMPKPSTDQIH
C_PHYCAscaff	VLVVVPKQDGTLNETSVAQTPLQDEEASAYSFSELNSAMRDQIVRKMRLVENVPDVKE
old_21063	PEDTSIGGYSWIPKIEENEESQRAGYMAYLQQHLKTLIDRGDFLLDDIADDKSVLDIVDP
	RLPFAMSGTADVLLINRTSKNPLIKLAGVSLVIELKKKVEPDHVPQAIGQLVSCSMKAPL
	NCYPLSLLTDLNDRWHFSWFSDNHTLTQVTLKYPKNAFRLIEAAVLRRTESVSLPPSFIP
	GPFKIIKVDDFLLQPDDGYAEEMMERYELMADVVEPEFLMARRAEYAQHLVQSMPMY
	AHTFK*
>jgil563460lestE	MMKLFCAIVGARSAFSVEVGEDQTVEDLKLAIKNQNRNKLMSVDANDLQLFLAKKDK
xt2_Genewise1.	GNGMEWLTQLDVVKGVMDANGFAHLLFVDAKLRAVGLDSSELGKVNDKDVAVGKG
C_PHYCAscaff	HVHVLVVVPKTGVSKDGGRSIQKFSAPIEFSEMEHEGGVPSTFRTERKDTVAAIASLLDR
old_120265	IPVVFFRAPPLSGKTAMCHLLYNHIVFSKPDALVASVRANRMARNETFAEYFKKMYGC
	DFEEFCAYRCDRVLLIDEAQITYNDEQLWRGFVKDTLESQIPGLRLVLFSSYGSFDVYRK
	QERPGTPILVPTDNTFGLNVTPSKPGLQLSRVELEEMVLNSIGASVSDLIWVLCSGHIGIA
	RAVLVFLRWKFGSTTPNAEDVEMELRSEELLQYVRASYRGIPTADAFQRIVKNNDLSEE
	TILKMSEVLNGVASGKVTSLHDPDGQQTPRSQTAVELLTKFGFLYEDQAKQLQFASNM
	HLKIWLLSSRTDPIGYMVKDISHGDFIVACVQRMSASRLQKFATENTTRVARERQIQME
	LYGATTSCLPKGVLVTPEWRTDDGKGFIDLVIRGSGILWFWELLVNGDDAVCHSKRFE
	TGGTNYGSLTRNCRYMLIDFRQNMGVRKRKDGFLYVSFADSFTKAHVFGLDKPTVSVE
	LLS*
>jgil567378lestE	MVNLFCIIVGVAGNAFEVKIDDGASVAALKKEIKQENNIKLKGVDAGDLQLFLAKKDG
xt2_Genewise1.	AWLMSKDLLRMWNEETPEEDERDYMSERLDDPTVRIKEKFPSEYPDRSIHVLVQVPRE
C_PHYCAscaff	LLYNQPKPRDSSNEWLAEFFNHKVEPRSLPFVGKLSSFVTQPLPAKIRVKQEWLNEWAL
old_240430	SPGLQEKMFVLDDDAPCMEFTSLIFNKRTLNPFRRGKTENAFISMWDSIFRNVLDVLFT
	QAHIDRDSCNGSSTRQKRPDFLFVLDQVCVFRGEEKPPDVNISVPTEELCSKLVWAYGS
	VPYVFGYAASGYDIQLHALCPLDPLNVGGVVTKNIGTFNLEVKEHLFQIVLVMLNLSLL
	FQAIADECPASGRDEFRDITRSSGWPSKPLQTMLISGKNFSTNVGDKRTTTLKRASVPNV
	DRLTNLHFQKRLAVFKPRGTMVRPSNLLDLFGALKDVLQALVALHRLGWIHRDIRWSN
	VIRQRTGNSWFLIDFVDAATNPQQYPSGQHLSVEEHAPEIFVENGVHTTAVDIWAVGFL
	IETSGVEWLDFAGRTSLYRRLIAKDPAARPNAEEVLAELKALEEAAKSEKEACENSRRS
	ETQCRKRKLPDS*
>jgil570403lestE	MEVVKLFCVIVGEAGSAFSVKVGQDQEVDDLKEAIKDRSDGKIDVPRPDLQLFLAKNG
xt2_Genewise1.	NAWLSSDNDDVKALKEGVKTTLIDELTQKEKELQGESGLKKVLAGMLTPSTDQIHVLV
C_PHYCAscaff	VIPDQSKSAPSVSFEGVLDRCRDSFFLQLPTAGEDDSWLMFPQPLPLTERQKLYIRSSYK
old_370147	SIAAQALSKMDPKRRKYAVVTGTPGVGKSVFLFYVMWKLIKEKKRVLLMAEEPAIYFD
	GESMWEIQQLPYSGNRTFWSVDLWCLVDSVDPTTIAGFPIRKCCVLLASTPRRDCIGEF
	NKLEPTPDVFYMPLWTKEELSTIAPLYPNAQDQWENRFEGLGGVPRLVLKDLKVTPQE
	LLQTACSNCCLDGDQFEDQNGSDSDSHPQP*

>jgil573037lestE	MVTLFCAIVGIESDAFAVEVNENDSVYALKQAIHARKMYEFSADKLLLFPAKTEGPAW
xt2 Genewise1.	LSNSSEDVVKLEEGEKTPLIEALTTKCQELQAELPVSDMLKEIDPPSLSQIHVLVVLPDLK
C PHYCAscaff	KRKRGGTTRLSDLLETCSKEGSLPTEGDFLQMFEWDDQDCGKVKDITAIGDIVGFTGFR
old_510047	FFVRKEILCVLENLKHFKANFDRGEVGDQFIFLGSPGTGKSCVLALLCFYVAATSDHPV
	LWYRSVQYGREMSFTCLFYQKKYYRWNGAEVKIYDRLYDEV
>jgil573801lestE	MIKLFCALVGAQGSAFPVDIDASQSIGDLKDAIKDQKQNDLKNVDADKLQLFLAKKGD
xt2_Genewise1.	GWLASKDLPSIQRDMASPTIFEKLPLVDPTCSIQEVLTENELPDPQTRQIHVLVNVPQVH
C_PHYCAscaff	VLGKRTRADEWFINQIGMKKSRVIVDGNETDRITRYFEMAGFPPLAHPKAEYRKILERN
old_550284	AYIVIFTELMKKAKLSFEKGADCSLVVTGNPGIGKSRFYLYCIFTLFFALTWKLRSSPPSI
>jgil96357le_gw	MIKLFCVFVGAQGSAFPVDIDASQFVGDLKYAIVEKKKEDPNLKSVTAKNLQLFQTKTE
1.1.1010.1	NGDGWLSSDDGAVIAMRTGAIPEQVKKLLKDEMDPAEQIGDKFRNAPTKKTIHVLVVV
	PPAPENERKRKRMEDEVAPDAWIKAIKDEPVTTLPTCEGLKHHLLRALHVKIPINNRLF
	QIVSAQNSTGELFTVLEKLFEPQPRNVSDITGAVLRPIIDPLLPSGPTTKSSYHHFWDCVI
	ATLLKVVTDGNYHRSTNASASTGAYRPDMCFYSRKSNICVFRGEEKANGELDVPMAEL
	HEKLTWRYDDAPYIFGYAAVGLLVCLVTIQKDEKTSSRAKAEKIETYDLGNLKDRLLFL
	LALLNLSTLFDPVVDLIRPLGIPEYITRERTNGVRIEFAEDCVIKTYPKNMPSDGIIRNLKS
	LHRLMKEHSVPNVVELKNANKKKKHVKLAPIGIDRRPVNVDQLLMALCDILKALVAL
	HAINVMHRDLRWENVLKYSTEGDKWFLIDFDEGRLSPAATVTHLKAESHAPEILSSSSH
	TVKVDIWSVGYLLKTCCLQDLPPELKRIQSQCLQTDPSSRPTAKSLLAGIESLIES*
>jgil100678le_g	MVKDIKLRCGAYGEGSVFSVKIKQNADVEALQLAIVNARKGVNNRFNVDPSTLTLYLA
w1.5.1392.1	GKQEGEEIKWLKDEDSLGDELRGVYPKQYMKMRSSRILDEDYFGENFQPGRHDVHVL
	VELPEKVSSVQSVVRAVFWLVTGLVENALQTRGVHRLIYRIADAQLGYYDPANMLPDN
	KPRAFWYTNNDLQFHVLFKEGECVHCTCWCC*
>jgil102004le_g	CVIGVAGNAFSVNIDENLSVGHLKKAIKGENVNDPTLKNVAAKNLQLFLAKAEGGARL
w1.6.669.1	SSLTDEASEDATKVKKGEWLSNLTDDVKKLKKGEKTPLVESLTHENKGLQGESGFKRV
	LGPFLAKRKEWIWL*
>jgil106201le_g	IKLSLQCAIVGQTGSSFDVEIDDGEKVSKLKEMITEKNKQDPNLKNVAAKNLQLFLAKK
w1.12.351.1	GDAWLPDDDPAAQDLEEGKIHTEIKALIDGNKMKEAWTIEDVLVDNNMTGEGRAPKS
	RQIHVLAVVPGILTTIERERVDENQD*
>jgil108281le_g	MKKVSLQCVIVGIGSSFDVEIDDGEKVSKLMRAIKDRKPLTITCEADLLQLFLAKKGNA
w1.15.479.1	WLSSSTDDVKALKKGEKTGFIDELMHEKEKMEEEYPLSDYLANMNDPEVKQIHVLVV
	VP
>jgil108895le_g	MAKLSLQCAIVGQIGSSFDVEIDDGEKVSKLKDAIKTKNKDDPILKTVAAKNLQLFLAK
w1.16.149.1	QGNAWLPDDDPAAQDLNEGKVHTEIQALIPDEHRAALKLVNGESDDYINALTAGEQIL
	ASKTIETWLYEKIKMEEPSTAQTHVLVVVP
>jgil109378le_g	MKLTLQCAIVGQTGSSFDVKIEEGQTVGDLKEAIAVDQKFGFAASKLQLFLAKQPVEDD
w1.16.566.1	DGKEVVPVYHPCAEGMKKESFKWLPDKHRAALKLVKGESDDYINSLTAGEPILASKTL
	TIWLYEKNNMEDPSTQQFHVLVVVPE
>jgil113678le_g	MNRLERYAALNEIVQEKNQAENGKTSNQDTNHKRKKKALDNSVPFSSLSWDEIEPVLQ
w1.24.344.1	LNMFHLTAKPVPDEFVRKILAQLADLHQLYGDVSTGKEEKRKMFTMTVLEAVCLHLG
	DVMIFVDEELTGTKIHMHGSIEFVLQRGAKRVPIVIARRDNVEQGMAQCVACVEVLAD
	AEGLERTFGIVTNYLHWIFIRDEDESIELIDQPLNASMPSFESLKVILGMICGMLESE*
>jgil114206le_g	PLDDTLAPLNSVGLSEEDVRYRMTKEDIKAKKVPVHVLVVVPPADSPATILLETILPHVL
w1.25.261.1	THATTTFTEDNRDFTHNLCNFYGCYTREQSLVRCMLLDVPLPKSLVLASHLFRRSNEYL
	SFRMMQISDIDEVKNGLLLFKPLKYAFDHFQISFIRDDTDVFRLKLFDSTIKDTPLIDLSD
	RYGKKVLSEEQTGELVSVADNGSCLFDVGKTFGDVDGCALAFTGIERPYYRCLNLQAR
	VALMVALKKGWIDESYDFKDFWSEVSLDDKMEMFHRSILNSVAEF*
>jgil117855le_g	MVTIFCVVAGPGSVFAVDIGITQTIDHLKKQIKENKPNMIRFDADLLKLYLARDGGAWL
w1.34.341.1	NSNDDDFKALKRREVPARIKNLMQEQLLLDETAKLNDDDYFGKHFKPGDRDIHVLVEL
	PEDPTEVLHYKSELVCGDCFLAELLVVTNSFRL*
>jgil120859le_g	MVTVFCAIVGVPGSVFSVKIDENESVAELKKAIKKETPNIFQCNAMDLQLYLTKKGNV
w1.42.251.1	WLTEAHVKEGLRDTSGLKLLNSMKTKLKFLGLRDENDEEGEEEEGMGLVDVLV
>jgil122221le_g	MVTVFCAIVGVPGSVFSVKIDENESVAELKKAIKKENPNIFQCNAMDCLSDEGGRRVAD
w1.47.236.1	GGSCEGRLERHQWVEAVGFNEGETKSSGVEERKRRRR*
>jgil124124le_g	MGEELYDPTDKISAKFPSQIPGGTIHVLLVVPEGGKRHLSNEWFTESFHPLKKRRVGED

w1 52 167 1	VNKEKREEDMRDEPSI I EHPKVEEKTIVEREVYVVIESOI VOYAKTCEEEVPTSASEPDG
w1.52.107.1	KSKPGKDSNIVVTGNPGIGKSREELVCIEOLIL REREDVALL PPVELVVNHKTNVVKVDA
	VSKEEVELNKKDVD AL ODKDVVIDLVE ATSSELTGWDGVSVLEASDGVDGIDNESKVDG
	I TEIMOTWITEEEI EDVNSI I SDELKI AEDEI I SOVDOEGCIDDEVESOIMDOTEAKIOSAI
>jg1125925le_g	
w1.60.118.1	CDGLWLIENDVKNGVSSIAGLILLNAAQAPLQDAILGEGLHEPSKEDKVAGNGPVHV
	LVSVPESVG
>jg112/20/le_g	MMKLFCSVVGVAGSAFPVDIASDETVGDLKEAIKAKKMYCFPADELRLFLVNTSVKNP
w1.67.186.1	DEEEKKAH*
>jgil127255le_g	EDEDSSIDPYPWESDLAEDHEGQRKGYMQYLKDNLHGVLSEGGSTTIPPGTSTSGQPKY
w1.67.176.1	HLKDTSRMTSLLTCKASSLPFGLKGTADLMIIGEVAHSRNDIFADLQFVIKIKKNQCGPK
	ERKELLLELVAANWKSYSPCAPIGLLSNLNDYWYFMWFTTDRKIARMKLSCPANGLKA
	M*
>jgil127347le_g	MLNLLCAIVGAQGSVFPVLIGESESVGDLKKAIAKMKPLTVTCKADRLQLFLTKTEGGG
w1.68.101.1	WLSSKDEVVLATRKGGIPEEINKMLIDEIDPAKEIADVLGAAPTKMVIHVLVVVPRGLY
	TIHGDHVTARTDGWQSFSPTKLRLTRCRLLEV*
>jgil12764lfgene	MLLLNCAIIGGGDVISIIIEEWKTVALLKDAIKEKKPIKLNDVDAGDLHLFLAKKDGAWL
sh1_pg.PHYCA	MSDDLLQMREDGKGERGYMSEELKDPVAKISAKFPSELPDGSIHVLVVQRFRIEDVIDIL
scaffold_1_#_21	AELEKIDDVKNMTVILCVDALQQLVNDDAKTRNGQFRRSYHACSDDMAAI*
6	
>jgil128012le_g	MKVSLQYVVVGVAGSAFPIDIGKNLLVGHLKEAIKEKHDDIKCPARDLQLFLAKAGGN
w1.73.136.1	AWLANSTDHVKKLKKGEKTAYIEALIHENKELPRKDPISKYLERMDEPQMEQIHVLVV
	VP
>jgil128338le_g	MVKLSLQCAIVGQIGSSFDVEIDNGEKVSKLKEMIQVKNRETIKCDAKDLRLFLAKKGD
w1.75.57.1	AWLPDDDPAAQDLEEGKVHTAIQALIDGNKMKEAWTIADV*
>jgil128353le_g	MVKLFCAVVGVQGSAFPVDIDASLSVGDLKDAIKTKNKIKLKNIDASDLQLFLAKPKDG
w1.75.129.1	PWLRSDDSDVIRMRSGAIPEQVKKLLNEQIDPAAGIGALFGDAKPTMEIHVLVRVPDYD
	SDSEVNQQRKLTSFQKLRKESGATGELPVQGDFMKLFDLTDDDIGKVLNIKAIGDIVGF
	TGSDFYTFARPN*
>jgil128403le_g	GSAFPVDIDVSLSVGDLKKAINVEKTIKLKNVDAADLQLFMTKTKDGQWLRSDASDVI
w1.75.131.1	NMRSGVIPEQVKKLMNKWTRQMKLANCLVKNHQRRQFTCWWRLQRLI*
>jgil129581le_g	MVSITLCCVIVGVVGSEFDVNIAKGKSVHQLKEVIKAKNKKLLENVDTRELQLFLAKTA
w1.85.127.1	DGTWLSLTDHAVNSLRNGIIPTKVKALLKRE
>jgil129660le_g	LWNDEVIVAARIPREKVLMHERISRGGFGEVYVGVYNGRKVAIKMLLPEIRKRIQSVNE
w1.86.101.1	FLVEVKLMAALEHPRIVEFIGVAWDSLTDLCVVSELMERGDLRALLSQFQAENHPHGF
	DHDKVKIALHVAHALTYMHSFSPPIVHRDLKSKNILLTNEFDAKLTDFGASRERVDRTM
	TAGVGTSLWMAPEIMAGEKYDEKADMFSFAVVLSELDSHVLPYTKLRQETRASDVAIL
	QLVLQGKTQIDFSDACPSSIAALGMACAAKDPTARPTAAQALYELQQTLNSNEFY*
>jgil129685le_g	MVKLFCAIAGIAGSVFLVEIEEALEVDDLKNAIQKEKPLTITCDAHQLQLFLAKKDDGK
w1.86.155.1	GAWLTEVEVKNGVNDTTGLKPLDAVRAKLKNVQLSDSDVGGVDEADEVAGKGPVNV
	LVML
>jgil131615le_g	MKVSLQCAIVGQTGSSFDVEIDDSEKVSKLKKAIEKQEKISFKDVDAVDLHLFLAKVQK
w1.108.42.1	DMTWLYSRSEDVKKLKKGEKTPLIEALTMERHELQGEDPLENVLNGIDPPSVRQIHVLV
	VVPKG
>jgil131976le g	MDVRYESVAHYFKPINGTRVLKSIENCGEDFEPDEGRVHVLVVAPVTVLMLQDPPYCH
w1.125.17.1	FVVDGVNIPITDNMAFNLPGLTGFWKAFQEVDTEIEANTAIKLPEGTFLLGDSNRGSCIYI
	RSCYLQLWEITQKVVQDEVKKATNLVIDGNSGIGKTYFGYVMLLYLARLGETVVYESY
	GTKKRVLLSHNVVVEGSOODFSDILNLPTTFYIVDGVEPMHYOAKTIFLASLDHTLWYT
	FNEKRDQIRYIPVWSWDELSTCREVLYSDVPESVVEGCFHRWGGIPRYVLOYAOSEEK
	QILLEKTMEIADFFWLLNGYEKLKANNPEAHRLLHYRVNDHFAKEYFDFASPYVOOEV
	YDRAYKKDKRVLLGFIGGGDGWKW*
>jgil133238le g	MOLFLAKTEGKWLPDNEDLDTLLLTLHRICTCSHPTSWKLSNPDLFGPGVSLGEDVVH
w1.380.5.1	VLVVLKDAEAGVASVELSALPTVKQRHPERLKRWAAINEMVRQRNQERNEKTSTRDT

	NKKRKNRDIDCSIPYSNLSWVDLEPILPMEEDLKLEECRPSKCSSRST*
>jgil13996lfgene	MVKLSLQCAIVGQIGSSFDVEIDDGEKVSKLKEAIRTKKHLTITCEADQLQLFLAKQPVE
sh1 pg.PHYCA	GDDGKEVVPVYHSSAEEVKEESFKWLPDEHRAALNLVNGEPDDYINSLTVGKQILGSK
scaffold 5 # 21	AIATWLYTKNNMELPSNEQIHVLVVHHLTQFK*
0	
>jgil15135lfgene	MMHNLHGEAGNLKLTEALESPTRLVTDVVTQWLASMDEMRDEEAAPGRTSPKTQME
sh1 pg.PHYCA	KSKDEERDEEVQKKRPKRKFRKSTINVRKEEKAHLLEELSGIAYQDGGDQSSSICHÌHAL
scaffold 11 # 1	MSEYYLFNSPARGKMKTLTCAIVGVAGSAFPVDIDANKLVGHLKDAIKEKKMYOFPAD
21	ELOLFLAKKADGAWLSSKDPEVISMRSGDIPEOVKTLMNVEVDPTDDIEDVFEGAPTKK
	TVHVLVVERR*
>igil15224lfgene	MSDMVFLMLLFEHIVICVSLSSSIFFDAITEKNKKDPILKNVTAKNLKLFLAKTESGWLS
sh1 ng.PHYCA	YDEDLVNKLLLNRVDTSNTVTATIVDEEREKFDLLLSEKKGCLNVLFICATKYGDNLOE
scaffold 12 # 4	AFGPAKSVIYTEAMOSTEKKATTOYIHEVILLDLTTPONRAAFFGLAWDDTLOTRLENV
9	IHKAGGGNTAE*
>igil20945lfgene	MVOISLOCAIVGHAGSSEVVGIDDGAKVNKLKKAIORENPLTITCEADOLOLELAKDGK
sh1 ng.PHYCA	HFGSGFKPDEGOVHVLVVVPCAVAGSKAETIGONVEMNGDGEAGVDIFSL*
scaffold 77 # 1	
3	
>igil34764lgw1	MMKLECVIVGEAGSAEPVGIKPEDTVGDLKEKIKGKNTMTITCDAKDLOLELAKTHAG
83.9.1	WI SDNEDI DTI I OSEIDSSSYI RMRASWKI SKPNI EGPGVSI GEDVVHVI VVVPEG
5.5.1 Sigil49590lgw1	MI FLI CVIVGVTEWAFSVEVDETKSVDNI KGVIKNAKRNAI TGINASDI OI FTAKTTD
51 244 1	
51.244.1 Sigil502542lfgen	MVSSPI DSSTI DALADDEVDSMTWVDSITDDVVI ERDHSDGKSGEVLADDVEDMAEIDDEE
esh? kg PHVC	I EDPAEVPI SAMEMOEI EOVDEINEILAELDI MESHOELHYKI SEKTRELRSSSDVDAEL
Assaffold 1 #	VSMMARASKARNENQELEQVDENTELAEEDEMESTIQEETTIKESEKTKEEKSSSDVDAET
Ascallolu_1_#_ 2 # Contig261	I SIMINIAKASKAKINI SKULI VIILIIKINI SI HSKKININKSALIIQI KSVK
$2_#_Conug201.$	
1	
>jgilo03930ilgeli	
Assoffold 17 #	SALVDDAVAIOKOLSPDLSESEKLUIVIDKLSVNFULEILKVVPUIVSIEVDAKLSPDI
Ascallold_1/_# 12 # 1006622	
45#_4090055.	
2	ASTANIOEITELAOUDALTISTNLLEELTASTAALDAALTAOAATTOEALSTDEADT DI SMNEDAMATEVI AECIDCESADIVVI EOITVAVI SA*
Sigil509616lfgan	
>jgilo08010ligen	
esn2_kg.PHYC	GPWLKADNSDVISMKSGAIPEQVKKLMINEEMINPAAKIGDLFGDAKPIMEIHVLKVKSV
Ascarrold_36_#	55
69#_Cont1g46	
95.1	
>Jg1150894lgW1.	
/9.118.1	KGEGMWLIEKDVQKGVNDISDFNLLGIAGAPLKFVGLLKDDVEFKPILEDVESMNIP
1150006116	
>jgil509061lfgen	MGQGQSSIPTTEVAALVELYDALSGDRWRRRDGWKQPTRDPEQWFGVEVAMGHVVA
esh2_kg.PHYC	LELPANELSGCLPVASLARLPNLRVLDLSKNQLRGEIPAELGQLTALKRVDLSCNDLSG
Ascatfold_41_#	AIPRQIGACNQLQELNLYQNSLSGTMPKELGKLQSLRTLQLQHNNLCGALPETLCELTQ
57#_Contig18	LTKFSVRGNCLTGRIPTDIGRLQSLVFLSLRNNELTGVIPPSLGCCKALEFLNLSSNQLSG
45.1	PIPETLGELEDLEYLYLFDNALEGRYPGSIARLKFLKESDFRDNRLRGELPNFLDGCSSLE
	AVMTKWKNRKASYRHAILGDPMPSPDTPPTSSHQFLQTLEDPPSNSSATFLSQSFDHKG
5 1 1500 4 4 4 C	SEGLODUATEDDTANNSHLVAETSKKKVFQLTDSVVQAK*
>jg11509444lfgen	MFWSAALDLKIQKAPNKLSYGSFNKSASNSSFHIPQSIPEHQPVSQFTSPLPDKRREEYV
esh2_kg.PHYC	EKPDIAFAKIKASLQLSEHIQIALKMLLQGEKEFKKPSLGKTIENKPKPKFAQDMYSMS
Ascattold_46_#	KISSYSAHRAPMKELPEKTPLLKPIARAHGQEEPNPQENSNEHNYWMQELERRRAGLRR
29_#_Contig27	SKRKTCTPMSERATTSPLKVGVVMCSVPLVLVLGFVLFIFLFGGGSDASSNVMDVMNN
50.1	LLGIGRGDDAETIAARHAQDRDRIASTFARPPEKIQLEESIVMQLDVSSDHQKLRGSQKE
	KAGE*
>jgil509452lfgen	MESVVPCDLLDELMHGVVILHHEAALVLGTTQFSVETFFPLLAYVLVHCRLPIIHAQLH

esh2_kg.PHYC Ascaffold_46_# _37_#_4103400: 2	LLENFAITADNANGEESYYVYCVHAAVEYVCNAAGLSGSTNPLGPASSASSAVTTPGG CGLTPVAPMSSTSPPKQAAFSLDLELEMENELDVKVLGLSRDSGEEQPEESGTSALQ*
>jgil511091lfgen esh2_kg.PHYC Ascaffold_75_# _1_#_Contig233 .1	MKLFCAIVGVAGSAFSVEVGEDQTVDDLKDAIKVKNDDIKCPARELQLFLAKKDKG
>jgil52465lgw1. 36.563.1	MVELFCVIVGVTGSAFSVKIGNTKSVDHLKDVIKTKNKITLKDVDAPDLQLFLAKPKDG PWLRADNSDVIRMRSG
>jgil52481lgw1. 67.119.1	MAKLFCSVVGVTGSVFLVDIDQAESVRRLKKAIKAKKMYQFPSNELQLFLA
>jgil530721lestE xt2_fgenesh1_p m.C_PHYCAsca ffold_750003	MVKLFCAIVGVAGSAFPVTIDEGQSVGDLKEAIQVKNRETIKCDAKDLQLFLAKQPVEG DDGKEVVPVYHRSAEEMKEESFKWLPDEHRAALKLVKGESDNYINSLTAGEQILASKT LTTCKST*
>jgil532971lestE	MSISEACTCSLQARLHQVENARSREMKAHGTLVQSLEIELESRRICEGESRLALAKLRGV
xt2_fgenesh1_pg	NCKLEAELQAAHGKARQLLEQLEAQRVETSKCEERLKVFCEESDCQLRAAREKSDNLM
.C_PHYCAscaff	EEKKQLNVEVSEVKACLEKAKAENKVLHAKYEEKISEANAKTADIVQLTSQITDKQQEI
010_90133	NOFISHRIFLNWKLRSAOAKFOSAAAMARCNDDTRSRLOLIEMNVLAOCSSAKEELNR
	VRQACINECVQMQFLRREICQTELPKLLTIVRNHQRSFDNQQRHLEMQLVNQRESFERE
	RERLNMSRQDFEHCVQKAIEEKRLFQKRQHHAMFQVFTSKKKWELHRRVFGAWKEFY
	LRSIVGHATHTAMVFQSQHLRASTSEVSRPIRRGESHQWTMPAPIRRGESHQWTMPAPI
	RPISLASQHWRRIDRASS*
>Jg11533583lestE	
C PHYCAscaff	L TTKESOFFARVMVEOFENNRI TSOVAAMI EDKOOOFKETOTI MODNOSI ONKVDDI
old 150061	MOOVAGAROLOEDTEONYLEVASKI NOVLRENDELKNKPGTDASVELOSOLRAVETE
014_100001	LAASADKWROEOLOLEOOIETLTAVNKKNTTELADVADIREKLLVHVGIDLTYASIESL
	LDTKNNEIQMLTEQLAAAEAPQEQIIREADKTVEDLRAELERVAEEHSLSLVTMENEYR
	LACEELKTEAEKLTGQLDAAQTQTQQLNDDFSVERIHKSEEENRLLVEKLEECEAALKA
	RASEIEKLVARARTASSDTDKSGSSEMAACKAENLRMIFEVAKTADSVSKLKLDHEELL
	EAHRKKSSEVDTVLVQLASLESANQTLESDLKKKSEELHNHLETCSMQKEDFESVVAN
	LNNAVIKAEKEKSKIKRDLEDMKTENDVLEEQISELHAVAETKTEPDESQEKDREVEEL
	KSSLVQAKVDFLEQKQQLIALEKKLVAVSKSSGPICIVNSISLDAEKKEFEAALIEMIE MEKKLOVAVEAKOCI ESTLOEDMEAKTDIETDISIAEDKITELEOOLEOKVALIATIEE
	MEANLY VATEAN WILEST LYCENIEAN I DLETKLSTAEDNITELEVYLEYN VALIATIEL
	NEELEEOLNELADRIAEIEAEKODLLARLDETVYRSEEDIHOLRERLYMLEEEKSGLDDE
	NFKLERTIEHLESKLDSLEEOKAGLEAANEASSOOLSLLEERVEKATSEIATLSAEKDSL
	VELQQSLEENVSVLQEDKVKLEQTLDETSSKLRDELDRVTEQMESFQAQLAQSTAEKE
	EVTIALAELRERSEADKLASGEVTAKLEAQVAENQTLENKISVLKQMAEKALQSLQSSR
	DELSEAEARASVLVEERDAVKVLLQEKQTAYEQLKTQREELQLRVEQLSSELESSQKKR
	TEEAAIAEETIQTLKLTEAKLTESLESVKHDLAEAESAVMVLVEERDAARKEITARDLKI
	EMMISQUGELDLAAQKLESELSILRSKSSADSEATKEMLRDLEESKAQAEETVRSLQGS
	SAQAKESLQSIQEQLAESELKVAILEKEKDAIKISLNENMSIQEILSALQEDLQKKVDA
	ATKKTLSKRDLTLEVLSSEHGELOLKSOSTTTELENLRSKSAADAOAAFEALOSI REFI
	KRATESLESTQNELAESESHVAALTAEYDMVTKAMADVOSERDSLSGLNEELKESVED
	LKNEMENRQKQFSSELESAEEAIRTLKTSEVKMTASLDSVEQELCEAKSSSASLTAEYNS
	ILATQKEKEAGISKLTSEIDSLQERIQALESELQELRNQRETEAQTAEETIRSLKESESQTV
	ETLEAIRVKLSETEARVVSADEEREAASKALSEMESHRDAQSTEVESLQERIQALESELQ
	ELRNQRETEAQTAEETIRSLKESESQTVETLEAIRVKLSETEARVVSADEEREAASKALS
	EMESHRDAQSTEVESLQERIQALESELQELRNQRETEAQTAEETIRSLKESESQTVETLE
1	AIKVKLSEIEAKVVSADEEPLEIGSLVEKIVPLESELVELKNVKEIEAVAAEETIKSLKESE

r	
	SQIVETLEAIRVKLSETEARVVSADEEREAASKALSEMESHRDAQSTEVESLQERIQALE
	SELQELRNQRETEAQTAEETIRSLKESEAQKVVELDSVKKQVVLLSESSSSALTDLASLR
	EQWVAKDESHAADIASKDEVISTLKSKLEEVMAAYKRLKGHLHELQDRLTQQTSTNDT
	LKASYEELSGOKIAAVEELEALKIELASSSEHSNTMAEELROAEEKVAVAEAORESOME
	NEKKRVI AYDDELAOMOKOHAVALOEOKETI LKTSAVREFAAATKI LEOKHDOAGT
	CHEEASOKUEAEDMEI ESOI SSANETIEVI DI ASVTSVCI EVELAEI VI OVETETECANA
	SHEEASQNHEAENWELESQLSSANETHERLEASVTSVOLERLEALERLEVEVETETETEUAANA
	ARAALEI Y KKAAH ALKKASSENKLNLKKISEA I AKLEKELISAKSKVNI LE I ELED I K
	KRMAEVESAGDLLAQSTREAVESEKRSLEVTLRLEIDSLKAEVNRLEEALENDRQPLEA
	QINQLSERNAALNQDVITLKEEIRSQTESMEQEVHTKEEEIRDLSKQLQAALAAAASLAT
	NEAGRRSYSPTYSPTEKERRSTASSSRSFDSDGNNSFLHRSSIEEQSEHMAAAVADSCPIP
	LASKMAPANGVNKSQTDEDEVCRLKLQLNELETASHLFQKKYEDTSALLEEANQQKQ
	RLOELCDGSTOAINIEYLKNVIMKYIESOVPSEKEOLVPVISTLLSFTPOEOOKVMAVHR
	PNDEGAGLEGGVESLEGGGAAAAPPPKPLAAPLNEKPSPTTAMSNTTGAALGSKDKNG
	VI SEGSDPSDDEEFATPI NPFAA*
Sigil527102lastE	
>jgil557105lestE	WI VETCA V OF OUT VEDDWIT DI VEDLEK TRIKERKI NATI TOASLEKET LANGODK
xt2_fgenesn1_pg	WENSI DEAMKAVKARQCPDRIKNEMQEHELEDAMNFQIPDGVIHVEVEYPREDELDG
.C_PHYCAscaff	SFLENWFPCWSSLQSLFSVEKEKRA*
old_740004	
>jgil537210lestE	MRPDFLLHYLGMVLLRGEEKSATTEIDEPVKELTAKMSWWNPMFYGDLPYILGYATSG
xt2 fgenesh1 pg	ARLRVVMIDRHLHSDAILEFQSIFQQRAEVIKLFYNLAFFFHKMSVLAKRTCPSSLMPFT
.C PHYCAscaff	PDVNRKRKIELMDDVIVRTIKRNOCRDRVDFRRLADIYATLOELNNRVRERTHLOIVRK
old 790005	I RI KKRI I RVOI SPI GTVRPPMNVDEVRVWI OGMI TAI KYWHSCDYCHGDI RWSNI
014_790005	VVIDVSSDSGEWVI IDMDESPKSNTTTIDWNHEEOCYTI IEEHDI EOI GOI MNSETESI P
	SDIEDVOVALLTAVITDACDIDTILLIOUDS
	SDLEDVQKALLIAVHIRAGOLI HELQLIN
>jg1153/308lestE	MKLFCAIVGVAGSAFPVIIDEGQSVGDLKEAIQVKNREIIKCDAKDLQLFLAKQPVEGD
xt2_fgenesh1_pg	DGKEVVPVYHSSAEEVKEDSFKWLPDEHRAALKLVKGEYDDYINALTAREQILASKTI
.C_PHYCAscaff	ATWLYTKNNMELPSNEQIHVLVVQGKRIEDVIDILAKLEKVDDVKKMTVILCVDGFQK
old_830012	LVNDGEKTFGSTNADQETVDAVDRVCLNKKSKIAKIIRGMYSDLSDEENLFSEP*
>jgil537717lestE	MFEKTMKGFGTDEDALSATLVRYHGVLNDIRPVYKKKYGKELRDRPVYKKKYGKELR
xt2 fgenesh1 pg	DRIHGETSGKLLLALPTACAIVGQTGSSFDVEIDDSEKVSKLKKAIEKQEKISFKDVDAV
.C PHYCAscaff	DLOLFLAKVPKEKOGEVERTEEVKDEVOEDMTWLYSRSEDVKKLKKGEKTSLIEALTM
old 1080013	ERHELOGEDPLEDVL SGIDPPSVL OIHVL VVVPKGEDVVL A*
\igil5/1/030lestF	MPEDDALOTI DIVDEDGOHOOVU VGJEGGKOTVELANEVGGETTEEVTAVKTSEGEK
>JgllJ44950lcstE	
X12_Genewiserr	LVIEDAQQATVVEV HELFAEVAATVIEDQVVIAAVIADVQVQIVVEOEDEUOEQT
lus.C_PHYCAsc	VQLIDEHGDV1VEQAKAIE1SEGVKLEIKIDQGPVSVVVAEVPEEIKEEVIKEQEEHQPG
affold_160416	EFGGNPFTEEEPAVLQTVDVVADNGEHQQVVLVGGIEGGKQTVEIANEYGGFTTEEVT
	AVKTSEGEKLVIEDAQGQATVVEVHELPAEVAATVTEDSQVVNAVTADNQQVQIVVE
	GEPEHGEQTVQLIDEHGDVTVEQAKAIETSEGVKLEIKTDQGPVSVVVAEVPEEIKEEVI
	KEQEEHQPGEFGGNPFTEEEPAVLQTVDVVADNGEHQQVVLVGGIEGGKQTVEIANEY
	GGFTTEEVTAVKTSEGEKLVIEDAQGQATVVEVHELPAEVAATVTEDSQVVNAVTADN
	00V0IVVEGEPEHGE0TV0LIDEHGDVTVE0AKAIETSEGVKLEIKTD0GPVSVVVAEV
	PEEIKEEGPVSVVVAEVPEEIKEEAAEDSMOGSKFMTKPEFREWIRKHYEDRLESLKVEE
	OKLEDEERANAGRVKGLEDCIEOASNKEGYYGVYEOAPYYONAVDWVEEECWTAOO
	WEDESEVENTAGIA KOLLDELLQASIARI GITGATEQATTIQAAADA VLLLEA TAQQ
Sin:15/72071an4E	
>jg1154/38/lestE	MSEKLDDP I V KIKEKFPSEIPDRSIH V L V V V KREFL I NV PKPKDSSNEW LAEFFNHK V EP
xt2_Genewise1P	RSLPFVGNLSSFV1QPLPAKIRVKQEWLNEWALSPGLQEKMFVLDDDAPCMEFTSLIFN
lus.C_PHYCAsc	KRTLNPFRRGKTENAFISMWDSIFRNVLDVLFTQAHIDRDSCNGSSTRQKRPDFLFVLD
affold_240409	QVCVFRGEEKPPDVNISVPTEELCSKLVWAYGSVPYVFGYAASGYDIQLHALYPLDPLK
	VLTKNIGTFNLEVKEHLFQIVLVMLNLSLLFQAIADECPASGRDEFRDITRSSGVVVRLSP
	TFVEKIFPDISTFGHLELVYGHLKRASVPNVDRLTNLHFQKRLAVFKPRGTMVRPSNLL
	DLFGALKDVLQALVALHRLGWIHRDIRWSNVIRORTGNSWFLIDFVDAATDPOOYPSG
	OHLSVEEHAPEIFVENGVHTTAVDIWAVGFLIETSGVEWLDFAGRTSLYRRLIAKDPAA
	RPNAEEVLAELMALEESAKSEKEACENSRRSETOCRKRKLPDS*
Sigil563419lastE	
>Jg11303418lestE	
xt2_Genewise1.	KGNGVEWLTEKDVQGGVSDTSDLKLLARARARLRRVGLSGEDIVEVDEQIEAEGGGPV

C PHYCAscaff	NVLVELPPGTNRAPLSDGTDLWLSRFHHARISVLPTLGDLGEYIEHLSL*
old 120143	
>igil576078lestE	MVTIYCFAVDVEGSSFDVEIDDVAKVSKLKEAIATNOKFDFAASKLOLYLAKKGKGKG
xt2 Genewise1	VWI TEKDVOEGVSDTNDI KLI GAAGAPI NI VGI SEKDVKEEPTI EDVESMNTPVHVI
C PHYCAscaff	VVIPSPDVGPKR
old 830005	
>igil576081lestE	MKLECAIVGVAGSAESVEVGEDOTVDDSKLAIKNOKPNDLKDVDREKLOLELAKOPVE
xt? Genewise1	GDEGKEVVPVYHRSAFEMKEESEKWI PDEHRAALKI VKGESDDYINSI TAGEOILASK
C PHYCAscaff	TETTWI EKKNKMEI PSNEOIHVI VVVPDPSSNAAI EYKDPMSNI RKRGAEVMTDVVM
old 830013	EDDA EVA FENR VR VNR VR CIIVGIA GOSECINIESNAR VENI OGA KK SR CRORI KDVEA
014_050015	DDLI JELAEKADGAWI SADGAE
>igil577802lestE	MVELSI OCAIVGOIGSSEDVEIDDGKKVNKI KEMIOVKNRETIKCDAEDI RI ELAKKGN
xt2 Genewise1	AWLPDDDPAAODLEEGKVHTAIOALIDGEKMKEARTIADVLDDNNMTGEERAPKSROL
C PHYCAscaff	HVLVVVPOOWTVKRVTKPDDDI SRAKRARIVRI EVAYOGPVPTDFYSVPAFTIDTYOO
old 4740001	I KEAFI SHERI VRPI CI I YGPRREGKTTIGHRI VSI I DAEPSII VIYCSI TPI SVESEFAF
014_4740001	WVALGEFIGEHTRSFOFF
Sigil59630lgw1	DEMDSSVAPYAWMETNEAVHDORNAYMOVI KDNI HGVI HODGSTTTPHAIROPRYH
67 177 1	IKDTSHMKTI I NCTPEAI NHGI KGTADI MIVGETAHRMNDVETDI OI VVEVKKGOCK
07.177.1	A VEOOOI TI EI VA ANI KCEKRCAPIKI I TNI NDVWCEMWETPDSKKAPI TI SCPANGE
	KAIKDELAGAVDA
Sigil60673lgw1	MVKI SCAIVGVPRNVETVTIEDAASVSAI KEAIKTEKKNVMANEAAEDI OLELAKK
22.466.1	
>igil70314lgw1	MVKI SLOCAIVGOTGSSEDVEVGESOTVGDI KKAIKTKEPI TITCEADOLOLELAKOPV
16 564 1	GDEGGKEVVPVYHPSVEEMKEESMKWI PDEHRAALKI VKGESDDYINALTAGEOILAS
10.504.1	KPIATWI YTKNNMFI PSNFOIHVI VVVP
Sigil70333lgw1	MMI SI OCAIVGOTGSSEDVEIDDAEKVTKI KEAIOPKNROTIKCDAKDI OLYI AKKGN
32 357 1	AWI PDDDPAAODI NEGKVHTEIFAI INGNKMKFAWTIADVI VDNKMTGEGI APKSSO
52.557.1	IHVI VVVP
>igil71308lgw1	MKLECAVVGAOGSAFEVKIDNTESVSAFKEAIAAELRYKGRPDMLELELAKTESGWLS
16.631.1	NDDDLITKLLONRIDTSKLKALWPTWKLNRAKNPOLFGRDVSLGENVVHVLVRFP
>igil71310lgw1	MVKLFCAIVGVAGSAFSVEVDODOTVDDLKDAIKTKNKIKLKKVDASALOLFLAKKG
13.804.1	KAWLSNDKTGDTVLOSEDVASFEOMRGSWRLNDPKLEGTSVLLTEKVIHVLVVIP
>igil72559lgw1	LVMVSLOCVIVGKEECSENVKIDDGANVSKLKDVIKEOSGGLITVPPPTMOLELAKTGR
33 407 1	EGWLTGDDATAVVSDDI NHFTI MDPTI EVKDPKHEGEGEKPREGOVHVI VV
Sigil73233lgw1	MVKI ECAIIGEEGSTEPVDIDKGOTVGGI KKKIKEENEDDPTI KTVAAKNI OLEI TKTES
1 815 1	DRWLSSNDSVVIAMRKGDVPEOVKKI
>igil73236lgw1	MVKI FCTIIGDGSVFVVTIDEGOTVGDI KVAJAAAI SYTGRPDI LOLELAKTEKNAWLT
1 816 1	YDKDLVNKL
>igil73237lgw1	IMKLECAIVGVAGSAESVEVNEDOTVEDI KTAIADNOKENEANSTI OLELAKKHEGAW
12 600 1	I TOI
>igil80099.0	MVKI SI OCAIVGOTGSSEDVEVGEGOTVGDI KNAIKTKTI KTVAAKNI OLEI AKOPVG
10 758 1	DESCREVVPVYHPSVEEKKEESIKWI PDEHRAALKI VKGESDDVINALTAGEOILASKT
10.750.1	IATWI YTKNNMFI PSNFOIHVI VVVP
Sigil81603low1	
5 1157 1	NGKGVWI TEKDVONGVSDTSDI NI I GTMGAĐI KEVGI I EKDVKEEDTI KDIESMNTD
5.1157.1	VHVI VVVPKKRTSTAIVSFFKD
Sigil81615low1	MVKI FCMIVGEVGTAFSVDIADGDSVDDI KKAIAFNOKEGEAASKI OI EI SKKKNDTD
2 1184 1	FWI TEKDVOGGVIRDTSDI KPI VVAROII STATDVEVKIETKAEEAETSOVNVI VVVDE
2.1107.1	RDSTA
>igil83070law1	
-jg105070lgw1.	
12.944.1	

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	MKTATAFATVLALIVATNAAQVPPNTPTLRGLRPTADTPSVEDDKEDRKDHHHVKKVKKIAIPVPVPVEVPQ
4	FIPVPVSVPSTVVANSNNAVVGPSTNVAGPGAAAPGAPGAPGAVTPAPTTLNGRPAATPAATSTTRSRPTP
	APTNFAGARASGAQLPAAPVQAGAPGMTGFPNSAGVGDNTNNNAFGAGIGAAGGFPMAGNGLAGGFG
	GNPMNRFGGANGMGSGGIFGGAGTGPAMGNGFGGNFGQSMGFGMQDSNGLGGFGGQGMMTNFG
	QAGNNGFGGGQGGGGGGGFNRRERRL*
>scaffold_63_R3	MVNAMTMAIVLCTIVLRCSSRLRAALKQEVRVLRADAAQAAQTASNISSGLGTRLNALIARVSDLEKAFQAS
81	QASHHP*
>scaffold_22_F1	MGPLTGISLTFVACSRLGSWICSSAFLIISLDSTAAASWSRVASRQRNKKGNACMRISKLRLLRSLKTTKTMSY
619	PTS*
>scaffold_52_R1	MRLGLFLLVALAALFASCDASAARPVNDLPSGRFLRGESFKESLKILPKTYVKELVDDGDKLKAALQSWSDMK
150	LSRKRLAKSLELSTNRIKKLRKIANRHKVNEVFILDQYVKHLKQLKRSPRLPRVVE*
>scaffold_8_F31	MSLIWRILFVAAILLAGCNAFSDEAPSHLTLTESALMLRLIDTDPDHIGSRDTGKRFLRRDNDEALTDFDTEEG
04	KSSHPSAKMDKAVISPSIQGLSGQIPTALYNFLLKI*
>scaffold_438_R	MRLFPLFVAIATFLIATDAFLMTGDSNQISNVDSPGGSRQRLLRAHAEVDFEKEAKEMMRMMKNKVTKED
9	FAKKLKIAEEIDDIINKRAPGMHEFMQTVKYKRYANYMNFLNDMAKTSEYGDLVKEIKAKSRAQVALKTFRK
	PTTSQSRWQSILASLKGKRIGK*
>scaffold_87_F3	MKSIYRVLLLTVFALLCGLSNAVNSGGRLLRVVDSTDERGGFPSYLKDSFTKWRINSKIKSWVKKQKTDEYVLS
00	KLGLSTLTGKDLVKAPKYSQFQDFKVGMWLKEATPTTTVFNTLGLNKVEGAVEKADDFGTYVKYVMALGEK
	ADDYPLVTWRKLFGGGSLEQLELKRKILLLVKRDEIDISVMLG*
>scaffold_29_F3	MRFAYVVLAAAVAFVASTDSAVASEQAKLTSVVVAENSPLTHSLTAEQEGQRLLRAASDGESSFMERTKFYY
65	WYAMGRTPSYVYEDFFKGMDKSIIAENPNYKVWERYKAYYEKRKAN*
>scaffold 10 F6	MFCPLCVVYLPGLLLCSSDCIIPMYSSVQESAERLPLENRRRPYRNRPTSYITLEIPQGIGRPQVDRHLRLP*
08	
<pre>>scaffold_52_R8</pre>	MKILRSLTFLLIGTVAQAENSSKLLRGPDPNYDVAGLFPGPADTVVDTPRTKEFEPEVVDEPVIPSTLLNELDE
22	NSVERLVTEVTTYDDYTSEYNRHLRSVNDPAFQGQIQLAEPRHPKEHMIPPSPHDDQVPNLRGWT*
>scaffold_27_F8	MQIRVIALALSAALGSVAAVQTTDRSLGTLVGCPAVRSAKSPCLWAGENGEVVDSRALRELFLERHYVAHSD
34	REAYGRNLQEHMTYIEGEDDKAFIGLDTVY*
>scaffold_623_F	MHLGHLLFVTIVILLASSDGVMSTQNAGKKPIEDTVVGPSNSNGIKKTRALRTDNGSEERGFLSNLVKEAKVR
5	WWLETGKSESQVKKAQLLDELSGKAFETNTKLFQKFATKAYANKLNKWLQDDATTYSVWKILKLENVSSKQ
	LKSSPAYRTYVDYVNQFDELLQRKWGAYKLPEMVGSSKKEMMAKSSIWGEAKRYEAYVKMALDMEGVTG
	KALKEHKNYAYYNNFLEALKGN*
<pre>>scaffold_25_R1</pre>	MCVAPPTICLIFQLCQGSLAEILNDQVCRQNHPARQQLAISIGYMLDAARAVAYLHSFSPAFVHRDINPSNFL
072	VDAECNVQLSGFGESRSTVKRNGRALRPKMKVPELEAKYFTLNATVASPLGTTRTQLDSPCTHMEKNSAEYI
	APELIDGLSEPFFYGEAADVYALAITLWDILHPNGDKYPQANDDYVQVHVFECVLQGLRPRINSNLPLKLCSII
	ERAWQQDPGLRPSAQQIVKSLEDFQDDLCARLVLDLISDFHDCSSGSSLRTTEDLHVRTYTGAFIVDRMIDRR
	FVRCPAEAIRLGNALMDSSVLHHANHSTSFENNSASRYYFDVDEAQLALPIDSHIERSVRSSSSLSMNESSIA
	MLNVGHNDAHPRSSFSVGRTRSEPPCRCRQLGQRLINRKSTHFHRQVQFLTALEPTVDGNTLTTALLGGDE
	ASPQNSYNGLDMTPGIAATMA*
>scaffold_25_R3	MYRVLLLTVFALLCGFSNAAKSGGRLLRVVDSAEERGGFPSYLKDSFTKWRINSKINSWVEKQKTDEYVLKKL
36	GLSTLTGKELVKAAKYRQFQDFKVGVWLKEATPTTS
>scaffold_68_F6	MRLNLVVLAAVITLVSRCTAVSAASSTNQINLSNLNQALNTMQNHAGDGNRFLRSAKIVNDDIDSDDDIDR
89	DEEERGGKTWAEKFAKWHARGESADDVYARFALEPVVRQAYKYGQIGRL
>scaffold_24_F4	MRLSFLLLTATFVLLSSGIVASPTTKDESIPSPNQVLSEGRRLRVHKSSIDDVEERGFNPEKFNRLMNERGYRST
44	RFSNWVNKNYTDRYVYNLLRVDSNPNYKRIFNYYQTYLENFAPRLISS*
>scaffold_11_R2	MAVWIVIAAATALVRVSGGGRILVNCSWIASICVLPRPLRLADVDMAKMAARRRVFNGIAALYDCLTSS*
784	
-------------------------------	--
>scaffold_50_F8	MRFFYFLLLTSAALLSNSNATATVSGEGHVMTSADAPARALETNNGKRSLRYYATDEDVETDKYDQKNGKY
48	DNDDEEERNLQRHNSPNGLKRRRAGSRRIGRREYESDGGR*
>scaffold_46_R1	MRLSQVLVIAVASFVFASDTVAVATSNQAKISKMEQSPSQRLLRSNHYPVKEEEDESEDSVDFEERGFTTPDE
328	EDLEERSPLSSATVKKLENIAKGWGTTYSSVAMGTSSVSQTKAKALLALRDAYISGIKSEKNAAKMAILMANK
	S*
>scaffold_24_F1	MRLLFAVVLILAACLAAADVPALNDAPSKRLLRSTVRVDEEEERGAWDTLSNKFAKVILKPNQFAVRNMDDP
432	KVAKAAKTTLLKTFKSVDPQKFNSVENFFRGKAFNNLENYVLRLNKQDINKQTSVAKVLSTGLGDEKAFHLFL
	TATQSSDRAVKKSGKFFRDQLLTQWAVEGKTSAEVTKLFPKGLGANYYLHLENKYSGILMDLARDSQKRATR
	LAKLERARAAAT*
>scaffold_25_R1	MCVAPPTICLIFQLGQGSLAEILNDQVCRQNHPARQQLAISIGYMLDAARAVAYLHSFSPSFVHRDINPSNFL
150	VDAECNVQLSGFGESRSMVKRNGRALRPKMKVPVLEAKQFALLGTVASPLGTTMTQMDSPCMHMEKNS
	AEYIAPELIDRQSEHFFYGEAADVYALAITLWDILHPNGDKYPQANDDYVQVHVYECVLQGLRPRINSNLPLK
	LCSIIERAWQQDPGLRPSAQQIVKSLEDFQDDLCARLVLDLISDFHDCSSRSSLRTTEDPHVRTYTGAFIVDRM
	IDRRFVRCPAEAIRLGNALMDSSVLHHANHSTSFENNSASRYYFDVDEAQLALPIDSHIERSVRSSSSLSMNES
	SIAMLNVGSNDAHPPSFSVGRMRSEPPCRCRQLGQRLINRKSTHFHRQVQFLTALEPTVDGNTLTTALLGGE
	EISPQSAYNDLDVTPGIAATMA*
>scaffold_6_R18	MRVLSLVTLLAFVSAQLIAANADQTGIAAVDSNTALLPRVLGVESKRTLRRYDPSEFDSEEAVDSDEEADPVE
84	VADSDEEVVSEGEERVGIPGMEKVASKATKADDMVPNAAKGAMKWKILVQKNMGQLVETAKLVKKLKVG
	SFYSNVELEKMSLSALRQLDDIQQLQKADIKSNVFGTKATANGMRRKMTRTENMKLPPEQFLVSHVGRGA
	QRLGENGQRLLSAAVISKGDDINSKVLLISSSDTKKGDFLLPKGGWDHGETIEKAVLREVIEEGGVNGQLLHKL
	GEYPFKKGATAYAYMMKASTVYDDWAESIRYRIWVRTEMLWKY*
<pre>>scaffold_10_F2</pre>	MRLSVILLVVAAFVAALDPATAANDANTVVAVPNVHESIATGRFLRAHLEDDYPVKDERDEDDDNEKDEER
067	MFSFFQEKATALSAFKNLVSQSGDDLVEAVSNLTKKEFVALFNQGHTHMAKMVPGFHPGMSLDEFGAVVR
	AAGLSDDMKNALMVGYGKYLAHLMD*
<pre>>scaffold_58_R5</pre>	MVFKCWRITLFVVLTVTAPEILKGEEYGTAADWWAFGAVLYELLTGLPPWYSENAREMCKRVLRTPLSVPEY
67	VSTEAKNLLQKLLTRNPYERLGSLLGGPEIKEHPFFQHIDWEMLSFRDAPAPIQPCATSDTVVRPVIIYPYVDF
	VSNFLPCL*
>scaffold_13_F2	MVSVSSVYLFSAVLAFATLQGTTATSLSLDPYTTCTSYENFPGRDTDIKDGGSCTVIVPEDPSTSTSKRKLEWV
050	DGNDIADLEAHFGVPMERKLRNLPTSTTHSSPPWSGGNWLVNVDSINYVWDQGQPSAAEKYATAFGLDV
	KTFMDDVSAQNGIDSVQNATECTEDKECYEDLICAKRAGKSSGRCLPTWWGMGNAWAAAATLEKEPKCP
	VTFNGVTFQPMDIKALVTDIYDSANVSYVFTGSRYNGYEDSIDDYGRHTDASYRDLNPGFLHIAATNMLGLL
	NTTFIVDKDPDYVVWNQPVVGFEVREQMNMTPAEAAKKLYGLDTYPWNVNASSIVYVNSYLSWVTESLN
	GPIVTLNTTETGNYTYLLELNDDEEIIGGEWLYDSNDYHPDFLWLLEAKPAPDTATSFGLSYTNVTMLLEKATE
	C*
>scattold_12_F1	MQTIRAIVTALTLATIVAGDEGATRTLRGAMADNSELDQLEGSGSNNSILWVNASKPEERRPIRPNYHHGH
/55	
>scattold_81_R7	
5	
Scaffold 81 E1	
05	
>scaffold 47 R6	
72	WDFLNKFKVDTVISKLKGTKYGAFLLTYINSRPKAVRAAAK*
>scattold 17 F1	

914	CVPSLSRMLRILSGKSYREK*
>scaffold_31_F1 334	MRLLLWALLVTLVAFVSSINAEFTADSKVTQDSKEEINALTRLLAVDSSDAAKRFLRGDAKDLTTANDDSKELS AEGEERGLIPSSITNLVSKVKTGWANWKAKALEKAFQHMVKNGENPTTLAKRLDIGRTVEGRHHRLYEKFTA WWINYHTVAGT*
<pre>>scaffold_143_R 10</pre>	MRASIVILVAVIAVFTRSGLASASNTAQLQQTVPGATNKVENAEGGRFLREGLKDESDDLTSEERGIWGDLFK KAKPYLNGDIMYARLFKKSNSQLWRREITPDELKSMVIHLEKAGWSGEKLQKLKDKSHGYEKYFYTPIKD*
>scaffold_40_R1 065	MAFSSFLKFLAMATLLMVQINAEPSQQTRNLRIQSDDLPQEQARRLGPNWKALIINVKNNPGVSDGVVNLL KKADFVTGTSKKIKEKVKDTAGKVKDKVTGK*
>scaffold_51_F6 03	MQLTNILWMATVILFSCNDAAASRPELPTISGVDAVQLNPVNQRFLRSDYNSEERAVTLPFAKLDDVAVAKL TTKLFNTEKLKAALGKTKEAYNAREALFKEVNAVNPAVRKAFLAKLAKSGDYNWLLPYWLGFKLKGIRKVETT EIKNAVRAGNAVS*
>scaffold_2_F40 6	MVVVGVGVCGVASSATYAVPTDPDCAIFSSPQGVWNKYTGILDIPYGRVWNGRCWAVRPIRKRPAKLRKR RSRNQQHRSPRRMLRLNWLEPLIVKTKARAMTQMKFNLLTERRRSRKRDVPRQRPGQYAPGTARSVEEKP AALRPVGQVICYACSKPGHFAKECPDAEANARNDAYLASRIQRSAGKYNGERSL*
>scaffold_107_F 190	MRVLSLVTLFTFVSAQLVAVNADRTGIAAGDLNTALLQGVLAVESKRSLRRYDPSEFDERESEEEADSDEEVD SEDEERQIGLPGIEKLDDAVSKFTKTDDVVSQAAKADDVVEKAAKASPKWKALMLKTWMSLLRPEDW*
>scaffold_1_R39 4	MKTATAFATVLALIVATNGAHVSPNTPALRGLRPTADTPSVEDDKEDRKDHHHVKKVKKIAIPVPVPVEVPQ YIPVPVSVPSTVVASSNNAVVGPSTNVAGPGAAAPGAPGAPGAVTPAPTTLNGRPAATPAATSTTRSRPTP APTNFAGARPSGAQLPAAPAQAGAPGMAGFPNSVGIGDNTNNNAFGAGIGAAGGFPMTGNGLAGGFG GNPTNRFGGANGMGSGGMFGGAGPAMGGRLNGQSMGFGMQGGNGLGGFGGQDMMANFGQTGN NGFAGGOALTOGGFGGGNSFGGGFSRORR*
>scaffold_17_R1 815	MKACKFLVLLPSQALANHAVLHVVCDRLFRYTAIQRELRKKLVGDYVSMRTEMNQCQQKWLSALRHEHTT TLDRLEKIETRLDSLVEAATISMNKLQ*
>scaffold_11_F1 315	MRIGYVTLLATTAILASFGNVSGDTDLDHTQVAKVASLDAVLPINTNRFLRRRENYDDEEAATEERSAFEALA AKLDKSVLPAFLKIANVKDSERAFHILQQYPISLEKRLDILALHGLSKSDRKKVLLLIQ*
>scaffold_14_F2 398	MQPTLVIFVNVLLLLVGLIGRSDAVHFDETAQASPSDSTIESKTATNERFLRGNDKTIRTTEEEDRDITVGSLVI ASAKAVNMRNKINIMLARGLSPTRVLKKLNVVRMSDKNFNNFARFYARYLDKYSRKKPNLPKTAEDVIVLPK MKDWLAQKLLPFQVEQNLKDLASRNVNRYMQLYFKDADNIIILPRLERWENQKLLPSHFKTNLIEIGVKDTT KYMEWYMRNGGEQVVKAKLQKWLRQGIDPREIGAKLRKIGVTDTAKYVAWYIDAPVVTKLKTWVNTGLR PPQILDELIRAGVTDIQRYYNTISVIYGQRQVRLYRKNLP*
>scaffold_53_F9 3	MRFTYFVAVALTGLLASNDLVVASTADKTVVHDQVLSDRELIDTGLNDNEKRSLRTNEDLEDDSEDLKKSDEE RFSLIQLSNQPRYYVWFEDEMTPKDVRRKFGLTRHSIKLVKRSIYRGYVKYYDEHCSYFENRKKDFCRAQEY*
>scaffold_18_R4 67	MRLNLVVLATVITLVSRCTAASAASSTNQTNLSKLDQALGYQVGTMQNHAGDGNRFLRRAKTVDGSDDDI DSDDEERGGKTWDEKFAKWHARGETADDIYGRFFLEPIVRRAYKTANIGSLKQHEYYQKWLDYKAFLKAKE K*
>scaffold_25_F8 10	MRITYILAVTVAATLHSSVTAIPSVKSSKVATENGAVPAVIDSTHAGAGRMLRWVKEYEEDLDDSDDLDDDL DDDLDDDLDDDLDDDLEEERGFGDTLKKANPLKLVKKGTKLTAEQAAKVKEALKDAADYQKMIENANKLIRS D*
>scaffold_6_F39 33	MRFCFVVLATVFALLSSETPAIATAEKHSEALSFINGNNYGGTANRSLRSHDKVNAVEEERAIDADKLVKLLNP DDIAAALKNYDDEVVLFTRWWQHEDDVVNKLMEWPNRMKNMPILTKFNDFRTSMHYDQVLTPWLSTK DLDEVATALRASKMNLAKEQFNVWHKSGVKPAEISAAIEKVKNPLKRKGYGALNGLYKMYVSGEAKKTARA AAKKTEEFAAKRAQAKKIAEAAAKRAQEIKTQKETAKKIADAAAKRAAELKAKAAVAA*
>scaffold_193_R 7	MSKATPNLLGISWLLGSLVGSHYHLLNRPLCAWNGRLNWPPPRLGVGLRFHQQRRTLIYLEMHRRYLLLLQL TPCFLPPTLHWLRHLRRQRSTKT*
>scaffold_686_R 5	MRFHALALLAAFLAFGSGNDAFELTADGAAAYSRFLRKSEPKTNEERVFEGLTSKVDDLIDTTKLQKYKLQWS KLQLGDDLTAVLKSPDVAKIAKFNLNRASGSQVSMFQRITAKYSDDILARTLVSIERSADDNPALLAMVKQLR DDQIANWLKNRETVPGVVSKLKLGTDESIFRSKALDVLEDFIKKYNTARNGDESLLKTLTTIYGGESELVTMVS RARATYPYAQMNPQSVDKASNIENQ
>scaffold_5_F19 44	MRLSYALPATIIAVTFLSSGNAVATTDGGTMDLSAMTSPNAVASIDAAVGGGKRSLRYHNKDPEDESDDEAL LEEEERKYTNMFSTTKLDEMLDGTKMMSRFKKWKEKGYNTYNLPAETQGDKYTWIRQKYRDYLYHN*
>scaffold_2_F15 74	MGCLHLLLAATTFVACTNAADQLEMPVKRGLRANEERAFGAGVTEGLSNWVARAAPKLLLTDNELEHLA MKVTSTDKVFKMLKLDDGLDGILRNPNLKAFASYIRKVHATNPDQVLITTLITRYGDDTLAKFLFEAKQVRRTE

	ESAKMLQAAQFIKWFDDGKTPNQVFNLLGLKHLTAYEDKFHKLWWEYVVAYAHLASKLKKPLPVEL*
>scaffold_24_R7	MRFTYFLLVATAALLASCNAAAATSNNQNKLSTMTSTDAVVSALESSNDKRFLRSYRKEDDGDDSDDEEERS
96	VMTAEQVAKWTKKVERWVKKGHTPSYIKDKLTALDGTMNAKNREKYRMFGAAWGRANPHELGRM*
<pre>>scaffold_10_R1</pre>	MRLSVILLVVAASVAAVGPTAAANEANTVVAAPKLPTARFLRAHLEDDYPVKDERDGEDSDDENDEERMFS
115	IFGEKATALKAFKKLVSESDDDLVMAVSALSKEEFVALFNQGQAAMAKMVPGFRPGMSLDDFATAVSSAGL
	KPDMQNALMVGYGKYLAHQMG*
<pre>>scaffold_3_F29</pre>	MATLVVCASNFALASTVADDLHQEAEGNVDRRLRAAAPANKDNVAKIAGGFLTKIKEGSTLTKAEQMIKNA
10	NGDEAAVKKAILLASTAKESAKMSDESIAKLSAMITTAVVKDPKSWPRLQKFVKVTLGAGVGGLALYGAYKLL
	FNKNSSSGPVTTTTGSADLVAASTSGSGSA*
<pre>>scaffold_26_R1</pre>	MLLCVVILAFLNTVILVLCPRIGRVFVSFSTVAAVWFDAGDTVSLSPVLAPSSWFVVIRLAAAFSGEWILGEDS
068	RDLRPDDMDSRSLVTILTEELAEVGGVEAEGRFCSWLVGFLLPSAIVDCEDRMIFELLPAWGRFDTSVSLATC
	R*
<pre>>scaffold_8_R11</pre>	MWLLLCVGFAFSVCVDAVRLGSRVLLCPPSSPPYRTPYCPQQFSHPRPRYEDTPQRGLRAFGAPALDGFVLR
92	CRIVPRSSRLTLQPVLLTV*
>scattold_93_F9	
1	
>scattold_24_F5 45	MQSAAYIAAFAAMLAAVKAESGSILQGRQVLRGQLRVHQCADHALGHRGRQAVPAVHEGGRRVLRHLPG SVPVLLHVEDGLPVH*
>scaffold 2 R11	MHFHSFIFIFIHHTSMTTSAAASCSEAARASRTIELLAPVYEDVCQAIKGGSTTVPVTLERTADDWIQLVFPQT
64	NVRKDVACVNGASGCRCDFVRLLRALLTLTEQLVVKVAATYVLAHRSADVATLEMRVQVHEEARRGTSLRD
	FDLVLCHLQATVLSVVSRSGKKKIEARHEALESARSSSCHVVGCRLHPWDASSGQFNPSIN*
<pre>>scaffold_214_R</pre>	MRLAIKTLVALAAVLLATSTEAAKAVQTGGDVNVVQSSHILPGENKRLLRSEHDEGKLLEDDEEEDDEEERKY
4	GANLFSTAKMEKMLGNDWYRYQVARRWKRDGYTWETLPKDVPVDLVRYFKGFRERHG*
<pre>>scaffold_7_R12</pre>	MVKVVRVAVVGAVFMASVDATSMKCGTTQATPLGAPMNDCNPAYGGEIDPNCMPEVAITDAEGCEHAT
23	GPAMDEIIGSITGASAATRSLRRMEDASNSDVADLETYFGESLELSFTTLKEQYSSASVPTTPWPGSYWPTYQ
	DGINVIWKTGDVSASEKYAIAFGLDPTDFMNKISAKTGIDSRSSSTKCTADTDCTYRNDGSVCAKRKGVESGY
	CVPTWYGICHAWAPAALLEAEPQCDVVKNNQTFHVLDIKALMTDVYDGSSISTVFTGARFNGPDIPEEMDA
	YGRYISAPRRDLGPGFFHIAITNILGKHKKPFILDVTAGSQVWNQPVRSYQVQTMELVDANEASQKYFGVSP
	YPFNSEMVFLAYVKTTVSWIVEAYADGPLVSSGQVDAYTVSNDYQYVLELDANYAVIGGEWVEGSKTDHPD
	FLWFPTERPDASTITISTGLSYAHVKELLELSLACGSSGGTSENASASTSTGGSTSASISGSASSSASKSTNESTSS
	SDA AVTDEDSEGESTGTTALAHDSC*
Sscaffold 568 B	MRI VIKTI VALAAVVI ATSTEAAKAVOTGGDVNAVOSSHII TGENKRI I RSERDEGNI I EDDEENI I EDDEEE
2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	RKGGANI ESSAKMEKMI GNDWYRYOVARRWKOYGYTWETI PKDVPVDI VRYEKGERORHG*
<pre>>scaffold 81 R2</pre>	MTVRDIVELAAVVI I VOAOAI SVPTDSSTRIJ RTSVDTEDI EDEORAGI AESVIKKATAAI RPNVTPEKI ANW
17	
17	VDGTEAIATKLEAAOIOKWLSSGKSADDIFTILKLDOVDDSLLASPVFDAWNNYRKVFAODNPTNLHKPSVR
	VLEFVPEPLQQGPTS*
>scaffold 81 F2	MHLQIALFLVVVSLLVNVEAVPAKAQSDSVVSFRHIGRLLRDEREERGVSANAVETITDAVESKINTAQLKSW
24	LESGESADDVFKLLKLDSAADKVLGHAKLDEWIEYMKLFNGQKGSKKTTLIKTLTAHFEDDGVARMIQKALQ
	VDSTAKMAKRLQFEQIQRWLGHEKTPEEVLTLLKLDINRYDLFEKPELLTWVKYLDDWNKMYPDRQTTLFA
	RISPLLEEGILANLLIKAKSVASTEKIALRIQAEQTASWLKAEKTPDDLFTLLRLNRAEDSPLLENPIFDAWVKYA
	DDFREMYPKVSFDPIATISEHYTAAQVATMIVEASKSPSTSSIAHRLNTEQFRDWLNTRQSPVRVFKLLKLDE
	AGDKLFQSPVITTWLNYATFYNTKREKVSITTLLRKRFGDEVLAGILTEAQQVPATKEEATKLLTSLVGRWPKS
	RVHPDNVYKWLRVQGREKTDGFRLFYERYAAAYKAARNG*
<pre>>scaffold_37_R1</pre>	${\sf MRVASLVLLAAVTLLASREVVCATTNTLAKADTDTINAAIPIDDARPLRNLRKKTTSVGYDAEERAAEEELVDV}$
044	ELLDKVISDHAYAKQVFLSWLQNGQTSQDIENRLETLGVQDKYGKVVTQYAHYLTLLEERSA*
<pre>>scaffold_18_R1</pre>	MRFSFIVLLAAISSLVTTSHALPALPHEQSTVSKVSSTNPDQSDTGAQPSERRFLRSAGVGSKMLKSESFKDAK
184	FAEWLKQNLSSFTVFESKLGLKEKYRGLYDEYAKLHKASGQYP*
<pre>>scaffold_175_F</pre>	MWRLSCWAMAETLSVPSLVCSFAVPAVLELPSEVPAAVIVRGSRQVCLRGHRRHRHRRHLRWNCCCSSLR

27	RLRHPFRLRRASLVVLVESSLLPSS*
>scaffold_13_F2	MTALSAWCTAVNSVDAGKCCLLVLMLYCAAQRLVNPSQLLPTGVMGVCRHLRWNHWISRRDTVLQASDT
21	LQDAAYATGVSACEVTKLCELLSSGRSRGDAGSRLCPV*
<pre>>scaffold_81_R2</pre>	MRVHSLVLLAIAIISATDAVAAQSLPEEQNVSRFLRTNKAKETDSEERVNLAGFELKIGFVDDIIEKMHLSPTFK
28	ELLANKGDAKKAFKAFHVDDVADDVFQSAQWKEWAEYVKYVAVKKNENADEALAAAMSVAYNPDGLSKL
	LAKAAQNPNTKDIATVLENARQSSWINGGFTPGAIFKTLNLDKADDIFDTAPFATWTNFLKAYNEKHPKKA
	MTEFEVFSSVYGQQKFAKLLASADDGKYAQAMKKELVQSWIDDGVHPTNMFAILKLDKVDDLLTNPVLNL
	WVRYMKEFNELYPKQATTMIKTFTQSYGDEKVALMIQEAAKSSDEKVVEFAKNLQTAQINQWMVDKKTPK
	EMLTVLGINSQTLTENPLGNVWRAYNKEYTKKMANGDFAFQP*
>scaffold_50_F1	MRLTCILLMTAAALVGVLDASAATTGNSVVANAAMVISPLAPESQGRRSLRLVYDDEDDSADEKDEEEESA
228	DEVDEERGWFSDKVALTSLASKFVGKSTDEMGEVIKGLTPAQINTMFESGEDSIQQILPGFKTGMDFTKFDE
	LVQKLPQEQQALVMSAYTKYLHNNGKIL*
<pre>>scaffold_38_R2</pre>	MRHNFAKVLVAIAFVVTCSAEAEYHTTKSDPLALTLSLRGGVPRTRSLRSDKYAGNEERAAGTSAVESLVKAT
61	TNVDDVETWLKRGDTTNKVFKALKLQKAGDNLLDNPQLPTLMRYLRLFNEANPTKKTSLLATLTTHYGNHG
	LTKIIEAGLASTSKKTVATAKHLQTEQIQYWMARGRSPESVFGVLKLDKLLPFTWISSDLFEKPGLKTWIRYLD
	EFNAENPKQKTTLISILSKRYNDKTLANMLIAANKAESTSSIAKRIQSEQTLQWLSKGKKPEDIFTLLQLEKAGD
	SLFQNPLFPAWIKYADDFRLRYSETDLATMSTLINHYSDEAVAKMIMAASEDPNTKMLAKRLQSELQKDWV
	FSGQTPNDVFIMMNLDKTLDKLLENPLFDIWYQYGLFVNKMSAKANWNPITALLRTYGSEKELSGILMAALK
	QPSTKAMASKLLQIT*
<pre>>scaffold_13_R7</pre>	MRLSQVLVIAAASLLFASDTIAVATSNQAKISKMAQSSPNQRLLRSNKYSVKEEDESEDSVDFGKRGFATNDE
93	EDLEERSPLSNAIVAKLDDIAAGWGTTYYKIAMGQSSVSEAKINALLAMRDAFISNNKKEKELARGMLMLAN
	KS*
<pre>>scaffold_29_F7</pre>	MRLALLFLLVIATLVATGNTLAVEHTEGKITSSDTLLQPTAISNPDTTQTRRLRTPESPNDGEERRLVNVDLAIK
71	DAVHEVRKLTKWKLQFAVWKGLKKSPTKLIQEWGMKYPYTSDPRWAKILAYKAFWGKGPLKYP*
<pre>>scaffold_25_F1</pre>	MQRRMSTPFFLMITTIGDAQGEVEWVMIPADNSPLIKSDNFSRNCGATRRTRWATGVALPTSMSQTAPRA
705	AGGRAFPFPIRRCLRFQMICLNRSICSGVLPSASSIWISSILTSSVVALTSGRMAI*
<pre>>scaffold_102_R</pre>	MQQHLVFCKVANFILSTVLFFRVALPATHADLNFPQSHEQQLRPLTSSVLDQTRAAVARGRHSQLRRRRLRG
91	WR*
>scaffold_83_R2	MSFTIGVCLVVLLQSEVHIICRKTHKKFSKNSFVNFLVIAKPEGYGQPYGLHHPTQVLRLGRVRLLRRGRERHF
91	YILDSN*
>scattold_3_F32	MRLTCILLVAVATLVGVLDASAATTGNTVVANAAMVNSPLVPESQGRRSLRLVYDDEDDSADEKDDEEEES
92	ADEVDEERGWFTDKMALSSLASKFVGKSADEMGEVIKGLSPQQINTIFDSGESSIQKLLPGFKTGMEFDKFD
<u> </u>	DLVRKLPQEQQGVVMSAYSKYLHNNGRFS*
>scattold_81_F7	MRLHCLLLLAAVALASVDASPLQLNQQSISKANGFLSKRMLRTAESSEDSEDEERMNMPGLDTVTNAISSTV
0	
(()) AC 50	
>scattold_46_F8	MVMILIFIIILETGEGFVPDGDNRHYADTNSAQTLKQTDIGEMREQGASGKEIIQKLVENSSTWETKTEFSKQ
62	
<u>(())) ()) ()) ())) ())) ()))) ()</u>	
>scattold_6_R31	
08	
83	
1	
	EQSKSVFATLSAHYDDLPLATIVIKTAKEDVNTKKIAEYVERGLLPKWAADGKAPAYVLNKLATDKQDKERVL
Scaffold 110 D	
>scaffold_118_R	EQSKSVFATLSAHYDDLPLATMIKTAKEDVNTKKIAEYVERGLLPKWAADGKAPAYVLNKLATDKQDKERVL TLFMKEIRIVEDKITRAES* MQTRYRLLLVLLAVFLASTRASGASLQETTATSENPRPDGPVPTRSLRESGAVRLLLSPSEEERMGVKEAIG

	ARLQDWLDKNLSPPQVFKELGFSGTFASAQGHPDYKYFEQYAKMWSDLQVRISSGHA*
<pre>>scaffold_31_R7 46</pre>	MAIFVACSTVATAESVALTLGDNEVRRLRLNQQNIAKAAGDLISKSKESATLTKVINIAKTANGDEAAARRAV
Scaffold 8 E20	
12	ELTA*
<pre>>scaffold_18_R6</pre>	MSSSTGSQVPSSFLLGSLAGVPFALIAGTHARPLRAVVTVAASNTVPVVSVGMNSPRSPSFAATAISMGVTA
89	FAGICALPAIRAGAVVRPASASLVRNGITIVRPFSPYFTSLRSKSIFAFLSVRKSTPSRNSSVQPEMTHASTQIVS
	PNTFWNWKRQTPRTSSRNPPIPSTYSTGASRGSLFPYCIHAVPATEYLAQLSSLMNFCSPSIITHAVGNPSSAS
	TYCTASPSPGSPCRCTSLTRANSVSVPGLSPHFRRSCRCRNCSSASVACFSLTASRVSSKVRFSTSLTRSSLERA
	RRTRLRFLILFLRRLRRRACCNCFSGPSSLPVSGLSLSATTSNSASSPILSGSAASPSSEIACTTSSFAGVVRLLPPS
	APTPPVASCCAPSPDRASSRTRCSSLAGRCR*
>scaffold_3_F37	MPRATSPSLVLLSLYNSIGLGGELEPNRSWISVSVGIHAAQKLWWISARRDERVILARKLRPGSVRGLFAGAT
06	RLLGWVVRVFGRRHGGGGFFFLELPIFSIRIVERILFRVGSEVGLLRGMLLLVESMVLGDREVTLCRETGSSSA
	NGHCRGCACCRGSRRGLRVRRWERGLERWRVHSEMDSSVLPET*
>scaffold_2_F13	MRFFWVVLLTVIIVLAGVDTCTARGQSKLSSSDFVRPFVKNAPNTQLESKATRLLRTDVTKYDNDEERAITLPS
82	GLSNMISNVKKWFSNFISKIKLSFGEKRKLNAWLKQGKTPDEVFEILNLHKGTKGLLANKNLRTWSVFMTMY
	NRKNPTKMVNMLGTLTKYYGNEAVAVLLEVSRQGKTRPLANRLQTQQLHGWARNGLNTDVVFNLLRVGE
	GSVKTLAQNRALNVWVYYFRQMNFYNKGWEAELTKKLLTVYDDIPMAKAFAVAKTDPGTKYISSQLQRFQ
	FKKWLADKVEPATLLEKLKMDKTKLSFEPTVEVYVAYSSFYKAYSKTVR*
>scaffold_60_F7	MSWMICWICLVTKTISLMQCIRDQCRLHELERTTRNLRQLSFKLVALKSSTDFFRHYYLRCSHDPLLFVGAPK
08	γ*
>scaffold_24_F1	MRLSFIFLLAVTFSSLVATSHALPHEQSTVLKVAATNPDQSVAEGQTSQKRFLRSAGLGSQMLKSESFKKSKF
319	DELLDDNVSSFTFFVDMLKSKKKYRDLYDEYAAIRKARGAYP*
<pre>>scaffold_18_F2</pre>	MRPNLVVLAAVIALVSRCTAVSAASSTNQINLSESNQALDTLQNHAGDGNRFLRSANIVDDSVDDIDSDDDI
148	DNDDEERGGKTWAQKFAKWHARGETADDVYKRFKLEPFVRYAYKHRQIGMLKENEYYRKWSITWPS*
<pre>>scaffold_5_R12</pre>	MQFTYVLLLALCIAITASAAVQDNEYQAVVSPFAKRSLRSVDTGTESDDEEERGIIPSSLKDIVKKGTSKVSDW
81	SLLRKYKSLQKAGRTDDDIYKLWVRQRKNTDQIYTRWIRLGKSEEEVSQLFLKHGLNAEVLYNILSRQGKSMD
	DIYSLWKKLKLPEGQIYNIWLKSKTDDQIFSAWYKAKMDPEDIGHVLRLGKDPNTKRNVPFWEKYSEFYRAK
	KYRTS*
<pre>>scaffold_81_R2</pre>	MHVRLLFLPVLLLLVGSDTSADKSVPVTTRLLRSQKLLQITTKSECSESTSDPSTTSSKTGSLNRGYPRRSDPTK
13	HDTNESNILFYLKM*
>scaffold_7_R22	MRLHRTLLAPIAVFLAWNCFDPTTNAQVVAQDLSDKAFAANTNALRSLRSQKSSTSEDDPIINEETRNTMYT
62	QLFPAWYAAGKTPEDAFEELKNPSTGDENWPIYKNYKMYYDTYKTMG*
>scaffold_39_R4	MRTPSFLVLAAALLVNADAASSRHLANPNVQNLAHVNDNWLDTKRFLRVHVEGEERGISAPSVKMLQGW
58	VERGLISDEAVKLISLGDKADDLLSASLLNAWFSYVKVFNKENPSDKMNMIKTLTARFGDEALSTMIETAKRS
	SKTSAMAEKLQAKQLANWAALKKNPDDIFALLHLNSAKSLLFDQPSVNSWLKYMDDFTLKSSSVEFSAISTLR
	KYYNDETLAEMIIVALKSSKTSEAAKRVETELLRTWLNSMKSSTDVRRLLNLSTAPQSELSLIWRNYVALLNKV
	DPKFKTEMLQAWVKKGLITDDTFRLLTLGNAADDLLNGSLLSAWATYIKVFNQQNPTEQLNLIAVLTARFGD
	EAVSTMLEAAKKVPTYHTIANRVQTEQMKLWLDAGKLPDHIFVALKLNTVKTKLFDQPQLNTWVVYLDEFN
	KANPNSKTTLFSTLQTRYSEATLAKMLVVAKQNPNLESLAVRIQGEQLQFWIKANRRPGDIFKMLRLNILGSD
	LMRNQLFTAWVQYTDEFRKLNPGTKLTTLATLRKYYSDETLVTLFLKASQSPNTAKMGKRMESEMLREWFS
	AGTKPTHPVGVFALLNLGRTGTKVFESPLYNVWTNYVGFMKKAGAFRGDTITLLRGIYGDESLAKVLIAAGK
	VQSTKNVANTLEKELFALWKAARVHPTQIHKLLRVENVSRNSPIYKFYGDYVLAYTRA*
<pre>>scaffold_129_R</pre>	MRLSFVVLLAAAVSSLVATGHALPHEQPTVSKVTATNPDQSVTEGQTSEKRFLRSAGIGRQMLKSDSFKKTK
17	FDEWLADNLSSFTVFIDKLGGKEKYRSLYNEYAALHKASGQYP*
>scaffold_73_R2	MRLLYLAGVAILSFAADATGAKVLVSDDSNHNGAQAASASVVSTTRLLRARSVIDEERAGGISASASDKLAKL
12	FKSSKVTAVAQQGENCGKCLLSHESREHSLYEGI*
>scaffold_46_F1	MRLSQALVIAAATLLLASDTVAVATSNQAKISKMVQSTPSQRLLRGNKYPIYEEEDESEDSVDLEERGFATPD
215	EEELEERSPLSISQVKKLEGYARRWHTTWEKVALGRSSISEDKVQALLALRNAYISGVKSEKNAAKMLILRANN
	S *
<pre>>scaffold_15_R1</pre>	MRLSQVLVITVASLLFASDTVAVATSNQANLAISKMAPSQRLLRSNKYPIKEEDEEEDLSEDSIDVQERDFTAH

	ANKK*
>scaffold_77_F1 29	MRLSYIFAVVLVAILQSSGTAHPVYEDSKQIVSAGAGDATKVDSGRLLRGVEEKMEAEDEERLFKNLGTYLKKI PQKIKDSWEVKKAKEQLERSRNRRKWIREQNAVPTS*
>scaffold_244_R 1	MRLTFVLLAVAASVLTRTDATPATYGEVTSTIESPNRSALKNMAETLKERRPLRRPRCPSTIYRKKSAQTFSETR CSGPACPQSLSRGPSR*
>scaffold_5_R34 48	MMRWRLLFILHALMATLTLPSSCWTVEQILTRGPRTRKLRCSTLPRRDNTEVFDCLYEGNATLQRRIATVTW RRKKLVMLKL*
>scaffold_107_F	MRVLSLVTLFTFVSAQLAAVNADRTGIAAADLNTALLQRVLGVESKRTLRRYDPSERDSEGDADSDDEDEERI
161	ITIPGIEKLDDAVSKVTKTDDMVSKAAKASTKWKALIQKNINELAETGKLVKTLKANVLYKDVELEKMSLQGLR
	QLDDIEQLRKVDIENKVKGTKKTPDGMRRKMEHTANMKLPPAQFLESHIGRDAQLVGESGQRLLSAAVVSK
	GDDVHGKVLLISSSNPKKGDFLLPKGGWDDGEAIEKAVLREVIEEGGVKSQLI
<pre>>scaffold_10_F2</pre>	MRLSVILLVVAAFVAALDPATAANDANTVVAATNVHESIAIARFLRARDEDDYPVKGERDEDDSDNDKEKDE
271	ERMFSFFQEKAAALTEKATALSAFKKLVSQSGDDLVTAVSGLSKSEFVALFNQGQAAMAKMVPGFRPGMSL
	DEFETVVRSAGLSDDLENALMVGYGKYLAHLMPK*
>scaffold_73_F3 95	MRFGFFLLAAIALFGLCESTVLNTQQKRNLRLSTTNDNEERAIGNISKVDDVADDASKNFAFMINMFKEWDE LSQAQIIARMFKQTTPDEFAAMWIMYKVYQKLGADDLIKTLKAKAK*
<pre>>scaffold_6_F17</pre>	MRLFIKTLVALAVAFLATSTEAAKAVQTGGNVDVVQSSHIVPGENKRLLRSERDEGNLLEDDEEE
3	RKGGLNLFSTAKMEKMLGNDWYKYQVARRWKRDGHTWEHLPQHVPADLVRYFKGFRERHG*
<pre>>scaffold_52_R9</pre>	MRFAFLLFVAAVSLIASGDALSTQADATSRHLRSHHQTNTYDAEEEERGLDKSIVKSLPEQFKNMYKYPSNM
85	ENVLESWRTGLQSVDDAVMYMKSLGMDFDAISHFVDAYRKHINKKGLPY*
>scaffold_78_F7	MHFISSFIAVALPFNVLAEELAWELDCSVPLGFGPGLRLGVDFDRAVTLEFAIGFSFELTDFLEWPRCWRRVLL
7	GHLSSSNLRCWRVSSLRCVLPRGRRQLRRSFTSAEIACIGEPGYTS*
>scattold_26_F3	
00	
>scaffold_431_R	MKIQLLHLHVLFLVQSLDVLHVLRGYYHAILREVHPVPLGLLPDANNHSLQRPLREILVTRISINRTGRPENHYL
1	
9	
Scaffold 30 B7	
9	KTRI KWRMSSDINI F*
>scaffold 11 F1	MKIYTVGAMI EVAATVSYAI ESROI RSOOOOTGARIVAGDGIEETAGDDWNVI I SDRKRGAKOAGTI VRD
229	SGKDEERLVADSRRTOTYPPVO*
>scaffold 10 R2	MALGTVVVAHTTLFTHASSACLHFSGCPAVKRTAVEDFDSRYSGASTSPPRVRALRRKRAKRKEISYGVWRL
236	HQMTAHTRGMRWSTGFCGPSPPLLVPFHLVFVLATYVIRCSSAPKPSKR*
>scaffold 29 F6	MRLNCFLFVVAAALVASTNAAASSIQQNHVSQTTPDIQAATQTDSVVKRSLRYVKDEEDSIDTKDEERVQGI
61	NVKELNKILSAKRIEIPANIEALSKKAQRDLVHTFKGQDLTKKMFATKLGMRDVDDVANRHYPFFEKWEHLF
	RPGKKEKKVPEMLIGNHEYYY*
>scaffold_46_R9	MRLAHLFVVLAATFLVSSGALAATNTDSVIKTDSYVKGSKPATHRALRTYKADFEDDDDDDDDDDDEEERGIKDI
61	PLERLQSLGRKVGVSADDVLSDAAGVMRTMSESQIKQWTKGLNKLKQLYKKAKSPRISDY*
<pre>>scaffold_46_R7</pre>	MRSSFVLLVAVVALLFRIDAASGSSMKLRSGQNLIPSTETRETEQIGRVLRGEAESDPESEERGITDLAKLLGTK
83	SKTMFDNNALATIVRLKSQFATADDAVQAFMTRNINPDAVYKWLKLYKANNQVVKKTGDAAPEYLLWIRY
	AKEYKDAHPKWVSKLPKGA*
<pre>>scaffold_29_F6</pre>	MRLAYVVVVAVVALFASQDAVSAEADVQLGNTLNVNSESTTRFLRVESTIEAEDEERALPSSLTKISEKLKPVG
55	DKVKPLTNKVAPLTEKLTDKTMAIAVRLKPIADTFTGKLKPIADKLMPILKPFAAKLKSAPVVQKIIEKFKTFAEK
	IKNYKVGGHTLSERYTMAKFENWFKQNKSPDDVKTLLKVGEGATVNTKNYDLSVQYNAFFQWATRDKEM
>scaffold_710_F	MRLTYITIVATAAILACCDGKSAVGDSKPLAVPQSHVQTGNNANRFLRVHDDEEERIGTQFITSLSDKFASAA
2	
>scattold_2_F16	
79	KSAAELEKAAQAPKNVNPEKLLAMMKANAEEHEMNLAKSGVIPKIMRKEYNIKRMRKVLSKDALMRNAN

	YQEYERFKEFWNKLPKKLKREFLAKAASLH*
>scaffold 225 R	MVFSVFFSAILLSVSASFNYNVNLETPRVQYRGWPTTWFRSPIAKILWERYLRNLADSIDIPLNPDIRGWHMS
6	NPPLEVCNRYLLIYPFVEQRIVHSLGQFCSAGDLLRFV*
>scaffold 237 R	MORRMSTPEELMITTIGDAOGEVEWVMIPADNSPLIKSDNESRNCGATRRTRWAIGVALPTSMSOTAPRP
14	AGGRAEPEPIRRCI REOMICI KRSICSGVI PSASSIWISSII TSSVVAI TSGSMAI*
>scaffold 614 F	MAESSELKELAMATI I MVOVNAEPSOOTRNI BIOSDDI POEOARRI GPNWKAI JINVKNNPGVSDGVVNI
1	
>scaffold 44 R8	MRESYVLLVAATALLATANAIFADPTSRSI RAHKSHEKSODEFRAFTYTENESI WDDI ENSI PEOFORMRKE
18	PWYI RRIERSWRSGMGTSDEAVAEMKSOGI SOKAIDOEEDAEI KYRAHKI AKGK*
Scaffold 29 F6	ΜΒΕΙ ΝΙΙ Ι ΙΙΔΑΛΙ ΤΔΑΛ/SASEDAKPIOIKGANNSPNE/SAHRI I RAHKESDI DAFERVIPD/VHI SKI EKVAIVP
63	KIVGWRI RKEDVNAAMADI GVMAREVGTKNMAIIRWYOKIVKNGGEKHTPKGKYITORK*
>scaffold 111 R	MDSTASTAVLIAMTAI VDAORI I RSEASTATDSTVSTADESTI SSGSGETPGNVEAOSNEGAI DKEGSTGGN
171	RPSGSMOGGAGRGEGGSGHNGMONVGSGEGLAHGSGSEGGHGGEGRHGGHGRDEGSESGGDEAGSE
1,1	GGHDEGGSGEGGNEDRSGNGEMPSIGSGEVPTSGSGEAPTAATTTSSTSSVTASSSA*
>scaffold 443 F	MRLSVILLVVAAEVAALSPTAAASEANTVVAVPNVHESIATGRELRARPEDDYPVNDERDEDDSDDDKEKDE
6	FRMESEEOEKKTALKAEKKI VSESDDDI VMAVSAI SKEEEVALENOGOAAMAKMVPGERPGMSI DEEETV
U U	VGRAGLRRDMFDALMVGYGKYPAYVM*
>scaffold 29 R1	MEPRI YAIAI IVVAI ISTOGGTRMDEKOPI ALI TTRSSRHI RSIDDRPRDTKTEERGPNIDIMISDAAKKVKKAT
159	
>scaffold 52 B3	MRICEALLI AVTALTAVVSGSSVNI RASOKIOSGDVAOYKTSGRELRGDI NTDEAAFERVNEGELKPDPI KSE
	AKOOMKYAYNERIENDI I KKEDDPDAI YTTI NI SKMKNI *
>scaffold 57 R8	MAESSELKII AMITI ITEOVNAETSORARNI RIODDDEPOEOARRI GGNWKAII INVKNNPGVSDGVVNI IK
93	KVDEVSGTSKKIKEKVKDAKDKVKDATEKVKDKI TGN*
>scaffold 592 F	MRI NI VVI AAVIAI VSRCTAVSAASSTNOINI SNI NOAI NTMONHAGDGNREI RSAKIVNDDIDSDDDID
19	
Scaffold 73 B3	MRI SYIVI VVVASVEAYCEGASSVSDSERI AVPOSRVOAEONSNREI RVHODDEERAGVEKII TOSASKI KRT
84	
0-1	p*
>scaffold 18 R4	MRLSI VVI AAVIAI VSRCTAVSAASSSNOINI SESNOVI DTMODHAGDVNREI RSANMVDDSVDDIDSGD
58	AIDDEFRGGKTI TEAWWNAI GKTADDIYKKYNI KPI VRDAYKYNOVGEI KNNKNYOKWAAYSAYLKDRGL
	N*
>scaffold 16 F1	MSLLLSWRSRLSSLCLLSLNVSVLCLHGFDIVDFTRCRLRLWFFGHCCFSGRHLRFLSCNRSFLFVHWLNVNF
314	NLRFLNCSLHFFHRLNFSFRVNLVRFSGLLDLRFLVNCIRLHPFRKWESGFQPVVKVFKHRLPFRIVEKRVEISV
	EHLKLLVTGRQSVVKDLRQFQTRDLVVRAHHHEHRQFA*
>scaffold 63 F2	MKLCHIVALTVALLVACVDPASASKDSKLATQGDNIPTKRLLRTTEDEERGISVSLGLEKISNAFTSGKTKEFKN
87	LVKAGQSADDSFANLGLGHVLNFEKGGEVRTKMVAKFFTSDKFKAWSKYTHKMNQKDADSAMLAGLTRA
	YGEKNAAVIILLGKDSLSSHTVAKKLETAQFTKWYTVDKIPNADELVKKVLDVPRSKLRKYPREMSIWDNYSK
	FVGKYVLNPRPGPPVRK*
>scaffold_25_F1	FVGKYVLNPRPGPPVRK* MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK
>scaffold_25_F1 242	FVGKYVLNPRPGPPVRK* MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV*
<pre>>scaffold_25_F1 242 >scaffold_12_F7</pre>	FVGKYVLNPRPGPPVRK* MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV* MRALKLLLVTVFLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK
>scaffold_25_F1 242 >scaffold_12_F7 08	FVGKYVLNPRPGPPVRK* MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV* MRALKLLLVTVFLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK STSGATNWLKDKKLAAKLKLRALPMNLDKSVQKTMKDGIDPDRVFTLLNLQKKSNRKINGFRTGEYNLWKE
>scaffold_25_F1 242 >scaffold_12_F7 08	FVGKYVLNPRPGPPVRK* MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV* MRALKLLLVTVFLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK STSGATNWLKDKKLAAKLKLRALPMNLDKSVQKTMKDGIDPDRVFTLLNLQKKSNRKINGFRTGEYNLWKE LTVEWTKTYPNWVSNISKKPK*
>scaffold_25_F1 242 >scaffold_12_F7 08 >scaffold_25_R6	FVGKYVLNPRPGPPVRK* MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV* MRALKLLLVTVFLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK STSGATNWLKDKKLAAKLKLRALPMNLDKSVQKTMKDGIDPDRVFTLLNLQKKSNRKINGFRTGEYNLWKE LTVEWTKTYPNWVSNISKKPK* MRLSFLLPVAMAAIYCATCNATADSDQNKVSMVQSLDARLNGQADGTRFLRAHHENEEESDREERGLTDL
<pre>>scaffold_25_F1 242 >scaffold_12_F7 08 >scaffold_25_R6 73</pre>	FVGKYVLNPRPGPPVRK* MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV* MRALKLLLVTVFLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK STSGATNWLKDKKLAAKLKLRALPMNLDKSVQKTMKDGIDPDRVFTLLNLQKKSNRKINGFRTGEYNLWKE LTVEWTKTYPNWVSNISKKPK* MRLSFLLPVAMAAIYCATCNATADSDQNKVSMVQSLDARLNGQADGTRFLRAHHENEEESDREERGLTDL FKTEKAAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYAIHLDY*
<pre>>scaffold_25_F1 242 >scaffold_12_F7 08 >scaffold_25_R6 73 >scaffold_74_F6</pre>	FVGKYVLNPRPGPPVRK* MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV* MRALKLLLVTVFLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK STSGATNWLKDKKLAAKLKLRALPMNLDKSVQKTMKDGIDPDRVFTLLNLQKKSNRKINGFRTGEYNLWKE LTVEWTKTYPNWVSNISKKPK* MRLSFLLPVAMAAIYCATCNATADSDQNKVSMVQSLDARLNGQADGTRFLRAHHENEEESDREERGLTDL FKTEKAAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYAIHLDY* MAGRLCVVMAITITLVAASVTTPSQSLLTERQENDTSGRFLRVASTLKEENDEERAAAPVEAVTSLIKNKNVA
<pre>>scaffold_25_F1 242 >scaffold_12_F7 08 >scaffold_25_R6 73 >scaffold_74_F6 14</pre>	FVGKYVLNPRPGPPVRK* MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV* MRALKLLLVTVFLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK STSGATNWLKDKKLAAKLKLRALPMNLDKSVQKTMKDGIDPDRVFTLLNLQKKSNRKINGFRTGEYNLWKE LTVEWTKTYPNWVSNISKKPK* MRLSFLLPVAMAAIYCATCNATADSDQNKVSMVQSLDARLNGQADGTRFLRAHHENEEESDREERGLTDL FKTEKAAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYAIHLDY* MAGRLCVVMAITITLVAASVTTPSQSLLTERQENDTSGRFLRVASTLKEENDEERAAAPVEAVTSLIKNKNVA NAAVEAVEKKYPLVIPGVTDLLIKARMKRWFWKRTLPTKVFKKLGLRGHSEDRLKNHPFRKFYAEYLEKWKD
>scaffold_25_F1 242 >scaffold_12_F7 08 >scaffold_25_R6 73 >scaffold_74_F6 14	FVGKYVLNPRPGPPVRK*MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEKAAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV*MRALKLLLVTVFLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKKSTSGATNWLKDKKLAAKLKLRALPMNLDKSVQKTMKDGIDPDRVFTLLNLQKKSNRKINGFRTGEYNLWKELTVEWTKTYPNWVSNISKKPK*MRLSFLLPVAMAAIYCATCNATADSDQNKVSMVQSLDARLNGQADGTRFLRAHHENEEESDREERGLTDLFKTEKAAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYAIHLDY*MAGRLCVVMAITITLVAASVTTPSQSLLTERQENDTSGRFLRVASTLKEENDEERAAAPVEAVTSLIKNKNVANAAVEAVEKKYPLVIPGVTDLLIKARMKRWFWKRTLPTKVFKKLGLRGHSEDRLKNHPFRKFYAEYLEKWKDAQAHLNVDWGKAPAALPGTITTGNTVLKT*
<pre>>scaffold_25_F1 242 >scaffold_12_F7 08 >scaffold_25_R6 73 >scaffold_74_F6 14 >scaffold_1_F54</pre>	FVGKYVLNPRPGPPVRK*MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV*MRALKLLLVTVFLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK STSGATNWLKDKKLAAKLKLRALPMNLDKSVQKTMKDGIDPDRVFTLLNLQKKSNRKINGFRTGEYNLWKE LTVEWTKTYPNWVSNISKKPK*MRLSFLLPVAMAAIYCATCNATADSDQNKVSMVQSLDARLNGQADGTRFLRAHHENEEESDREERGLTDL FKTEKAAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYAIHLDY*MAGRLCVVMAITITLVAASVTTPSQSLLTERQENDTSGRFLRVASTLKEENDEERAAAPVEAVTSLIKNKNVA NAAVEAVEKKYPLVIPGVTDLLIKARMKRWFWKRTLPTKVFKKLGLRGHSEDRLKNHPFRKFYAEYLEKWKD AQAHLNVDWGKAPAALPGTITTGNTVLKT*MKTATAFATVLALIVATNGAHVSPNTPTLRGLRPTADTPSVEDDKEDRKDHHHVKKVKKIAIPVPVPVEVPQ
<pre>>scaffold_25_F1 242 >scaffold_12_F7 08 >scaffold_25_R6 73 >scaffold_74_F6 14 >scaffold_74_F4 31</pre>	FVGKYVLNPRPGPPVRK*MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV*MRALKLLLVTVFLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK STSGATNWLKDKKLAAKLKLRALPMNLDKSVQKTMKDGIDPDRVFTLLNLQKKSNRKINGFRTGEYNLWKE LTVEWTKTYPNWVSNISKKPK*MRLSFLLPVAMAAIYCATCNATADSDQNKVSMVQSLDARLNGQADGTRFLRAHHENEEESDREERGLTDL FKTEKAAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYAIHLDY*MAGRLCVVMAITITLVAASVTTPSQSLLTERQENDTSGRFLRVASTLKEENDEERAAAPVEAVTSLIKNKNVA NAAVEAVEKKYPLVIPGVTDLLIKARMKRWFWKRTLPTKVFKKLGLRGHSEDRLKNHPFRKFYAEYLEKWKD AQAHLNVDWGKAPAALPGTITTGNTVLKT*MKTATAFATVLALIVATNGAHVSPNTPTLRGLRPTADTPSVEDDKEDRKDHHHVKKVKKIAIPVPVPVEVPQ YIPVPVSVPSTVVASSNNA
<pre>>scaffold_25_F1 242 >scaffold_12_F7 08 >scaffold_25_R6 73 >scaffold_74_F6 14 >scaffold_74_F6 14 >scaffold_1_F54 31 >scaffold_37_F3</pre>	FVGKYVLNPRPGPPVRK*MRLSFLLPVAMIAIYCATCNANVASNQNKLSMLQARLNDEAGGTRLLRVHHENEEEADREERGLTDLFKTEK AAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYMTYRDYV*MRALKLLLVTVFLLVSLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLKK STSGATNWLKDKKLAAKLKLRALPMNLDKSVQKTMKDGIDPDRVFTLLNLQKKSNRKINGFRTGEYNLWKE LTVEWTKTYPNWVSNISKKPK*MRLSFLLPVAMAAIYCATCNATADSDQNKVSMVQSLDARLNGQADGTRFLRAHHENEEESDREERGLTDL FKTEKAAVKKMAKAIMADPSKADDVYQKWADKGYTLTQLSDFLKSKTRGKYDRVYNGYAIHLDY*MAGRLCVVMAITITLVAASVTTPSQSLLTERQENDTSGRFLRVASTLKEENDEERAAAPVEAVTSLIKNKNVA NAAVEAVEKKYPLVIPGVTDLLIKARMKRWFWKRTLPTKVFKKLGLRGHSEDRLKNHPFRKFYAEYLEKWKD AQAHLNVDWGKAPAALPGTITTGNTVLKT*MKTATAFATVLALIVATNGAHVSPNTPTLRGLRPTADTPSVEDDKEDRKDHHHVKKVKKIAIPVPVPVEVPQ YIPVPVSVPSTVVASSNNAMALKIATLLSCLSLWLLFAETFYWKETLSLEVSFAFRPQTSLACVDSTPSNRRSTYRRWLRVNVPWQRLRTQS

>scaffold_7_F13	MRLTYLLLVIATTLFSCNNATASGSSSAKLVISDSADPLHNGDAAEIVGKRILRGSNAEKNKYFDEEEVEEEERA
70	LPNLSKLDDEANLKILVDIFKYFDDAGVPGLKVFIKKEAMDLDDLASMMTLYQRYRGAGADDFMDLMKKKI
	GVAS*
>scaffold_2_R34	MYKPLLVTVLILLASYTAASLELASETKALDAVPRDRRLLRANTEEGVDGTETQHEERGITTKVGNLIKDWKIK
10	QMFKNGKSFEEIFNKGVTSGDVWDAFKIAKLQNKMSLTDLQFNPKYANWFHYDVWYAWRTAK*
<pre>>scaffold_16_F2</pre>	MRICFLLLPVVALIAAVSGSSVNLRASHRIQSRDVAQDNTGGRELRGDLNTGEATEERAKFDFVKKLVSKAKG
450	DPLESFAKKQTRYVFNGRVFNDLYKKFPDPDALYTTLKLSKYENRWDKFGQATPSFKIYGKFRNAYVEKFPD
	WTSKLNWL*
<pre>>scaffold_24_R1</pre>	MASGIFKVIALLVIFAVATTVNAEPVRHLRVEPDHEPVTRKLSRKVPASMLKNFWGIVRTNLENQPTINKAQE
723	MYAKLSHAGHVKPPAFHVPAKAGEDVLKHNYRFLDFA*
<pre>>scaffold_21_F1</pre>	MMVVAVRALLVASDLLFLCLHENLGADDWVRNDRCHCLSNSSQDEGLQCRQRTHALGIASRHLRVLQLEL
267	QFVEDSVVDRRVTHQYE*
<pre>>scaffold_11_F6</pre>	MIATATEVLLLISGLTHRGVASAVGTDNAEGVIDTLADLVTPCKSQSFDVRVGVGGLAAFSPTPITKGSSSTAP
83	TRSLRSTGLHKPKPTRASNAPTKAGPKPRQKTAHRDDSAETSTNTKSMLTSSSRLGGGDLRAFRDTVGTKRA
	DVDGVLDRCEASFAKAKRTRALGASDELKQKISSIQAASSGFGGSIMKVMLFLREEN*
<pre>>scaffold_475_R</pre>	MRLSFVVLLVATISSLVATSHALPHEQSTVLNVAATNPDQSVTEGQTSEKRFLRSASLGSQLLRSETFKKSKFK
2	DLLDENVSSFTFFVDMLKGKKKYRGIYNEYAAYRKAHNAYP*
<pre>>scaffold_50_F3</pre>	MRFFLSLAVMILALVADGDAKGSQVLVSNFPNFQQYGASTGVVSTTRLLRARSDEERGGGISVPATDKLSKLL
14	KLPKVTDEQLQQWLSKGKAAESVFYRMDLGNTRITKLFENTQFLRWLQYADDLSDSGKGISAISILSTQYGD
	DTLYKMIDSAKKTTNTEALGLRLQSEQLKHWVKIGKDPDEVFKLYDLNYAGRRILSTSQFSAWTKYVDDLNA
	KNEGAFVLIVPTLRKYFSDDELFHIALAAKRSGETEAMGAKLEDAFVQFWTRRKDTPDNVLVELGLKKSMETL
	LESPLLNILTKYTEVYNVNYAAKKTTVIETLTRTFGDDKVARMLLAGRTESTTKKIAKQFQTDQLEMWLSSGQ
	SVDDVYKLLNLPSRRDLLSDFGNQKLFNTWVTYINTVSLKNPEKTSAMLSTLASSFRDKPMMQILEAANKFPS
	GSYQGGVNKIIDKAMKDPSIMKIASLAQKEQLNEALIRWKYSPEKLFRFLNVGEAGEKVFSAPKFELWVKYL
Scooffold 111 D	
25Callolu_111_K	
LJ4	
5/	
54	
Scaffold 22 F1	
370	PLI DSAHRPIROI SGRTRHARI HROKAASIORHYROHPERSYSEI TAVYI I HPEPPRHI PVPOVEOHSOORTS
570	RSSPORCSI PPSRPPARTNI ATSAC*
>scaffold 48 R1	MVKOPFLILVCLDLGFCRVIPVWSADAPAVTNGSVARSLRAAHLELPLHWLLSEPRTLAHCAFCALOSTPAVL
250	AKLERSQTWNRWPFSRK*
>scaffold 12 R1	MRLSYFVLVAAVCIFACCEQVAAVKDSQGIAATHTDSLRKIDNRFLRGREDVMGDESDDEKDSEAENEERAL
094	PKLDVAAEKLMTKAAAKLTRSKSLSSLAKLDDEAFHRAVVTNNFFTLERIEKMGYNPDKMFLKMKEMGKNA
	PLTDPQKVLLEYYSMFWKIKYPSWVSEVPHMF*
>scaffold 16 R9	MRLSLVVLAAVLFASGTAVSSADPASVAAVHSSRVLSDEDKRFLRRHPTEHDNEERAFGQNMFAALKLSKM
43	KTDAEYRVKVFHRWKKHGYTADDVAKHVPAKLADQYSHHVNTFTTSGWVMNPLRIFSPP*
>scaffold_1_R45	MRFPIIALAALVAFLSTLDASSLLVQAASDKLNSGRSLRIAAGTGLEQDEERGSLITKIRTTFSKKARVDEWAKA
69	EKSDDFVRKALKLENVADDHLWAMKNYPYYLRFLEKTEAKKINDWLIGDVPTYGAWVKLGLGSIDDIEKVKT
	TAAFKTYEKFVKEYDDQAIALWKDYKIPIPVAKATSPTEMNARMDILAAAKRGDDYAKKVLGLDNVTGSALIS
	HVNYQYFEHYRRTVKRLQDPKKKLNRLPTITER*
>scaffold_24_R7	MRLQAVVLLVLSAALAVADISAATSGTGGHHVNQNDVSANRLLRSEADVSDGEERAATWLQSLKNLVKPN
35	TAPKTMTKLERENPDVAKLVGKSLDGAFRSLKLNKIKSVDDLFTSKNFETFNGFMIYRNSQSRNKPSTIAKFFT
	KKLGDEQASKLFSQAAKSSNTKVKKMGESYQTQLLRQWGKEGKTLKQVTKIDASLQTRYQTVLDDLARAAK
	KKAEQAAAKAKAAANTAT*
>scaffold_1_R25	MKTTGSIMLTAALLLSDSVCTAYYLRGTGIDEDHALDKARVATVQVRYLRKSVEPSFLDSSSTINTSEHHVDNT
2	KAINSSSPVSATKLPDVTLSDLDETERTSTVSPTNVEIGISSGNMVEFMVHPSERSSSVVSDIERRVVPMAGPN

	RARPTERNFWLGSHYPSRYHMKPQHFFMNRQMDRFPVNLPLILPRARQRQHRYFVGV*
>scaffold 29 F1	MRCILYIALALAAFARSDAVEAFTNADLSSKISHDFAANELVNESSQKRFLRVDYPKSVDTEERMKISSLKKIIKE
478	LDVKDAKAKKLAAIKDLAKARKDAQKLATNVVKSKAQRTRTKFSSLEYDNFMRMFKSNLAPEKAKSTGKIKT
	PEQFTRYEDFYETAKAADILTRMAHA*
>scaffold 93 F1	MRLHCIVVLAVIAFATNGNEVSAGKSRVAITTTGALDTPTTRLLRTQYTDEERAFGLNLLPGSKKISSFMTDKK
27	LSKYLKSNQEFDDVFIKLKLDKAGDKLFENPKFLAWAQYVDDFNQKHQTQNSMLPTLVRQFGGDDLSIMLE
	KAKQADKTYGVALRLQGEQMKLWRREGLTTDMLFKIYKLDDGATNLLENPGIKIWMRYADELFPGDSTLLF
	KKLQKTYSDEALSKILINGKTVASSEYGEVGDGFAASSLAEGPCASGESFPVAVTQQGGRCVW*
>scaffold 13 F2	MLVERDLLATEASQTDLLLLLLLLGEPRRVCNFHWCSTFLLRCRLLRRCTCITSTRLLLVFLTSLRWFLRLSLLW
760	SL*
>scaffold 8 F32	MCLQSIFCGAAVLCGEWLVIRAFAVKCRLRSHSLPPAHAALSRGVLPRHLKCLLRYSSYVSTKWHLRYVVQRR
13	RWRNLRHQSRLIFLFVRLPLCQDKSPRDNHQFLVQVWDSIPLRWFGSRSQTTTPL*
>scaffold_18_R4	MIHICTFILLLLSGERDFGVNLNKPFNNHLPLDLPPFSGNHADLLIVCLHKIIVSGYEKLNSVYNCFLTIICNISPYC
28	KKLNMVSAVRLLRLFKLFAQTRYLFDNEANHHLVFFLLETFDNIIQYQYEGNQQVVYAMIQNKNVFYQLNDL
	QLPPIRASTEGKKEATEKTEEEKATDGEFVPTVEWLSAWKKKLPLTTSLRLLQYLIPQLEDACKKAGGSLDEDA
	MLYFLRTTTMVGLLPVPHPIVIRKYQINQFTHLWFTTFTWGVIFLRNQVLPLFDGGAITLFTISVL*
>scaffold_8_R13	MRLSFVVLLAVAVSSLVATGHALPHEQPTVSKVTAANLDQSVTEGQTSEKRFLRSAAGIGRQMLRSESFKKT
70	KFGEWLKDNLSSFTIFVDKLGGKEKYRRLYNEYAALRKASGQYP*
>scaffold_2_F23	MMGSVSAFVVWLKSLLNLPTTCTCSCPCDVQTCYRRELLLLLPVRACLRRSLLRLLLRLVPRRLEWDLLRFLR
3	ADLDLDLS*
>scaffold_4_R23	MRLTFLLPVTMAAIYCATCNATAVSDQGKTPTVHSLDARLNDDTGDRRSLRAFKEKEEFDTEERGFLEKAAV
6	KKVGQGNYERP*
>scaffold_29_R9	MRLASILLVSALLLCGSVFSTEGKQAHTVKNTDGKRLLRSDDYDEERGFNWPWLKKIKQLDETTAIKQLATDF
03	ADLKSLNREALDVFHLMREGGMSPKKATYISNLFAKYMENPRLYH*
>scaffold_50_R4	MRFSYFLLLTSAAILSNSNATATVSGEGHVMTSADAPARALETNNGKRSLRYYAADEDDKYDQKNGKYDDD
62	EEEERNFTAAQLAKWTEKAESWVSKNRTPAYIKDKLTGMNGLMTAENRKKYELFTAAYGRANPHALDRL*
>scaffold_25_F4	MRPILLHAEFWTLWLVLAWKACCLREPSGLAFTTTYWSMTKTSYRLTWITSFASTRLSLLRMTTTRRRLRRLL
87	HLLQQTPRMYPLYLKRMRVKRALYRWLQLTCGASTDASARFS*
>scaffold_7_F31	MRSLLALAHHLILCLLPHPAYFHTLMSASRPLDTSMGLGFWLLPSSTSVRSKIQSVWAGDGGTRPLRSILIADS
0	TPPPPEAATLPGDE*
>scaffold_50_R8	MRLTYILAVAVVATLHSSATAFSSVKDSNAADSALAEGGRMLRGEYGNDALIADDDAGKKKVSYGDDALDY
12	DDLDDDDLDDYFEEERTLGDVLKKLNPVRAVKKSAEKTKEHAAKVKEALKDAADYQAMIARAKEMVNKD*
<pre>>scaffold_30_R3</pre>	MGVWFLLLLDVALVAATSSVRTALFSLPVELCSFVFLTDAGRGLRPVEASLVLAGDAVRGPWSKFCDPFGAK
24	LDSTVRGGNGPL*
<pre>>scaffold_111_F</pre>	MKVFSILASAAVLIVMTVLVDAQRRLRSEASTATDSTVSTADFWTLSSGSGFTPGNVEAQSNFGALDKEGST
98	GGNRPSGSVQGGAGRGFGGSGHNGMQNVGSGEGLAHGSGSFGGHGGFGRHGGHGRGEGSFSGGDFA
	GSFGGHDIGGGNFGGSGNGEMPSIGSGAVPTTGSGGAPVAATTTSTTSSVAASSSA*
>scaffold_25_R8	MRLTQVLVIAVASLLFARDTVIATSNQAKISKMEQSTPSQRLLRSNKYPIEEEEDEFEDTMDFEEGDYTTTDKE
04	DLEERSPLSSAVVAKLDDIASGWGTSWARVAMGQSSISEDKIKALLALRDAYISGSKKAKQAARMAVLRANN
	S*
<pre>>scaffold_81_F2</pre>	MRLNCIVLIAAVASLSTAAATSATTDIKPVRPAINLQPPPLVVGRLLRTVQDEERGFTLPGAGKLADLFESTALK
85	LTQSARINTWLAKGTSTDDAFIKLELNTAGSRIFENPKLLTWAVYVTKVEKKNPEEIILARLSKQFTEGSLAKMI
	ASAKLDSKTEGLATILQAQQRQVWVDAGKSSDEVFKLLQLDEAGTKLFKNQQFSTWTSFVDAFNRKYPEEA
	VSIFSKLAKTYDDFTLWKMLEAAKKVPKTEIIASKLQAQQIDAWLDAGKSTDEVFNLLKLQRTGDKLFKNSQF
	FKLLKLNTPSLENLIGPRLDAWTSFMRAFNMANEGKETTLIATLTTHYKDRGLAQLLQEGTKFASTKKIAEELQ
>scatfold_50_R1	MIVSFKKTLFVSAVAVALAVSVEGYSGAVESTYETKRMLRLEAQDVAEEEMADDSECGSLEMAEDDSECGSL
069	EMAEDDSECGSLEMAEESGTGDKGGDNGGNTWTQAPSTGDKGGDNGGNTWTQPPSTGDNGR*
>scatfold_29_F7	MRLVSLFLVIIATLVATGNTLAVEHTEGKITSSDTLLQPTATFNPDTVQTRRLRTPESPKDGEERRLVNLDIAIK
68	DAVHEVRKLTKWKLQFAVWKEIGKKPQKLIQEWGMKYHM*
<pre>>scaffold_10_R1</pre>	MRLSVILLVVAAFVVALSPTAAASEANTVVAAPKLHESTATARFLRVRPEDDYPVKDERDGEDSDDDKGNDE

Lamour_MPMI_SuppFigs/Tables Page 47

133	ERMFSFLEKKTALSAFKKLVSESGDDLVMAVSSLSKAEFVALFNQGQSAMAKLVPGFRPGMSLDDFVTVVKS
	AGLRRDMEDALMVGYGKYLAHLMD*
<pre>>scaffold_50_R9</pre>	MHLLYLAGVAILSFAAGATDARVLASDDSNQYRAHAANANIVSTTRLLRARSVIDEERAGGISASASDKLAKL
34	FKSSKVTDEQLQQWLSKGKAAESVFNRMDLGNTRITKLFENTQFLRWLQYADDLSASGKGTSAISILSTQYG
	DDTLYKMIESAKKTTNTKALGTRLQTEQLKHWVTIGKDPDEVFKLYGLNHVGSHILSESQFSAWTKYVDDLN
	AKNEGAFVSIIPTLRKHFSDDDLFHIALAAKRVDETKTMGTKLEDAFVQFWIHRKETPDNVLVELGLKKSAETL
	LENPLLKILTKYTDAYNVKYAAKKTTVIETLTRTFGDDKVARMLQAGRTDSTKKKIAKQFQADQLEMWLNSG
	QSVDDVYKLLKLPSRRDLLVDFGNQKLFDTWLTFMNAISIKHPEKTSAIFTTLAPTFRDKPMMQILEAANKFP
	SMEKAATKLQLEKAQSIFSTGISPYEAFKLAALDDVGDSVLSSPLFKKWMLYVEDFNKKNPANKESWFLPLRG
	NYQGDRLDEIIDKAMKDPSTVKLAKSVERERMKEWLEKWEYSPSTAFRKLHLKTAGEKVFSNPNFELWVKYL
	DDWNKAYPSNKMTMIDGFRDNYNDLNLVRLFTKAEKTPSSKKLASQLKDALADKWVAEKKTLDYVKSWYS
	HVPSSDDMLDRFTKKLNNA*
>scaffold_2_F36	MRLLFWTLLVGLATSGVQPIAALSKVADTSETSTLDHGQSTIIEGTARHLRLAPHEDTSITDVLVERGLKVHDE
47	ERGNFAAAAGWTANKAFNGLGKIVGHDNSNKLKTKVAEFFFSILRKKGKTAGDIFAHARKQTDPSQRAKFET
	VATMYQQYLTKVL*
>scaffold_644_F	MAGRLCVVMAITITLVAATVTTPSQSLLTERQESDTSGRFLRVTSTLKEENDEERAVAPVEAVTSLIKNKNVA
7	NAAVEAVEKKYPLVIPGVTDLIIKARMQRWFWKRTLPTKVFKKLGLRGHSEDRLKNHPFRKFYAEYLEKWKD
	AQAHLNVDWGKLKATAALPETITTGNTVLKT*
<pre>>scaffold_104_R</pre>	MRPNLVVLAAVATLVSRCTVVSATSSTNQINLSNLDQALDIVENHAGNGNRFLRSANIVDDSDDDIDSEDEE
67	RGGKTWAQKFAKWNQRGESANDVYMRFGLEPLVRKAYKYGQIGQLQKNEYYRKWAAYSAYLKDKGLN*
<pre>>scaffold_36_R1</pre>	MRFFMVAAVAAFAFASSCETASAEAPNAFLIDDNTNAVFSRYLRSTQTDSEERAIAQVLKSEDRTVANVMIK
210	YQSLYRAKITPKQAKLMLGVSDNMVEFTKMTRPIQRFYTGYYSYFTTMEKRKKWKKELKNQVMV*
<pre>>scaffold_379_F</pre>	MNIAVLLCFFAFPSGLFFSFRRLRAASLAPFLLSDIRWHVLGGPSLQHFTPKQQQLHALPLQAAYLTSSAAFCS
2	QPLHRRALISLRL*
<pre>>scaffold_20_R1</pre>	MLGFLLLVMTVGSTTELILGGWSSAAITPDTRAVLVQALSTTNVCVDSIISVRSQVVAGTNYEFRINGCRGLR
237	GGNCVRASCVAPRTFVVNVFDQPWTRTTRVMSVSEI*
<pre>>scaffold_24_F4</pre>	MRLSFLLLAATFALLSSGTVASPTTNDEPISSPNTVLSSETEGRHLREHKNSIDDTEERGFNPAKFDRLMNERS
23	YRQTRFGNWVDKPYTDMDVYKLLRVESNPNYRRIFHYYQNYLQEFAPHLITP*
<pre>>scaffold_2_R38</pre>	MKFVSTLVFMTVALQSSCSTAQIDFVRSYDHHAIDFSTSSKPTQPSVADVNTVEANQPLRSGNRRLRDGSAA
00	GIVIVGRNGGGFEPLRNGETPD*
<pre>>scaffold_11_F1</pre>	MKAKLLLVVLSFLLAVTSVSAERHLRVADDANPLKPYEALNAVIEKFKNKAKGKPERRRLTGEKAFDKFVKAIA
235	VLKEKDHARRVEEAN*
>scaffold_55_R4	MRLFIKTLVALAAVLLATSTEAAKAVQTGGNEDVVQSSHILSRESKRVLRSEYDEEKLLEDDEEELLENDEEERK
	NGANLFTDAKMKKMLASDWYRYQVARRWKERKYTWQNLPPNVDPNLVAFYKGFRERHG*
<pre>>scaffold_55_R6</pre>	MFFLVKQLSAVNHMYRFSLASFLGLFKGTLATKMESSSTKDRILRLIPILEHKVLMFVGRALFKEHRPMFGMH
23	LVHGMHPECFDKNEYEFFCGEVVELERGSGGHSTLPEWASPERKEAFTQFVETLPRLTQLCKFESHDMWIR
	WSKSMECEQNFHPKMDKSGSAGGLSPFQKLLVVQALRPDRLQSAIIQFICGVMQLKSLTPPSLDFKVIGTEE
	ATNTTPVLLLTTAGADPSKELEEVATSVVGKGHYFEVAMGGGQQEKALNLLKSTAEHGEWLCLQNLHLVVA
	WLPVLEKEFSALNASHKFRLWLTTEPHDAFPLVLLEQSLKITFESPPGMKKNLQRTYAAWNPAFIAKGNSAR
	AQLLFLLAFFHALLQERRTYIPQGWTKFYEFSFGDFRAGSNVMELACQTSGAAIDWQTLHGLMENAIYGGRI
	DNPYDLRVLRCNLTEYFSQDLLSGHKSLTRGVKLPQSTQHADFLDIIDRFPDIDAPAMFGLPDNIERSMQRSL
	SGQVIAQLKALSSNEAEATTFDREKWRAQLGPLLETWSKLTTGFQLEGTSLSSSTGKNLQAMTPADAFVALE
	NEYALDLVQQVNSILQALKKVIYGTGLLTPAIQTVAKALLKGIVPTEWAA*
>scattold_325_F	MLSVLSLALMLAVVTGAFETSRDVKNGVVTPDSRRLLGFTADQEERKFGGPATGTDHSNSHAWENFKAWF
4	
>scattold_50_F4	MGLFYLVCVAILAFVARDATGDRVIVSDGFNQHRENAARASVVSTTRLLRTKSVIDEERVGGIPVSATDKVAK
3	FLKSSKVTDKQLQEWLRKGKTAESVFYRMNLKTSQLFENPQFLRWLQYADDLSASGKGTSAISILSNKYGDEK
	QSVDDVYKLLNLPSRRDLLGDFGGEKLFDTWLTFMNAVSIKTPEKTSTIFSKLATSFEDRPMMQILEAANKFP
	SMEKAATKLQLEKAQSIFSTGVSPYTAFRMVALDNVGDSVLSSPLFKKWMLYVEDFNKKNPGKEDSWFLSL

	RVNYQGNRLDRVIDKAMKDPSTMKLAKFVEKENMKEWLVRWEHPPSVAFRELHLNKAGEKVFSAPKFEL
	WVKYLDDWNQAYPSKKETMIDGFVDNYHTLDLIPILAAAEKVPSTKKLASQLKDALVDKWVAEKKTLAYVKS
	WLNGVPSSDDMLKRFATKLNSA*
>scaffold_11_R1	MRSYHFLLLATAALLSSCNATAAVLCEGQVMTSADAAVPVRALGTSNSKRSLRYYDTEEEDETDKHHQKND
430	KYDEDEEERTWSAAQIDKWTAKADEWVDLGKTPAYIKAKLTAFNGVMSDKNRKKYELFLAKWGRANPDEF
	GRR*
>scaffold_22_F2	MALCSCICTGIACFPVSSHFAVYFHLSPHSLLSSLRRLCHTDTAERLLRRVLLIARRTQEQRRVAEWLQVATQR
08	AERSGSRTIQHGHITGAGWTCLLLLHRWVISLFPLRLHSSTGHQLVSL*
>scaffold_29_F1	MRSVYVVILAVLALLCNDALAATSKTDLAEAELTKNERNRSLRGEVIADEDSYAIDEERVQTSGISKVAKELKLK
480	KANLKKVSAALEKAKSNPSKLAVNAKKMKAIAEKKILAKLSSMENDNFLRLFKANFTPQRAKSTGKIKNPEQF
	AKYEEFYKLAKLIGLK*
>scaffold_3_R39	MLPTELVLFAFVVGLRQAGYAPPPATTSEWPLLFSPLGGLVNWHTFAFLRINRFLRIVHLRPLSDQLQRFLLYD
1	${\tt RRLKRLTPGICYLVRLALDFLLGTHWLSCLFYGVSYLAYDDGEMSWLTTPDMLAFGDGVRDLADIRKVPLLQS}$
	YLRTYHFSIGAITTVCYGDIIPMNAQETEVTMAVIFISVALFSMLSGGFYKYFDMELGRRAEYEERVAQVGHFL
	KFHRFPSDTWRQMQVYFALSWRESRGRRERELLSGLPPSVRQDLAQHVHASLLKNVALFTRCDPTFARAIIA
	ALQHEFFVRNDVIIQRGDMERSLYIVESGIVLISAVRKRQVHAQAGGAEEEQASNDPVGADDNRSPGVSVN
	EWVRASVGASLRLRKAKKAEKGQANTPRQRQKSLVALMSPTGTITDINNQTEREEKIYKGPFDYFGERSLLFG
	TPRNATCMALCVTSLFVLTSARFEAILDEFPHERSNSVSAWVMTRTPSLPPDERPIDN*
<pre>>scaffold_7_R68</pre>	MKACVVLPSLFLPCPSTSAAEPTDPDEDNSTPHLDALERLRRRQRLVNRLRPDQEIDTHDSIRWH
1	VRYGPERIPSPPQSPPTPDFIETYGDSEYAVSDDNSSDSGTESEQSC*
>scaffold_3_F35	MALVVFFRILLLVCLQIGLDEEAIEDHPTRKTRRQLQAGPHEASSEERVTEETHPEPMARHLRQFPTQHFDIV
36	QSIFGRIGQQIRRAGIDS*
<pre>>scaffold_61_F2</pre>	MRLLFLLVAIATFLIASEAFSTTGDSNQIYNVDSPVGPRQRFLRAHAEVDLEKEMKKMMKMMKRKVTKEDF
09	AKTLKITDQIDDIINKHAPGMHEFMQTPKYQRYSNYMNFLNDMAKKPEYAALVEEIKAKSRAQVALKTFRKP
	TTSQNRWQIIFASLKGKRIGK*
<pre>>scaffold_23_R4</pre>	${\sf MQRRMSTPFFLMITTIGDAQGEVEWVMIPADNSPLIKSDNFSRNCGATRRTRWATGVALPTSMSQTAPRA}$
90	AGGRAFPFPIRRCLRFQMICLNRSICSGVLPSASSIWISSILTSSVVALTSGSTAI*
<pre>>scaffold_25_R1</pre>	MYRILLLTTFALLCGFSNAVNSDARLLRVVDPTEEKGGFTTYVKDSLTKWRINSKIKSWVKNKKTDEYVLKKLG
22	LSTLTGKDLVNAPKYSQFQDFKVGMWLKEATPTTTVFNTLGLNKVEGAVENADDFGTYVKYVITLGEKADN
	YPLSQWPRLFGGGSLEQLELKRKILRLQKRNVLEINFMLG*
>scaffold_640_R	MRICLFCYLLSLTAVVSGSSVNLRASQKIQSLNVVQARMSGRELREDLNTDEATEERVNFGFLKADRLKSFAK
2	QMDETHIRTKSMNHKRN*
<pre>>scaffold_38_F1</pre>	MRSYCVLLLVYTAVMSTAGTESTAMDSTQPSTQYDIVLHVRTLRTGEYTVSEERAAGASAVESLGKATTSVD
173	DVETWLKRGDTANKVFKALKLQKAGDNLLDNPQLSTLMRYLRLFNEANPTKKTSLIATVTKHYRNHGLTKIIE
	AGLASTSKKTVATAKHLQTEQIHYWMAHGRSPESVFGVLIQYTKDALRRSVLLKLDKLLPFTWISSDLFEKPGL
	KTWIRYLDEFNVRNPKQKTTLISILSKRYNDKTLANMLIAANKAESTSSIAKRIQSEQTLQWLSKGKKTSDLFT
	MFQLQKAGATLFQNPLFPAWIKYADDFRVIHRDTQLETMTTLMKHFDDDVLAKMIMTAYDVPSTKSLASR
	LHAELLRGWQWRQETPDDLYILLKLC*
>scattold_25_F1	MRLSLLLVAAAALVANSDAAPQSTSSLTKFSTDVAPVRSLRGASKTKTEVDEDDSFDPEEEERGISMYGKLDK
656	
	WKKLQLSPLAVSRDLRKKGIPRTTDNVMWEAFRLYSATSGRKFNSLRPTGL*
>scattold_77_R2	
49	
>scattold_1_R40	
/	
>scattold_43_F1	IVIKWLUKUUVWLUSHULIVIVVSPELLVUCPULSDSNVETSGWTMMLLGSUMWKSSLUNLEWLAAISSACL
201	
>scattold_6_R23	
12	KKAGKVPSGLKNLLLKNIDEFAEHSKLAKKLSGLKLYKDAGLEKMSLSTLRQLDDIEVKRVSDIKNGITGNKDTP

	GGMRRKMDHVVGDVAPAKYLTSHIGRGDQLYGADGSRLLSSAVVSRPAEQGGGKVLLISSSKPEKGDWLLP
Secoffold 405 5	
>scattold_405_F	RSPLSSAIVSKLDDIASRWGTTYARVAMGQSTISQNKIDALLSLRDAYVSGSKKAKAAAKIAVLRAND*
>scaffold 9 F15	MPTU VVIAVSTAMAII DMTRTI RIRISICWYTRSIEELI SLPL RWCDSRU INNSILIHVHRDREIRCHHGRMV
5	WVILFLFSYRTVVSHWRSSGRRVSRVVFIFTMRNSIHR*
>scaffold_42_R1	MSKVFLLLVLSVFALVSCDALSAPVGSKLSLSKTDELNAQPIDAKRMLRAQEEPTNAADEERGMTELANKFK
91	AWAAAIKTWVTNSKLVQSMNNKLASLTQKGRVGQIEKLLKQDNVNVNVLYQNKVKPDELFLALKLDPKLKL
_	IADAPAAWANNPGLSMFYQYATYYAKMTTKA*
<pre>>scaffold_73_R2</pre>	MRLLYLAGVAILAFIAGDATGAKVLVPDDSDHNRAQAASASVVSTTRLLRTRSVIDEERAGGISASASDKLAKL
70	FKSSKVTDEQLQQWLNKGKTAESVFYRMNLENTLYTRVFKSPQFPRWLQYADDLSASGKGSPAISVLSTKYG
	DEKLYQMIGWAKKESSTKALGTRLQTEQLEHWVKVGKDPEEVFKLYDLNYAGWRFLSNSQFSAWTKYVDD
	LNAKNEGAFVSIIPTLRKYFSDDDLFKIALAAKRSGDTEAMGTKLEDAFVQFWVHRKDTPDNVLVELGLKQST
	KTLLESPLLSLLTKYTEAYNVRFATKKTTVIETLVRAFDDETVARMLLAGREKSTTKKIAKOFOADOLEMWLNS
	ESSMEKAATKI OLEKAOSIESTGVSDVKAEKMVALDNV/GDSVLSSDLENKWMLV/EENKKNDGKEESWELD
(()) 000 F	
>scattold_300_F	
13	EPVQTMWEARPRKYKSILFEV
<pre>>scaffold_39_R1</pre>	MMLVLLAICETTIHVAAGTSLLFESDFAFGVDSTTSACAIDRPFLDPDKHLLYSNLTNNLIALARCTVLERLEVS
000	CKDMDQRNKFSRLLRSYYLLHVPVALYSTERTSYHQGRNHIHYHVGLI*
<pre>>scaffold_7_F14</pre>	MSSKAGLISCALWLLLLVSTLGVDAHKRLLRSEVSHDNLEDRVLDNWKKLSLLVKVGSRTKADKLYAKHVVG
94	KITGNFFDTNAFAAWFVAVQMAYAKTPAKAKVDMVSSLTARYGDQALAKMLATTEDDKFIREMKAIQLDN
	WQKDKRTVGSVYKLLKLDKEQDELLQSPLIATWIAYATKLDNEDPLGAVFSTLKTQYNGKDFATMLLNVKDT
	DDSFVVAEKLETLLMKSWQREDKSVVDVYKLLNLDNEGDLFFQHPLIDTLIRYATVVDKKDSFSGVFSLLQAR
	YNEEKMTDMFMTMRDWWPRNILTDQLEDLLLKTWQRQEKTMDDVFKLLKLEQQGDSLFSSKLLSTWVSY
	VAKVETNPYNVVFSKLKSTYGEETLTSMIIQARDMPTADYVGTGRRIRESSVQGLGEGQVHCP*
>scaffold_17_F9	MQSVTSLACVNGAFLSSVAVIAGSDRSLRVLDVGAGGGGRTMRVVRDAHSRAAHTVALPRPTCYTSHPSNF
40	YDLLLSSAPDSTTHLWDIRADNCVMRFCEHVNRVHTLGVAFSPCMRYVATGSEDRAAYIYDIRTGRRLVKLK
	GHTDVVTSVAFSPLHPQLATAACDGTVRFYSSTRSD*
>scaffold 21 F4	MSSRLNLLFTGSFLLMARKEIVRIFFHFLLLVNCSAIHATVSSSSGSKSITATRHLRRQAKPSVSRARVDG*
92	
>scaffold 2 R24	MGLPLMSSVVFPFSLAAGLGGSHATTTWNLADFVSRLTSELPISSLYGVMAGDATHLHHVLAHIGFALFSGV
30	DVPRVTRPSIRTWSODI VGELRRI LRSHALPTEVLEOVTGSNERRSALGNELLRVVEPEVSELVDELVRATSAS
	RAAAFGTNSATEI RTMTOOIVROI RVYARGESTESEDDSDERI KRI I RGI I VWI GMNENIMAREVIDSI VCW
	AFGDNSSIRRGRTRORFENSTDSPAIKRORF*
Scaffold 20 P1	
833	
<u> </u>	
>scattold_40_F1	MVVKKVGRLHLFSFLLELGIDCFADFHCRLLISKVHSIASSASLRPSRWLQSSRSSLIWICSRLIRSSGSIRIIR
027	TCMTNCNRCWRTIRRRSRTLCVRRSRYHCRRLRDSAVTLV*
>scaffold_5_F25	MRFHYLVIIAAFVLSISDTFSTVSATNTVPSRGLRRIFDEERAGGAISVSTSEKLATLFKSSKATDKQLQKWLQK
81	RKPAEDVFYRMNLAKTKTGIFDNPLFIKWVQYADDLSATTSGKGKSAISTLTAQYGDDSLYKMLNVAKQDSK
	SKELASRLQSDQLEHWVTIGKDPSEVFKLYDLNHVGGSLLRNPQYNSWTKYVDDLNAKHGGEVSMIPTLRK
	YNYDEDLFAIVGAAKSVDALKSAGVKLENAFVQYWINDKQTPVKVLAELQLGATPKTLESPLFSLLAKYTDVY
	NVKFPQSKTTMIETFTQAFGIEKVAKMVAAAKETEGKAKKIATELEAAQMQMWIRSSKSVDEVYNLLKLPPK
	TLVIDLGSSPLFSTWIAYMKILSIKNGDEMLQLIKTLSMQFADRPMMQLLQAMEKFPNIGSTATSLQLRKAD
	DIFATGVTPFRAFKMTALDTVGDSVLSSPVFTKWMSYVDDFNRKNPTKEESWFVSLRSTYEGDFMDKLIETA
	RKSPKTVKIANTVESERMKDWLTRQKAPEHVFHFLKLNKGGEKAFSSPNFQLWAKYLDDFNLQYPGEKTTM
	IDSIRANYRDIELMPILNEAAKIPSTEKLANKLQNALRDKWVDEKVTVTQLKGLFGHMPSSNDWIQKYAEKL

	NKLS*
>scaffold_39_F7	MRLHCFLLAVATTLAVLNNGITTTEASSLRKVPASAPIDSINAVQPETRRLRSAETVYRHEDSYKRRPFIEEKLH
U	
>Scallolu_46_F1	
199	
seeffeld 77 D1	
>scattoid_//_R1	MILLCVF1VVIAAIAVSD11A1NAIDDDGVLQ11SKYLLGFNQEERGRVGGPA1DGG1DKVSDWWPNFKAW
78	
>SCallol0_15_K3	
220	
25Callolu_10_F2	
149	RGFD*
<pre>>scaffold_9_R77</pre>	MRGSQSSQQLCMVLLIRILIVRQLRSLAHQEALLPRAHETSFLEHPARVRHRIVAPHSPQAVGYFGYEPLEW
1	MTVLWEIRSV*
>scaffold_42_R4	${\sf MRIYFVLLLAVTALIAAVSGSTANLRASQTIQSGDVVQDNTSGRELRGDLIADEATEERMNFGFLKPDPLKSF}$
07	AKQQMKYVYNERIFNDLLKKFEDPDALYTTLNLSKIKNRSNVNGVKTSRYNLYENFLISYLDKYPNWRSTLK*
>scaffold_5_F20	${\sf MRLSYTLLIIAVFVSSGNAVATTNGRTTELSAMASPNAVASVDTAVGGEKRSLRYHNNEGLEDESDDEALLEE}$
28	EERKYTNMFSTTKLDEMLNGTKMMSRFRKWKARGYNTYNLPAVTQKDKYTWIRQKYRDFLYHN*
<pre>>scaffold_73_F9</pre>	${\sf MRCFYLAVILAFVGCGDTKGTTVLASDFSNQHRDHVSSTISITRFLRTADASTGDEDRVGGVSISGTDKIAKLF}$
5	KSSKVTDEQLQRWLSKGKPAESVFYHGLGEDSLYEVI*
>scaffold_42_F4	MANKHDACQKSLLFLLFLLFSLQLPLSAQSPSCPTESPLPPPSAPDWQAAPPFWPLFSIYDVVTSSSPPAHWT
1	SRTLRPSATQPQP*
>scaffold_5_R21	MRLSYVVIPAAIAVTFASSGNALAAADGSNTGLSAITSPNVVASIDTAVGGEKRSLRYHTNEDLEDDSDDEGL
32	DDAEEEERRGNNMFSTTKLDEMLDGTQLMSRFKKWQELKYNMYNLPDTILASKYDELRKMYRRFLYYN*
<pre>>scaffold_44_F5</pre>	MRFSYVLLAAAAALLATANAIEADPRTRSLRAHKSHEKSQDEERAFTYTFNFSLWDDLFNSLPEQFQRMRKE
78	PWYLRRIFRSWRSGMGTSDEAVAYMRSQGLSQKAIDQFEDAYIKYRAHKLAKGK*
>scaffold_6_R32	MRLAIKTLVALAAAILATSTEAAKAVQTGGDVHVVQSSHILPGENKRLLRSEHDEGKLLEDDEGEDDEEERKY
03	G
>scattold_11_F1	MKMTALLTVLTVMLAFVSADNSPRQLRSTLTTSIQDTDGNRLLRSEGLISNLLGGLNSLLNNQQSTNPPTTST DPTTSEONPDEIDLLRSRRVS*
>scaffold 31 F1	MRWYRVI I JAVVSEVISAEOSESSHIRTRSI RGNAVNYSEKDVEDGNADERAGEKEVIKKI TSSKTMKKI SRS
104	VTKAANSESEKISPALPIKAKI OVWSNTGKSVOEVROELGI TDI AFAALKKTOSEKYYDDEVTSOLPIWAKKD
101	I TPDEVIEOLGMKGI PAAWEKADPNEKYYDNYLKARVPYWSKNNVEVGDVVKMI NI NTI SGAARREAVNE
	OYYDDEI VSOI RVWVDKDI PVGTVMAKI DI DKI TGKEII THPNYEYYKYEVKNRI KAWATEGTSI DDVAVR
	LGMGDI HGOVI KAHPNEKEI EKYOAKAEOYOOEGWI KOGVTTEDMWNDI OVYRVPMSII RRSNTYNTY
	KNYVNVVDNYIIRMRKKGVPLDKLPGLTSKDATPHELREKTLIWTSAKRPEWYVKFALGLDGLGENALKEAA
	NYQFYAYYLEAVKFIA*
>scaffold 12 F4	MRAHFLLLAAAAIFLATSDVASANQAELSKVASPQSIETANIAPKRFLRTNKYELDEQEERGIGALDDVIVKAG
82	KTKMSNAELKKLVTPKVLENALSDRNMKLFKDLYAGKVSLKRFAKVMKNNPNKEEVLKSYRWFRNYHISQA
	KARTAV*
>scaffold 30 R1	MVSAALASSGSLSGSLVAEPLLLVVLVSRSDSVALDVVDLPDDLPDPDPELLDVVPPLVVLALSSSPPLVTATM
705	IPTAAPMTTRIPTTMRTLRAATIVF*
>scaffold_5_F17	MQLLRNLVFLALFAGVVAAVQEEQPQQKTHLRGLLDKIVSSDPTPAPVRVLVASWEDGKVGEARKLRPIAEK
18	IVGGETVVMFNDRRLSTADDLESAVVNFDVRSNPNCVVSVCGELTKWYLGSKEQCFDLRIAPKSADQIENGL
	IRGVSAAYSGLRPVAKNLYLKFDETNFIDGSRCEYEILSGSKTVREEEIASGSQAAMA*
>scaffold_25_F1	MRLSFLLAAAMATVYCATCNATVDSDQNKVSMVQSRLNGQADGTRFLRTHHENEQESDREERDLTDVFET
308	KKAAVKKLAKEVMADNRRAKDVFQLWKEKGYTLDELNTFLKSAKYQHVYNQYMIFRGYV*
>scaffold_53_F9	MRFFLVAALAAFAFASSCEAAVAEAPNAVQIDDNINAGFSRSLRSTDSEDRAIAQLLSEDRAVASVKVKYQSL
1	YKAKITPNQAKVILGISDDMVELTKTTRSLQRFYTGYLSYYTLMEKRKKRKKELENQVKW*
>scaffold_11_R9	MILSTWLRLSAAFLLAAIACCTTQLGTPRPSMWINVTCSNTFGRSVSRKIMPDTRMYVRTGPSGCCDAGQH
69	KWLIDRILRAHTRRPVSRSTKESIYSRKHTSRPSESFDGGFHSDQRMAAPRNGLSVDGSPVRTAVASAPSPAP

	SAHIRSGPDTRLLPLPACRTGRMSSTWPWTWSCCRRGAPH*
>scaffold_98_R3	MRLSFLVLAATFALLSCDTVASPMTKDECNPSPNQVLSSETEGRRLRVHKSSIDDVEERGFNPEKFNRLMNE
65	QSYRRKRFPNWVSKKYTDRDVYNLLRVDSNPNYKRIFNYYQTYLENFAPRLISS*
>scaffold_66_F2	MRWTSILLVAAAALTGVLDASTSNTVANSGVMESAFSPVIHDHRSLRLVDDDTVDDDEDESVDEVNEERT
74	WFSNAKQALADKLALTAVASNFAGKSTDEMGEVLKKLSRDQINTIFDKGEDSIRQILPGFKSGMDSKKFDDLI
	KALPQEQQGVLLSAYGKYLFNNGLL*
>scaffold 5 F17	MTLLAFFLEFGFCVLAMAAKPTFFFSCCTVGPSALLLKPKRELVCTRSNATSSSPSSSELLSSSESSLPSSPSSRRA
02	LRFCSSCSSRSVISSRDVTNLRAVPFSRFLRPLPPAAPGFPPGLPLSLIIVF*
>scaffold 10 F1	MRVSVILLVVAAFAVALNPTAAASDANTVVAVPNVHESIATGRFLRARPVDDYPVKDERDEDDSDDEKDEE
990	RMFSFFQEKATALSAFKKLVSQSGDDLVEAVSGLSKGEFQALFNQGKAHMAKMVPGFYPGMSLGEFGTVV
	RAAGLSDDMENALMVGYGKYLAHLMD*
>scaffold 26 F3	MERVGGLSCIAVAGATVALVEALLSGLEQVRVGHHVSREVQQARKAIAYVLGGARRVEAQHSSASVRSHGL
52	RPRLKAEEGLANCAARSLRLARSLLLRRGRRNCEPRRERGAAERRHCKAAGWDQ*
>scaffold 96 F2	MRLSHFFVVVAAAFLATGAIADSEPNHRNLRKHHDTLVDEERGIPKILDQNIPIERLNSLLKKMDQRKKAELA
12	KLIKLYESSPKFGRTKPSNS*
>scaffold 92 F1	MRFSQVLVVAAVSLLFASETAAVATSNQAKISKVSQSSPSQRLLRSNKYPIKEEEDESEDSVDWEERGFATPD
4	EEELEERSPLSDATVGKLNKIAKGWGTTYGKVAMGQSHISEAKAKALLALRDAYISGDKSAKAAARMAILNA
	NHR*
>scaffold 853 R	MORRMSTPFFLMITTIGEAOGEVEWVMIPADNSPLIKSDNFSRNCGATRRTRWATGVALPTSMSOTAPRA
1	AGGRAFPFPIRRCLRFQMICLNRSICSGVLPSASSIWISSILTSSVVALTSGSMAI*
>scaffold 42 R6	MRFCFITLFVATTILASSGNISAATVLNRSVTTSIDAVQPIEAAQTNSNRSLRRRNNYKDEEVAEEERSGIESLA
7	AQLNKKLLPTIRQVAHLDLGRAALTLQQMSLSFDERQAIQALLQLSKKDRKAVLMLIK*
>scaffold 7 R19	MRLLLWTLLVILVTILSSCDAASVNENKSLORKLYTKVASHALAADDGFEHDKRALRGASNGVTEARAATVST
13	KEGSRLMAFERSIKDKYLAWELKILVPGFEKMAKKGTTYTOVREDERTRLNWSGLWGTPSGEKRYAKLYRT
	WLEKNHYSOLAV*
>scaffold 8 R16	MRVLGAIFSALFLAAGIDVGSSLPNSGVAFPVNPHSLAVETVESKRLLRSYEADSEDEERAGGOLGLVDKFAA
99	KVMQKLYKNPSDVFKRLKLKDVNLENNKVFEGWLVYVNKFRQVKGVENFPDQALFNVIHQSQYYQRDLVP
	LFQSLTHVQGMKDLARTMQFKLFEAATPATRTLMNKAWLEGFDTSDDVFHILKLQDGVFDHSDKLIQWLK
	FSDMYKKLPTSQSTSWLDELNLVLKTKKPNQQETKFGLLFQALKEEKGMETIAGKMESQLFERWMKMDSM
	TPDKVGGMLGGSATTNWKRIFEKLEFTDDRYIFLKAYTEAYAANRGANVLKSVEKLFAENKPVAALERAIKA*
>scaffold 87 R4	MRLSQVLVIAAATFLFASDTVVIATSNQANISKIEQSIPSQRLLRSNKYAVKEEEDESEDSVDFEERGFATPDEE
30	DLEERSPLSAAIVEKLDDIASRWGTTWGRVAMGHSSISDDKIKALIAMRDAFISGKKSDRDVARAMILMANK
	S*
>scaffold 15 R1	MRLTSIAFVAAASIFVCFQEASAVSDSKGPESRVQTANSDNRFLRIDQDEEERAGPQPSKSLLEKFTSKVTATK
164	LKRSKAVADLTKLDDVVYLKATPGYNVPLFQRIEKMGFNPDGMLLKMRERGKIDQTLLKHYTNYWKGKYPT
	WTSNPSV*
>scaffold 17 F1	MPVFDALLTLTILRAIGCDRVCANQKRRMADIVKRMEDGAFRTGKAWRILRKLNKSEMKNDAERWSTFLEL
758	SNVLDPRAFNSRRGRALMNEAEQVISVQQLELAAVRGPIVDVKPEGENVADELWDIAEQIVEFIERKAAPVA
	EPQDTTNDPTFSFDVVLVTDLEDYLKVEARTAPSGVNGMSDRETHIIDEGIELSVTEQLSRMCSSLKEEEYVIS
	SDILEKGFAELSAIERLDTVSIQVSFLLGMQMRLKSCIIDHCESAKSEFERRIKRVFNQLAGKGQDMDHAASLI
	VLSAFCPGQVIRECIRGARTGVLHHDLSLKVLQSSPLLLEWREPSEDGSGTLFELELQMTVLDISRNQSSFDRE
	SHNVVSFLLSLVGIDNSTSSIQSSSLMTVSKLITVCINPVWCCPDQSVEMQLNLLTLIQQLFHHFVATSADIDT
	DTLQISFNLAFRALSSANSNDTKVGALIREKVLLLLKSIMELMPDPVSTIRLEQFDSPLNAPLWTLISLFSDDLG
	NEELCKGLEDVTAVEAYMQTSSVDEADPLPLPIVISAIQALLWGLLWDSILSENVSVESTKAETWRLLDIIASFE
	FCGSESPDETIKGSSLIESAVAEMMLECGNVLFRALLCNIIPFLLEYELTENFQEEKLLIPQWAIDKVSENPEIKE
	QIPCRLVSSHVIMRYVAKCWCLSGVANRQLNAQSDMLVLEALTHVVTAHDQAITASKESLSGTLFCIQWLCF
	LVSAARELHLDTLSTWSTVRTQLELSFLRLLNQLEHLKAISEAEAKFSQVFVAAWLGYLPDDQFVQVFKYISTK
	RSN*
>scaffold 22 R9	MILTTNLATWIFLTWAATSSMNLSKCQCTCCLNKCENPRPLRGHLLSSWVSGKQSGWPYQANLLHVHLLEH
74	PRQIRTVGSRK*
>scaffold 65 F3	MRIPCSLLVVIIALLSTTTNAISTEADAGRRSLRSMKAKVQDNQVEEERGGTGTSSLKAFVHDFDFNLLDNIFL
81	PADFKRMTKEPEFLRHMMASWQMGFMSVDDIVLYMTRLNMSEKAINQFKLAYTAYLEYLKAVAKAEKAA

	KLAKASLN*
>scaffold_25_F1	
318	
>scallold_184_K	
0	ADI TVDOKVI I EVVSMEWIKIKVDSWIVSEVDRME*
Scaffold 1 E40	
0	
>scaffold 6 R20	MRVLLLVALAVFASSSVNTGVAGIQTKRSLRQYDFKSLTVADSKEEERNFVDKVDDIARVTDDFVTKVKIPTN
44	MNAAVEKVTTMADDIAAVAKTVAKTYPEGLSKGTLAQIKEVEQLRLKDIATYTKKTGDGMRRKISPFPGMKI
	APKKYLESHVGRNMQLYGDDGSRLMSSAVVSRSAKDGGGDVLLISSSNPQKNDWLLPKGGWDKGEDIQS
	AALREVVEEGGVCFFILYM*
>scaffold_60_R8	MFSLPTFVMFLVVPILVMKAEDWSTEGISHSLPNARSLRQERVVDNSDVDSKAQRISGHVATVLGHSAPMD
42	EVGQVVGMLLGV*
>scaffold 8 F32	MPQTPRCLLLKVLFADVVHCAALYDLLGHSNGSDLKYLMILGHPADLQLSGIQFVRLLRGQSPVALPVAHQL
65	ST*
>scaffold_10_R1	MRLSVILLVVAAFAVALNPTAAASDANTVVAAPNVHESIATGRFLRARPVDDYPVKDERDEDDSDDEKDEER
425	MFSFFQEKATALSAFKKLVSQSGDDLVEAVSGLSKGEFQALFNQGKAHMAKMVPGFYPGMSLGEFGTVVR
	AAGLSDDMENALMVGYGKYLAHLMD*
>scaffold_81_F2	MRFHALVLLSTALLATTDAFKLTADVTAPQSRLLRKSEPKTNEERVFEGLTSTKLQKYNLLWSKLQLGDDLSAV
33	LKSPDVAKIAKFNLNRAPGSQVSMIQRITAKYGDDVVARTLVSIERDASDNPILLTMVWQLREDQIANWLKN
	SETVPGVVSKLKLGTDESIFRSRALDVLEDFIKKYNTARNGDESLLKTLTTIYGGESELVKMIAKTRVLGRPPYSN
	HPASIEKANNIESQLIQKWKSQNLPDFRVMNKLNFDDDVSMALSAGKVGVLLKYSDSKTSAFRRLSAKYGEA
	EVAIAFAKAEGFLPEATALYRMQMNGWLSKGDTAGRVFSILKMKDTRDFVYKLDALETYVKFLKTKNPDDV
	TDVFKVLKKGFGAGENKLALAIVRPPEFEITGYHKSLFQDWVARDLDPLSVAVKVFKMSEADVAAAKYSDEL
	KPIIKQYTKFFKETVDLPEMPAVRVGRS*
>scaffold_6_R31	MRLFIKNLVALAVVLLATSTEATKTIQTSGKVDVVQSSQILTGENKRLLRSEHDEGELLGDDEEGLLEDDEEER
91	KYGANLFSAEKMKKMLGDTDTEKKEEKSCQQ*
<pre>>scaffold_67_R4</pre>	MWLLVVTLVFSGSIFTYADNSGSDSDNSIDDDYVESTYYYSSNAGLFVGMLVAANTIMHLRCLRTAQLLVAE
06	LVNAGVDRLQPPVVMLATTNPTSAV*
<pre>>scaffold_636_R</pre>	MQRRMSTPFFLMITTIGDAQGEVEWVMIPADNSPLIKSDSFSRNCGAARRTRWATGVALPTSMSQTAPRA
1	AGGRAFPFPIRRCLRFQMICLNRSICSGVLPSASSIWMSSILTSSVVALTSGSMAI*
<pre>>scaffold_12_F1</pre>	MGVWDWILCFTVCEEVGAACPCLTPSIVGRVLLHGKTKKIVSKPPTSNSQTMFCSFILDLIWTTYTTMSKPIQ
261	SDSDLDALRKLTRQLRVAKLVTDRDLKQTDRKNATQAVMRKWLPLSTTVLKMVTRVLPSPIAAQVKRAERL
	CTISSEQLEQSPQHAQVFRSLQSCQTSEDAPLVIYICKVISVEANVLSDYHQSGLAATDEVYVGVGRVYSGVLR
	EGQPVYVMDPKFQGVSGDMDVDTIDPSTVKHVARIDSGLIKTYMMMGRDLHKLDRVPAGNIVGIVGLQE
	HVLKTATLSSTLACPSLTKMPYQAKPIVRVAVEPEDPRNFGALEAGLQRLYRSDPTVEVHVQETGEHVIVALG
	ELHLERCIKDLKERFAKVAVQVSEPLVGFRESIVDGTISSFQENIVFKELLNPDVTKDDSSEKEDINASAEVQDT
	KVALGTTPDGTLTLKLRALPLPLETAKLLEESASLLKRIAVSKKANDDMKDKNTVVELEAAASEDVSVFKKKLEK
	SLQSSESSFLKALPLDQIWSCGPRRVGPNMLINSIPTYRATGCLFPSDSVAETHDSEKDEKIRKLENSIVTGFQ
	MASSAGPLCDEPVWGVAFIIEDVVFHDEKSEEDKSEEEKAEMSKYGPLSGQVISIMRTTCLMSFVKQPVRLV
	EAVYECTVQCQAEQLGKLYSVISKRRGDIYSEELSDGTALFTVKAHLPVVESFGFATDLLIQTSGAASNPQLIFS
	HWSIIEMDPFFQPQTEEEREDYGERVYEHNYVRRYIEAVRKRKGLSRDEKVVVHAEKQRTLKR*
>scaffold_81_F9	MRFPFLLLLLVTFLLVNGKADSQSTELRLLRDGNSPREEERALPPPPKVSLFADWLKATGLKVSDKVRARYWL
0	WRKQSAEDVFRLLKLDGGLEKLLGSRKFNTWTSFVNIYNKKNPNEKVTMDGILSKTYGDLELAKALEVAMGS
	MVKSERKMG1TLSLQQREGWQAAGKTADDVFILLKLDKVGADLFTTPQLNSWYKYVSMQADSKSLMASV
>scattold_6_R31	MPLLILIFIASILATTIYVNTKLLNILPGNSLGVSTYNHSHLDWKMDTAAFETSVGTSIHPTSSFYRRGKTFDRSA
37	SRERAVMVCLHDAMLNMGLSLLRELRCLGNHELIQVYHCGNELSDRSVELLFSLDNRVELVDVCSDLSSRGII

Lamour_MPMI_SuppFigs/Tables Page 53

	SKEMATKFRSWWIKPLAMYHTDVRHVMLLDVDDVILKDPAVVRTLDGYKNTGTTFFYDRVISNKRFLTGND
	TGEMYVHKLLRTFNYTRFNVSEGFNPSQHMLNTFAFNGKSIHEMDSSMVLIDKQRAGKVVMDILLWFITEE
	RFRFTYSWGDKETFWLAFEMARVPYFFSPWGVSVVDSMPNEDLKNHPDTLCGSILQFMPVKGMNDTAEV
	LYMNGKALIDPYPQGIGFIRKAKQNNLFNTIPTQMTPPQKRRQINTKAYPGKKFNTECLVGMGAVPLPTHFS
	PNLLRRRVHFLGLAMGVLGSLDHCETYT*
>scaffold 2 R11	MPVCYFMFLCVSYFASSSCQIKQHVEAFIGDLADAFVPVEGGVLVQRRRNLLTQGLVARQVHTHELEVPAH
14	VARYLRHEEVSRLDADLDAREGAMYDVRREHETELHELREQTCREPHEEQLQLDVALWPLQ*
>scaffold_5_F94	MRTTFLWLLLALVLCVCAAEEPKTPEPTSSANARDNDPVVQEIRGLRNSGMKLNDAKDFKGAIEKLREAITLL
7	HNRVFGEGRHAITDPSEISQDAALYAQILNDYGSVLIRAKQYDEAIEVLEDSVTMVEKIYGDSHPSLGLSLRSL
	ADAYMAKEEYKLAIKKYKTLRKHVKKGLEETHEAYIEASLRIAEGYKKLGNKKKNLKVLRDAVKAQGTEINGLT
	TGIAELYMELSTAHTAVGEIDDALRAAETASAIFLQRDGEETLSYAFSLNALAGVKMRQKKVDEAVKLLEHAH
	KIAVKIYGENDPITQASAKTLKEVKEYRLDMHAQKDEL*
<pre>>scaffold_48_R1</pre>	MRALCFIFSVLLLLLEIDASSVFNSGAAVTHPHLELTQVTTPTRLLRTSHEEENDERAGASVLDGIVAKAMQLI
407	NKNPEDVFKKLKLANTNLQNNAVFEQWLQYVYKFRAAKGEDKFNNHRLFNLLRKSQNHPDDLVPLFQSLT
	HVQGMEDLARTMQLKLFESGYQSTRNLMNKAWLQGLDTPDDVFHILLLEKNALESQQRLPQWLKFAEMY
	KTQNKISSWENELSLLLKTPHEKETQFGLLFQSLKKTEGMETIAAKMEAQLFTRWIKTDTMTPDKVGVVLAS
	PTNTNWKRIFEFLPVTEPRHVLLESYTVAYAASRGGKVLKSVEKLFANNQPVAALERAIKV*
<pre>>scaffold_59_F7</pre>	MRLTYIMIVATAAILACCDGASAVSDSKRLTVPLDRVQTGNNANRFLRVHHDEEEEEEEERTGPQFIKSLSEK
11	FAKAAETGNFAKTVEKLTRSKSLNGISKVDDVVVPGRAGKTLTEVLKIDDVAYLKAAGKWPKNLDDIQEAKLV
	REISTLDDQATLKLITKENAAEFKKIEDMGFNPDGMLKRMKQLSKNDKDYFDLVLLKHYTQYYMAKHPTWA
	SSL*
<pre>>scaffold_29_F1</pre>	MRVFSILLLVAAATLVASASAESESKQRLDVSPAAPQWRTIAENEVPTKRNLRKKEIEEERAITAISLFDDVVKA
645	KGLQALPYPELANLDSKLRTQYLKLLVNNLDRKQIVEITGQVPHYILTHGDKKARRIVQYNKWIWRKEPVDPD
	WVLKNYPAFFKGYEEFFNNRFTRGYKYA*
<pre>>scaffold_2_R17</pre>	MRFSHFVFIAAAILLSSENAIADTPVEGQALMTETDAETPVRALSSNNDKRFLRSYKEEEAYLTEDKYDEEKKR
01	EES*
<pre>>scaffold_29_R9</pre>	MRLLYIAVVAILASTNVYPAAADAEVSSFTFTERPHWDRSLTAATEGDNDGKRFLRKRKKTTTNETIRERSKQ
98	GSENTVSYCTACLHYP*
<pre>>scaffold_9_F34</pre>	MRFSHVLLLATAALISSCNATAIVPGEGQVTTSADVAVPVRALETRNGKRSLRYYDAEEEDETDKYDQKNGK
50	YDDDEEERVMTAAQIAKWTAKAGEWVEQGKTPAYIKDKLTAFNGVMSDKNKEKYRLFLATWGRAHPNEL
	GRRYLRPEAESSPRSFRKLIVFISNHHLLDVLFCVTR*
<pre>>scaffold_6_F30</pre>	
32	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV*
32 >scaffold_32_F2	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV*
32 >scaffold_32_F2 41	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW
32 >scaffold_32_F2 41	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL
32 >scaffold_32_F2 41	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI
32 >scaffold_32_F2 41	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI
32 >scaffold_32_F2 41	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE
32 >scaffold_32_F2 41	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA
32 >scaffold_32_F2 41	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNYKFVVGRSDSKPWGERLLTDVEAVTEKM
32 >scaffold_32_F2 41	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNYKFVVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFGFDFRVMRFESGFVCLDVDELVKKCANTSRRLFIFDYGGTLSSTANILDEEGARFSHRPG
32 >scaffold_32_F2 41	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNYKFVVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFGFDFRVMRFESGFVCLDVDELVKKCANTSRRLFIFDYGGTLSSTANILDEEGARFSHRPG CVEQSAENADSSDRSGGDCTGRSAARYVDGKVRKPISDETRASLRTLCVDPCNIVFVTSNTQRGALEDQFDSI
32 >scaffold_32_F2 41	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNYKFVVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFGFDFRVMRFESGFVCLDVDELVKKCANTSRRLFIFDYGGTLSSTANILDEEGARFSHRPG CVEQSAENADSSDRSGGDCTGRSAARYVDGKVRKPISDETRASLRTLCVDPCNIVFVTSNTQRGALEDQFDSI PNLNLIAENGLFIRMGGQTRWECVCEEDKRSFGWKDDVKRVMEAYAARTNGSFLVENAASLLYDYRNSDQ
32 >scaffold_32_F2 41	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNYKFVVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFGFDFRVMRFESGFVCLDVDELVKKCANTSRRLFIFDYGGTLSSTANILDEEGARFSHRPG CVEQSAENADSSDRSGGDCTGRSAARYVDGKVRKPISDETRASLRTLCVDPCNIVFVTSNTQRGALEDQFDSI PNLNLIAENGLFIRMGGQTRWECVCEEDKRSFGWKDDVKRVMEAYAARTNGSFLVENAASLLYDYRNSDQ EYGEIQSLELCAQLRKIVEVCIKVILSSR*
32 >scaffold_32_F2 41 >scaffold_399_R	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNYKFVVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFGFDFRVMRFESGFVCLDVDELVKKCANTSRRLFIFDYGGTLSSTANILDEEGARFSHRPG CVEQSAENADSSDRSGGDCTGRSAARYVDGKVRKPISDETRASLRTLCVDPCNIVFVTSNTQRGALEDQFDSI PNLNLIAENGLFIRMGGQTRWECVCEEDKRSFGWKDDVKRVMEAYAARTNGSFLVENAASLLYDYRNSDQ EYGEIQSLELCAQLRKIVEVCIKVILSSR*
32 >scaffold_32_F2 41 >scaffold_399_R 2	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNYKFVVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFGFDFRVMRFESGFVCLDVDELVKKCANTSRRLFIFDYGGTLSSTANILDEEGARFSHRPG CVEQSAENADSSDRSGGDCTGRSAARYVDGKVRKPISDETRASLRTLCVDPCNIVFVTSNTQRGALEDQFDSI PNLNLIAENGLFIRMGGQTRWECVCEEDKRSFGWKDDVKRVMEAYAARTNGSFLVENAASLLYDYRNSDQ EYGEIQSLELCAQLRKIVEVCIKVILSSR* MRLTCILLVAAASLVGVLDASAATTGNTVVANAAMVISPLAPESQGRRSLRLVYDDEDDSADEKDDEEEDSA DKVDEERGWLSDKMALTSLASKFVGKSTDEMGEVIKKLTPAQIDTIFDKGEDSIQKMLPGFYSGMDFQKFD
32 >scaffold_32_F2 41 >scaffold_399_R 2	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNYKFVVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFGFDFRVMRFESGFVCLDVDELVKKCANTSRRLFIFDYGGTLSSTANILDEEGARFSHRPG CVEQSAENADSSDRSGGDCTGRSAARYVDGKVRKPISDETRASLRTLCVDPCNIVFVTSNTQRGALEDQFDSI PNLNLIAENGLFIRMGGQTRWECVCEEDKRSFGWKDDVKRVMEAYAARTNGSFLVENAASLLYDYRNSDQ EYGEIQSLELCAQLRKIVEVCIKVILSSR* MRLTCILLVAAASLVGVLDASAATTGNTVVANAAMVISPLAPESQGRRSLRLVYDDEDDSADEKDDEEEDSA DKVDEERGWLSDKMALTSLASKFVGKSTDEMGEVIKKLTPAQIDTIFDKGEDSIQKMLPGFYSGMDFQKFD DLIRALPQEQQAVMLSAYTKYLHQNGRFS*
32 >scaffold_32_F2 41 >scaffold_399_R 2 >scaffold_7_F17	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNYKFVVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFGFDFRVMRFESGFVCLDVDELVKKCANTSRRLFIFDYGGTLSSTANILDEEGARFSHRPG CVEQSAENADSSDRSGGDCTGRSAARYVDGKVRKPISDETRASLRTLCVDPCNIVFVTSNTQRGALEDQFDSI PNLNLIAENGLFIRMGGQTRWECVCEEDKRSFGWKDDVKRVMEAYAARTNGSFLVENAASLLYDYRNSDQ EYGEIQSLELCAQLRKIVEVCIKVILSSR* MRLTCILLVAAASLVGVLDASAATTGNTVVANAAMVISPLAPESQGRRSLRLVYDDEDDSADEKDDEEEDSA DKVDEERGWLSDKMALTSLASKFVGKSTDEMGEVIKKLTPAQIDTIFDKGEDSIQKMLPGFYSGMDFQKFD DLIRALPQEQQAVMLSAYTKYLHQNGRFS* MRLVNVWSMLGLVLVLVMVTEAFETTKGAKNGVVTPDARRLLGFAADQEERRFGGPATGTDHSNSHAW
32 >scaffold_32_F2 41 >scaffold_399_R 2 >scaffold_7_F17 75	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNYKFVVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFGFDFRVMRFESGFVCLDVDELVKKCANTSRRLFIFDYGGTLSSTANILDEEGARFSHRPG CVEQSAENADSSDRSGGDCTGRSAARYVDGKVRKPISDETRASLRTLCVDPCNIVFVTSNTQRGALEDQFDSI PNLNLIAENGLFIRMGGQTRWECVCEEDKRSFGWKDDVKRVMEAYAARTNGSFLVENAASLLYDYRNSDQ EYGEIQSLELCAQLRKIVEVCIKVILSSR* MRLTCILLVAAASLVGVLDASAATTGNTVVANAAMVISPLAPESQGRRSLRLVYDDEDDSADEKDDEEEDSA DKVDEERGWLSDKMALTSLASKFVGKSTDEMGEVIKKLTPAQIDTIFDKGEDSIQKMLPGFYSGMDFQKFD DLIRALPQEQQAVMLSAYTKYLHQNGRFS* MRLVNVWSMLGLVLVLVMVTEAFETTKGAKNGVVTPDARRLLGFAADQEERRFGGPATGTDHSNSHAW ENFKAWFKETFFFWRKWEQKRRLRS*
32 >scaffold_32_F2 41 >scaffold_399_R 2 >scaffold_7_F17 75 >scaffold_18_F2	MSAFLVAFRCCHVVFAFHWELRSVEFALALGLNRARGSRWLRLVAHLLSLYEAVLDGPSGASTSVAFMPIV* MSRAGSPVALVPPKTVLIVVNVLPLVFERSETSASGWSVSWATGATAMLYRTLVRDADQYSPLFIGCPEVFV AKSEEAALEKQLRALRCVPVFLDPPVAHRYFQGFCKGVLWPVFHNVVDVYNSAELQLDCEENEGKRAKQW FPGQKEDSSGVKSAWCDPVSWNPAAQDMCWSDYCSVNRTVARRVVESYHDGDLVWLHDFHFLMLPSYL LRRLRTALVTMYLHVPFPSSEIFRCLAMRTEILRAMLCADHIGFLTFEHARHFLTSCKRLLGLDYKTCPNGMLVI EYNGRKIHISCSHVEPDVTHLHEMLDKNVAEVNGAALEFAAHVKNAIMAGRDSEQRRIVIGSVDRLEGLTAI PLKLRAFDRFLATHPEKCSSVVLVQIGISLDSRPNDYHRTRDNVLKFTEEINRRYAPPGGVVVYFEEKRKTTCAE RVILWRMCDIYLDTCVRGGLSLLPFEYMIAQHRNIQSKAKNSAGTGWSSPRGTIDVGGLDRSFGIMIVSEFSA YSRILSGSLAVNPWKTDDMVGALAKACGMSYYEKHNRFYMNYKFVVGRSDSKPWGERLLTDVEAVTEKM EDASTGEVVQVGFGFDFRVMRFESGFVCLDVDELVKKCANTSRRLFIFDYGGTLSSTANILDEEGARFSHRPG CVEQSAENADSSDRSGGDCTGRSAARYVDGKVRKPISDETRASLRTLCVDPCNIVFVTSNTQRGALEDQFDSI PNLNLIAENGLFIRMGGQTRWECVCEEDKRSFGWKDDVKRVMEAYAARTNGSFLVENAASLLYDYRNSDQ EYGEIQSLELCAQLRKIVEVCIKVILSSR* MRLTCILLVAAASLVGVLDASAATTGNTVVANAAMVISPLAPESQGRRSLRLVYDDEDDSADEKDDEEEDSA DKVDEERGWLSDKMALTSLASKFVGKSTDEMGEVIKKLTPAQIDTIFDKGEDSIQKMLPGFYSGMDFQKFD DLIRALPQEQQAVMLSAYTKYLHQNGRFS* MRLVNVWSMLGLVLVLVMVTEAFETTKGAKNGVVTPDARRLLGFAADQEERRFGGPATGTDHSNSHAW ENFKAWFKETFFFWRKWEQKRRLRS* MQRRMSTPFFLMITTIGDAQGEVEWVMIPADNSPLIKSDNFSRNCRATRRTRWATGVALPTSMSQTAPRA

>scaffold 50 F2	MRFLVFLALAIVALVADGEAKDSRLLRARSDEERGGGITVPVADKVAKLFQFSKLTDEQLQQWLSAGKTAES
79	VFYRMGLDNTFVTKVFKSPQFIRWLQYADDLSASGKGTSAISILSTKYGDEQLYKMIGWAKKESSTEALGLRL
	QSEQLDHWVKVGKDPDEVFKLYDLNYAGSRILSTSQFSAWTKYVDDLNAKNEGAFVSIIPTLRKHFSDDELF
	HIVLAAKRSGETEAMGMKLEDAFVQFWIHRKETPGNVLVGLGLKKSMETLLESPLLSILTKYTEVYNVNYAAK
	KTTVIETLTRTFGDDKVARMLLAGRTESTTTKIAKQFQTDQLEMWLSSGQSVDDVYKLLKLPYRDLVADIGTN
	KLFGTWLTFMNAVSLKNPEKMSAIFTRLEPTFGNRPMMQILEAANKLPSMEKAATKLQLEKAQRIFSTGESP
	YQLFKLVALDDVGESVLSSPLFKKWMLYVEDFNKKNPSDEVSWFVALRAGYQDDKLGPIIDKAMKDPKTMK
	LAKLVEKERMKEWLEKWKRPPSMAFRELHLNKAGEKVFSAPKFELWVKYLDDWNQAYPSKKTTMIDAFVD
	NYHMSTLVPMLAAAEKVPITKKLASQLKDALVDKWIAEKESLAYVKSWLKVVPSSDDMLERFTKKLNSA*
<pre>>scaffold_7_R19</pre>	MSVLCLALMLAMVTGAFETSRDVKNGVVTPSSRRLLGFTADQEERRFGGPATGTDHSNSHTWENFKAWF
03	KKTFFFWRKREQTRRLRN*
<pre>>scaffold_24_F3</pre>	MRLSFLLLVAAVALLSSGTVVSAMTNDEPISTPNQVLSSETEGRKLRMHNIASDDTEERGFNAQKFEQLMD
57	DGTYRSKRFANWVTKRYTDTDIYNKLQISSNPKYKRILNKYQTYIEHFAPGLISP*
<pre>>scaffold_17_F3</pre>	MAANSCVAVLVLVCGLPGAGKTTLVKQLVATACSISSRLHERISFDDLYEQHVTAEGKPGEFDPEKWKMCQ
59	QDMLKRVSNRLKEQNDPVHRNECNQLVLLVDDNFQYRSLRKRFFHLTAKRRRPSGFTLCTTTGMPLMTRSE
	LWFWCALCGCSFRYLSGTECWSKQTRTSTQ*
>scaffold_8_R31	MRLSFLLPVAMAAIYCATCNATADSDQNKMSTVHSLDARLNDEAGGRRFLRVNQEEEDVDAEERGYFEKA
93	AVKKMAKAIMADPSKADEVYNTWAEKGYTLSKMSEYLKAKKYDQVYNGYALHLDI*
<pre>>scaffold_83_F3</pre>	MHPPSLVLTLVLSPAALKSSTLHVAALELQDTVEPSSHLASEPSFSPRCRRPLRVASSTCAKAGADPLLAPNGH
73	LK*
<pre>>scaffold_25_F1</pre>	MRLSFLLPVAMVAIYCATCNANVDSNQNKASMLQARLNDEAGGTRLLRVHHESDTEERGFLEKAAVKKMA
083	KAIMADPNKADEVYKKWADKGYTLTQMSNFLKSKTAGKYDRVYNGYVIHLDY*
<pre>>scaffold_22_F1</pre>	MRLTFLLFAAMLAAFSSTSDAASVDQTTGVRSLRRYQAEDEERGVTSVSTSIDDVLNNLIKQKSSLSKLISQKK
823	LDNMLTKNFIKLLRSNKSYTDEVFTKATLNKMLTSEKFAEKKFVEWYALGLTDKLILQRLNGVGEHFGTLHSQ
	YVTFINRIHGVA*
<pre>>scaffold_157_F</pre>	${\sf MRVASIALLAVVTALASVTDSSAATTGTVLAKVVSNEAAPSVENEHATRFLRKHKDHHADTEREERNGISLLQ}$
14	GLKSTFEKVADLPFDRAWHHLQMLNLSWDKREALLKLHRLSAKDREAVLKLIT*
<pre>>scaffold_7_F23</pre>	MISRSFTLITAGVMASTTLIHASPLQYDPYTPVNISTPLTSSHPAYGAQTEGCIEPIVPEDPNQAKAESMIIQAD
10	IYRKLRSMEDTTNSDIQDLETYFGTKMEVNFQTLKQQYSSGHAPATPWASSYWPTFQDSINYVWKTGEPSA
	SEKYATAYGLNVTEFKDKISERSGVDSRRRSTRCTADSDCKDGSVCGKRDGISSGYCIPGWFGICHAWAPAAI
	LEPEPQCDVTKNNVTFHVMDIKGLVTSIYDGAEIKTVFTGARFSGLDSPANKDQYGRFTDAARRDLGPGYFH
	IAITNVMAKQNRSFVVDVTAGSEVWNQPVRSFNVQSMDLVDTRVASMQYFGVPSYPFNDKMVRLAYVKT
	TFSWVSESYKDGPLVSSGRIDRYTESKDYEYLLELDADYNIIGGEWVGQSKDEHPDFLWLPTAKPSASAVTST
	GLKYADVQELLNLSQSC*
<pre>>scaffold_39_R8</pre>	MPVLPRLSLAGSSSCSGAQVLSSLLALLDASSAQYASLPSFDELFYPLYLLLHALVKQLEDTKVSEVNAVISKLH
34	NRLETCWNARRPLRLQTFAPTILPTFAPQFDENYTVRKDKTAPKDTAQLKQLQRQVKRARKGAARELRRDA
	EFIHREKQKEEEARLSAKEEKQKEIRRWLEEQNATFNQQVRKGGHMLKGGGSARGPAPRARTPRK*
<pre>>scaffold_104_R</pre>	MRLTSILAAAVVVTLHTSATAFPAVKDTAAIENGAVADIVDSSVTKGGRMLRVNYNNDDDDLDDKKKKTRN
20	SEDPLDNYDEEERVGIVEALKKLNPVTAAKNSAKKLQSTLRRSRK*
<pre>>scaffold_61_R5</pre>	MKVTKVVVALAALCVALWTSPTDSEDISNILEVSASRHLRQTSAEFAAKPQETGKKRDSTNPLQRRDQALVS
21	AHRVYDPVSGLACSLVGECVACPQSERDESFCRETGYRQELDCPRPNDPKDEALLTKPEDERETRFKACSPA
	DSARPGVAVVKFELLMAAVLAASVVLLRRERRNHMSSFDLRKDPRQRTGLLGGSNSDKSSD*
<pre>>scaffold_90_R9</pre>	MANLYKLFVTLLVLVSWGSKVSVCCLTSRFLRTQNLKTRYLAKSTSCFQCGSGHHLHPLRHYAVSVSQLASLN
4	LRRLACC*
>scaffold_25_R3	MYRVLLLTVFALLCGFSNAANSGGRLLRVVDSAEERGGFTSYLMDSFTKWRINSKINSWVNKQKTDEYVLAK
32	
	DYPITRWRELFGGGSPEQLKLKRQLLFLAKRNAIDIRIMLG*
>scaffold_18_F2	MRPNLVVLAAVIALVSRCTAVSADSSTNLNQALDTLQNHAGDGNRFLRSANIVDDRVDDIDSDDAIDNDDE
130	ERGEKLGLKNLPNGMRAERPQTTSTKGSHSSLTYAKRTNMAKSDGSITTSTIASGPITWPS*
>scattold_74_F6	
12	
	TIVIGLEUKYKTIVIVIIPTVALSETATRIGVPSVLAARAVPTOTHOPLEPRRAKLIREEROGINVSRLEVAKSPRIRL

084	QGHLRAVVRYRTRRMIL*
>scaffold_11_F1	MRIGYVTLLATTAILASFGNVSGDTDLDHTQVVKVASLDAVLPTNTNRLLRGRKNNEEMEERNGFEALAAKL
319	DKSVLPAILKVANLDLGRAALTLQQLRIPFEQRLAIQALLRLSKKDRKAVLLLIK*
>scaffold_11_R2	MKWCRAASSASLALLACSLAGLQDQAADRWTVHTVQRTRSAEELVHEGTSRWPGDAGAVASVTCPSDLLL
801	ALQCREADGQRELRRDDQDLRRLTRMCMCCADHRRIKSGGLFALYMKASVRLTANCDRYELDAPLVSTLTC
	RSVDKDIHLLPDKEPL*
>scaffold 11 F2	MQRRMSTPFFLMITTIGDAQGEVEWVMIPADNSPLIKSDNFSRNCGATRRTRWATGVALPTSMSQTAPRA
657	AGGRAFPFPIRRCLRFQMICLNRSICSGVLPSASSIWMSSILTSSVVALTSGSMAI*
>scaffold_50_R1	MGLFYLVCVAILAFMAGDSTGDRVIVSDGFNQHRENAARASVVSTTRLLRTKSVIDEERVGGIPVSATDKLAK
378	FLKPSKVTDKQLQEWLRNGKTAESVFYRMNLNNPSTKYLFEDLQFTRWLKYADDLSASGKGASAISVLSAKY
	GDEILYLMIDRAMQEQSKALGIRLQADQLAHWVKVRKDPDEVFKLYDLNYAGRGILSNSQFNAWTKYVDD
	LSAKNEGAFVSIIPTLRKYYSDDNLIKIALAAKEVDETEAMGMKLEDAFVQFWIHRKETPDNVLVDLGLKKSTK
	TLLKNPLLNILTKYTEAYNVNYPSMRTTVIETLTRTFSDEVMAKTFLAGRTEYTTKKIAKQFQTDQLEMWLSSG
	QSVDDVYKLLSLPPRNSLVDFGNQKLFDTWLTFMNAVSIKNPDKTSAIFTTLAPTFNDRPMMQILEAAKKFP
	SMEKAATKLQLEKAQSIFSTGVSPYTAFRMVALDDVGESVLSSPLFKKWMLRPDRYHLMAHSHLCCRPAAQ
	HFLKRPRRRDVCGGNSCVLPRYVRGRGVAGVALLCRKPNELQLLLLSTTDC*
<pre>>scaffold_17_F2</pre>	MIVYTGLTVTTLVAFVATVCLELTEHTACVLSNDLGISSSTAQVLGLSLWSGVHLEQVEQSPFSRQLHGHVVA
587	LDQLEDGRYKGHQVFARLLRGALSRNLHNAQRATDHVLVLVRHTLHQYWD*
<pre>>scaffold_25_F1</pre>	MRLSFLLPAAMAAIYCATCNATSDQNKMSMVQSLDARLNGQADGTRFLRAHHESEEESDREERGFTDLFK
282	NEKAAVKKMAKAIMADPSKADEVYNTWAAKKYTLTQLSNFLKSKTAGKYDRVYNGYALHLDY*
<pre>>scaffold_68_R3</pre>	MRLNLAVLAAVIALVSRCTAVSAASSTNQINLSNLNQALNTMPNHAGDGNRLLRTAKIVDDSEDDIDSDDDI
1	DRDEEERGGKTWAEKFAKWHARGESADDVYQRFALEPVVRQAYKYGQIGRLDDNEYYRKWAAYSA
<pre>>scaffold_11_F1</pre>	MKLSMLVLALVCISQLGGSSANEATDIMRRQLRVGKAVASLFENQHQSTRELEENIMQDEDNKPNEVQAEP
227	TKFRMRRLRSDYVELIE*
<pre>>scaffold_117_F</pre>	MLRSVGGAHFDVHIVSLLFLLHGFSTHTLELLVANVVQSQGIETSECEVVDDQNGNTEPNVARALIDREALEL
62	SFALSISVSVTRELRQHIGT*
<pre>>scaffold_53_F9</pre>	MRLSVILLAVFALSSSSVSATRNHPGETATANTAMQGVDGMRTDTNQMRFLRTEADDDEERLAGKNMFN
79	AEKIEKALQDTSYAKTLFRRWKRYEVEHGAAFDKLIKFNIGKDDKVFGLYKSYVSWLEKHHPLGAETGGGPNL
	FSKAKLDKAMKDPKYENTMFGRWKRQGFESDAAYNKLLAFNLASDADVYKIYTKYVTWLNIHHPLAKTRKT
	_ AKAT*
>scattold_22_F2	MGGLRRPALLWIPHFLVADANCNAAVVEIDFKSGRILRDIDYAVISDEFARVAISRLSTFAVVSRPKVFRNDEE
097	GTGIVDRVRLLMLHPTDLSDSSRFGDVKLSFTLVRDVTNYCREAEVRLRQAREEIDGQLPCLTMHSFQLNIRE
>scattold_13_R9	
5	
Secoffold 72 D2	
	WISSKGHFFSTAFPILAACKFQSVSSSPFPHVALPFGSFLWISIANTFHAPGSLQYLVASLVSAGVWISSGTATSG
VZ	
407	
Scaffold 60 F2	
66	CVPSPIOMAWTCAVSARRPT*
>scaffold 5 R26	
54	
	PEGTEPERVYKRYAKEEDGHRISSDTYEIDESASVAGRYARAHIWAESKVDKEYVI EEIGI I YMKPIYVKHNPY
	YQYCSASSSVLVNAMVIKVFVEVHGRVEA*
>scaffold 13 F1	
424	ADQEQGEADPEPTGADPSTYQQAETLGCGEGDLTF*

>scaffold_7_F13	MSVLCLALMLAMATGAFESTGDAKNGVVTPSSRRLLGFTADQEERKFGGPATGTDVSNSHAWENFKAWF
78	KKTFFFWRKREQTRRLRI*
>scaffold_24_R3	MVLFSFCLVYIIILCCYAEKIRPLTLGQDRRLRCAVIAFPVHEKQRSEDTEVDTIPKVFLAKGSSVEMLWHFCAIS
31	ADRRLTFLQRGSLRLRLISRPFLFECAWGIPTFLDVVPAAPARWRRRTSRRAQNRAIRPRREVAVRLDQGGL
>scaffold_11_F9	MPTSRPCSWLHPAALCPVVFPMMLPLLDCREFSNGPFNLAYVIAVGSDSEPPTPIPITTARMCRSGQTPFCP
7	KFRLGCPFFIFSLNCRGRHPHHRALFHRCLRHFLLHYPRRHCPRRWQCPRSVRTS*
>scaffold_6_R29	MVSFFLPFFLFLAFVFFTFSCSSPHHDNEPPSRTDRSTPRSSGQPDLHPLRRLRGKSGSGTARRCILPGQTPSD
10	ASKDVPLRPPPHAS*
>scaffold_25_R1	MRITYILAVTVAATLHSSVTAIPSVKSSKVATENGAVPAVIDSTHTGTGRMLRWVNKYEGDLDKYEGDLDDN
205	DDLDDLDDDLEEERGFSDTLKKANPLKLVKKGTKLTAEQAAKVKQALKDAADYQKMIENANKLIRSD*
>scatfold_12_F7	MRASKLLLMTVFLLASLDATSGYNKLQGTNTEVVDGTQRGVNSERLLRGHPSTVDKVSADDEERASLPTFLK
09	KSTSGATNWLKDKKLAAKLKLRALPMNLDKSVQKTMKDGIDPDRVFTLLNLQKKS
>scaffold_12_F2 959	MRLISIVLVAVATIFACCQGAPTVSDSKSISVPQHHVQTENNANRLLRVHQDEEERTGAQLVKSLSEKFSKKA AEKLTRSKSFNDFVKLDDVAYKENFHSATQALYQRIEKMGFNPDGMLKTMRKRGDVDLDLLAEYTRYWMK KYPTWTKNQ*
>scaffold_5_F36	MFCGVLIVALVQSLFFNFLDLSPSEKKVRYLIELEWWEKATRQNAAKLLQTAWRSGNLRRGTDIGDQCHLFS
83	LMRTARRLRIDKPAIELSVEDQVAEMEATILAEVDRMEAQKVKVLQRIQAKATQLATLKHKLQMKKKK*
>scaffold_495_R	MRLSLLLVAAAVLVANSDAAPQSTSSLTKFSTDVAPVRSLRGASKTKTEVDEDDSFDPEEEERGIAMYGKLDK
1	VDDILKQLNLADDSVIKKLLAKGKGGVEQLITTDKGVKTLVITSKKTGKKVTFTNIDIENIRKIESTPAIK
>scaffold_536_R	MVLMLVSAVCMCSCLLYCAAQRLVNPSQLLPTGVMEVCRHLRWNHWISRRDTALQASDTLQDATYAKGA
3	SACEVTKLCELVSSCRRRGDAGSRLCPA*
>scaffold_5_F17	MALGVAMAITLGLLSLVSNTTAVGTENAVGPVTRHLRSLLLDQKVSEHGAALDQQPLAASMEATSRRSVPQ
32	NPGGPLGGGRTMEAIAPRYEKKAAPETIAEAVAKAEEAEEIAEEALIVALGAENDLQAIEAITEASILSLATNSIT
>scaffold_1_F44 48	MNCSQQSRLLFSLAFCQHSRLIFALVFSQQSRLQCVYDDRILPDEHLRAIRRDVSPLSLYSDRTLRRPSNAP*
>scaffold_53_F9 2	MRLGYFLLATIVGFLACDNATASVSESTSSKLTAREEHPIHGRIGDFTAGHDNKRALRSEDEDGDADDSDDEE RDLILSTIHRPKYWRWFKAGMTPYAVQQVLGLTGVRRLWKPFKRREYKGYVVFYTEQCHKPEYHDFCKKHA DP*
>scaffold_12_R4 2	MRLTSIVLVAAVSIFVCCQGALADSDSKSISVPQHHVHTENNANRLLRVHQDEEERAGPQLVKSLSEKFSKKA AEKLTRSKSFNDFKKLDDVAYKENFHSATQALYQRIEKMGFNPDGMLKTMRKRGDVDLDLLAEYTRYWMR KYPTWTKNQ*
>scaffold_3_R28	MMALLLVGAPWLLPLMLPLPRFDLAPDVTLTPRERALRFSSLRILPSLISSSLANLFTLLRLPMTAPPRGFLPLR
40	*
>scaffold_68_R5	MKTIIASLLLTAVAVNAANGDVNALSAISTISDAKQVSIRALRAAHGSHEHSSGMGSMEDSHDTTSSTHESTV
30	AGDDDDDHDSHDTSSKSSMAGTAGSAGPSNSTESTQAPDTSSAASITVAAGSIFLAAAAAAFL*
>scaffold_24_F1 366	MRLLLVVALTLAAFLAAADVSALNDAPSKRLLRSTVRVDEEEERGMWETLSSKVTKLIKPNQVAIKAMDDPKI AEVAGTSLLTSLKSVNPKKFDSVDGLFSSKAFNNLENYVLRLNKQDINKQTSVAKVFSTGLGDKQAFHLFFTA TQSSDSAVEKSGRFFRDQLLTQWATEGKTWTEVSKSVPKGLPATYYPRLENKYFDILFNLAHDTQKRAARLA RLEKARMAANTAA*
>scaffold_77_R3	MRLTYILALVIAATLHASGTAISTDKSVKIPAIAADDAGRMLRIVKEKPVSDKEAEREERFANPIKKLGAFLKKK
94	WDKQTLKEAIKRDENRRKWIREQGFEPRD*
>scaffold_5_R20	MRLSFVVLPAVIAVTFASSGNAVATADGRNTGLSAITPPNVVASIDTAVGGEKRSLRYHNNKDLEDDSDDEG
89	LEDAEEEERRGNNMFSATKLDEMLDGKQVMSRFKKWKAFGYNTYNLPDAIQARKYDELRKMYRKFLYYN*
>scaffold_12_R9 67	MRLQFAVLFALSVITAANGFSETTAQQFNGELSTGEKQHDEKRILRTEKVDEDDEEGTEGEERVQVSPVSWII DLFTPKTAEQIAEAAKKAEAVKFYTKLANSPSFRAERFPNWKIDGMQVESVLVHLKLWGLDGEKFKAIATKYT EFLASGKLS*
>scaffold_535_F 2	MRLTCILLVAAASLVGVLDASAATTGNTVVANAAMVISPLAPESQGRRSLRLVYDDEDDSADEKDDEEEESA DKVDEERGWLSDKMALTSLVSKFVGKSTDEMGEVIKKLTPAQIDTIFDKGEDSIQKMLPGFYSGMDFQKFD DLIRALPQEQQAVMLSAYTKYLHQNGRFS*
>scattold_29_F1	MIKVFSIFLLVAAATLVASASTESESKQKLDNSPAAPQWRTITENEVPTKRNLRKKKIEEERATTAISLFDDAVK

837	AKGWQVLPYAELANLDATVRTQYLKLLVNNLERKQIVELTGQVPRYVLKHGDSKATRLVQYNKWIFNHFKE
	AVDPAWVLKNYPAFFKGYDKFYQNRFTRGYKYA*
>scaffold_11_F1	MRLPHFLLVATATFLSSYNSTATVSTEGQMMPSADAAVPVRALEANDGKRFLRYYKKEDMYDENDGDDDV
676	DDDDKYDEEVEERGVMTATQVAKWTDKANKWVRLKETLVTIKEKLTNMKGVISAKNREKYNLFTAVYGRA NPHVFERL*
<pre>>scaffold_31_F1</pre>	MRFGLFLALLVATFVACSTVANAESVALTLGDNEVRRLRLNQQNIAKAAGDFISKSKESATLTKAINIAKTANG
184	DEAAARRAVMLAAGAKEGAKLSDETMVKLSAMIAESAKKNPKSWPRLKKFVKITLGIQVGGLAIYGAYKLLF
	DKGSSTAAATTTTTTTSSGAA*
<pre>>scaffold_40_R1</pre>	MAFSSFLKFLAMATLLMVQVNAEPSQQTRNLRIQSDDLPQEQARRLGPNWKALVINVKNNPGVSDGVMK
072	LLKKADLITGTSKKIKEKVKDAAEKVKNSAGKVKDKVTGN*
<pre>>scaffold_62_R1</pre>	MRVTQTLFAIAVVLFATTDNLTTASKATTSSKVQKDGLYAEIYNVEDGVVTKKMRVPLGDVEQYSDTDFEQL
2	KSTLTTEERAAVHLPFGVDSFFKGLEKFVGFFKFKGRRLRIEEA*
<pre>>scaffold_4_R32</pre>	MVKCCAFVVGLVMALASGENNVDVGDVGGNGVKTEAVVRVNATVESLRFPVLFDAIINRSVVLEHSRETRD
84	LRVLSIEREGVVEVEVEVDAATVSSCNATTKACPPEVNDTEIDENSVPPPSLQQTDRLVRVDGENVSTLSFQD
	VIDSAAGSVAVLPLFKPIGSAHVLSGFARVEILSPKLLEFEPAIPEREQVVMAWIEEKARLKAEREAREREIANN
	KELQERLEKERLLAEALAKKEEELLAKLDREEYERTRMTPHNLAAGKRRDGWEFRYEVEFKTKGPIGLNWDL
	NTRDKAVVSHLEPELPAQQLNVIAPRDQIISLNGVDTSKMGPQEVVEVYLSSAIPRKMVFLVQMSAERAAAK
	SAEKNPVKRVVMNWTLAFDAPEVLRGWEVRLHLASWSVPPQINETNASLPLQFELPTPITGCSSFPVPQSS
	NETAGVVYLAYRGACTLVEKAKNARTANGSALLIVNNANGEGRFTPSGTVVVERVDVPVTLYESTFVFIRCIW
	NLICGVCTAG*
<pre>>scaffold_211_R</pre>	MRLSQVLVIAVASFVFASDTVATSNQAKISKTVQSSQSQRLLRSNHYPVKEEEDESEDSVDFEERGFTTPDEE
6	DLEERSPLSAAIVEKLDDIASRWGTSWAAVAMGQSSISEDKIKALLALRDAYLSGNKNAKAAAKLAILRANWT
	RSQQKW*
<pre>>scaffold_80_F4</pre>	MSFLSLFCFTQLLWSCNFQTVKAVFEAYAQTYVRIPPEGTVVFSTALPTCTWKSTDSVFSFITTRILRF*
73	
>scattold_62_R7	
23	
50	
Scooffold 26 D1	
25Callolu_20_K1	
Scaffold 52 P8	
~3canola_52_10	TI OOVOGI FRALRVRHGVOKVI FRHHNHFRRAANRFALLRVVFYGII VGONDGRILLI DR*
>scaffold 33 F2	
96	LOOMPWWSSTHNENRSNNVAAFKPPETPSKRRSI RSELLSI RSS*
>scaffold 16 F1	
464	HPLTNRKLRSLKOSLEVGELLIRELLWVSOASRPRRLMRCLPCETRTCFATGLLSROOR*
>scaffold 2 R38	MRI OSIVI II VATI VATTOATTKI NAPSSETVOISPOREI RKHHTONDEERGAASI IEKAKRVEPSKITOKTI OR
99	WANKKKSPKQALTRLKLDIPGRTSLRSLSSASGPLSWPSATRTTPKKRWSRLYWRNTAMRP*
>scaffold 28 F9	MDTCLSVLLSLYVSACSSQTALPEVLNASVEMASVTVSLYGVGTPSLISRWLRAIAEFYDYPSISQLLDDHFCGL
19	WHRFISASHRPVPQEEENSTGWMKNSQLAPLPNGHSLLQQFPLCILLGEEPGMNTQETHFAKMLDKIVPIG
	VLHSFISGDMSGENGDRFKFVDEILSCFPVDDMNAERKIDFGAQLTTDLFAFSFMLLVYPDPDLKQLAQQM
	VDIAEERAHGNTLQLSHLGHIASKMARFTVWNIIQGCDEDLVTQSDLWRDALKMMKEKYSAFDWKLLNIA
	DLLGEFYVLLLRTEHIDPRAVCAVECFKIFVEETRDAVAESSVLQQLLLSICFQSIKRLAVRNRRRIGRVLSLLVRE
	NCEYFMKSTDKFGKYLGFVVQEISDILSKCNASSQTGRNSVGALSVSADDQAELEWVIFAVCNELGSGLGKH
	ALDIDMVSDGISSSLDKLNALIISSRKVVSSETSQSEEERKKNLSQSAMGTRNQAIQQILMFIQREQSRGAQFY
	NPHPFSGTASAISRETYVNAATQSPAGSLADVRLSAVTTCIDMVRDSSQSTMKLYGKLAQTLLYLSSSGTFGT
	${\tt SRYNCENDAASLANTLGQLGALHASEYEVSPAEQEGELSRLYWRHFHRGALREIKTTFGLVMHENVLTYLSSL}$
	LFEGTRFGHVDPTVVEETLKTLQTVLNLEEGLAALARSKDNELKAFLKPFESSSPSNWSSSYSSSGGDWGVKP

Page	90	of	90
------	----	----	----

	KRTEROFI KHWTSAFFI GEFAWVRSI TACI ARESSDPVI KACSALSAMRVDMAVEI EPYALERII RI DNRDV
	DNPDEEKKSSGVSEPSRIMKAANOGIREVLTGSTGDALSHHVLASLDGTORESEFSOPPEAVOLVVHSINFLR
	ETEKAOFVETNGRGPOTVSTIKGKGSRRSSTYTAASIGROHLNDLAYGCLVDVDFLAVAKAAVRVKMPYSA
	MQYVEMWLEKKQGGKITSLSSLDRDGMVDTVRDILVEAYSFDSDDDGIYGVNDGRTVKSQLVKYNREGLH
	ARALPLYDVSLQFSSQQLVSTDELTLNTPPRLVEGILTSLQSLGYNHLLTGYLQSLQSGDVAGNKSGTAIAQAL
	EHKYKLAWKSMQWEAVLSGLSASGEHSSHHQMIFQGLRAIAHGNFTRLQGITTKAKEQVLRSVQLSLHSFE
	STKDSYSALVRLQAIHEIEELANHIRNSVPPTEPLVFSATTTGSGPFLPSLGGTLAAPRQETLTVLPLLEQWHQR
	RDQIKNDFDKAESLLALEEVLVQVAKPSDNARVVTKLYLDLASLSRKAGRIAIAYRALQKLEHLDERGSLGIYER
	MQCQIQKAKLLWKQQEARSAIWTGKSVSSELTGYLRDTSISATEVTSLQLLLVKVLTFTGKWIAFQRSESSQVI
	LEDFFQKATEIMSNMDPEAVSERSRDAAKAHFALAEFMAGMYQQVSTRVTSQEWLTGKMVVQARHDEL
	QELQSMEQNMQNENRAHIFALNKEVIYDMNERSKVEASVDQFLIGAICSYGKGLTLSLQAELDMVFRVLSL
	WFNNQHKPDINRVVIEEVIDMVPSYKVVPLSYQIISRISSASGTFQTALRKLVMKLSEQHPHHTLIQLIALKNS
	GDVEGKGALQFRTNVGDAKAEGAKVYLTELMKTEQRELLQSLDSIANAYVQLALFDTSEYHGKKKKIPLSTVK
	IFETNSGRSGGTTFDQCLRARARRGDSVVLPAVLTSQIAPQPDMNYSNVVRMYSFEPQFSITDSGIHRPKIIYC
	YGSDGEPYKQLVKGQDDTRQDLVIEQVFETMNQFLMEEKATRKRKLRLRTYRVVPLSPIAGVLEWVENTMP
	WGSYLVSRTSKRLSAHERYHPHEWKHTECRQYLKNAPDKLPAFLEIEANFTPVFHHFFLEKFPDAAVWYQRR
	LSYVQSAAVTSIVGYILGIGDRHSQNILIHEKTGELVHIDFGVVFDQGMALYTPETVPFRLTRDMVDGMGISG
	VDGVFSRCCEVTLQLLRKKSASVVTILEVFVHDPLYRWTLSPLKALRIQEGQGHGKPTRSRSSSRSSGSAEYDG
	TGSMQDTQPADEMHAEPGSTDAAARALIRVKQKLEGYEDPNGSALSIEGQVKQLINAAQDPLNLCKLFPG
	WAPWL*
<pre>>scaffold_29_R3</pre>	MTSSSCRCLLPLWSGVSSVYSDAEIPLRRPFLVFLHRGFWYSRCQVPVVSSLCFRIRTFLSEVPWPPARVTRHR
08	VHRSSGRLRFTLLSRRVCCSYRARRSLRSACRLRAFVRSYRWYSIGRLFSTQVLVFTVLRMDHPHLGDVLLLRL
	LHCRRLLCGALDCGLRLSFSILSDRPIAPISHATIRYIPDAGEFIPNSLWVRENCPVGVSRCRRCRRIHRIPYANT
	WYYHGVSGSRPNRYRIRHLRSYPLLPDVLHPLSYIIKGVVHLRRLVYRLVKCLRLALSH*
<pre>>scaffold_21_F2</pre>	MQRRMSTPFFLMITTIGDAQGEVEWVMIPADNSPLIKSDSFSRNCGATRRTRWATGVALPTSMSQTAPRA
030	AGGRAFPFPIRRCLRFQMICLNRSICTGVLPSASSIWISSILTSSVVALTSGSMAI*
<pre>>scaffold_494_F</pre>	MRLLYLAGVAILAFAADAKVLVSDDSVNNRAQAASASVVSTTRLLRTRSVIDEERAGGISASASDKLAKLFKSS
2	KVTDEQLQKWLNNGKTAESVFYRMNLENTLYTRVFESPQFPRWLQYADDLSASGKGASAISVLSTKYGDDT
	LYKMIGWAKKESSTKALGIRLQTEQLEHWVKIGKDPDEVFKLYKLDYAGNRILGNPQFSAWTKYVDDLNAK
	NKGAFVSIIPTLRKYISDDDLFKIALAAKRSGETKAMGTKLEDAIVQFWNRAGGPVGNGPT*
<pre>>scaffold_100_R</pre>	MRICFVLLLTVAALVTAVSGSSVNLRDSQRIDVVQDKTSSRELRGDLNTDEATEERASFDFVKKLVAKLKGDPL
322	ESFAKRQTKYIFSDDIFDEMLKKFPDPDALYTTLKLHVIKNRSNKYGVTTARHKLHQNFLTSYIDKFPNWRSKL
	N*