Gene expression and proteomic analysis of the formation of Phakopsora pachyrhizi appressoria

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Abstract

Background: Phakopsora pachyrhizi is an obligate fungal pathogen causing Asian soybean rust (ASR). A dual approach was taken to examine the molecular and biochemical processes occurring during the development of appressoria, specialized infection structures by which P. pachyrhizi invades a host plant. Suppression subtractive hybridization (SSH) was utilized to generate a cDNA library enriched for transcripts expressed during appressoria formation. Two-dimensional gel electrophoresis and mass spectroscopy analysis were used to generate a partial proteome of proteins present during appressoria formation.

Results: Sequence analysis of 1133 expressed sequence tags (ESTs) revealed 238 non-redundant ESTs, of which 53% had putative identities assigned. Twenty-nine of the non-redundant ESTs were found to be specific to the appressoria-enriched cDNA library, and did not occur in a previously constructed germinated urediniospore cDNA library. Analysis of proteins against a custom database of the appressoria-enriched ESTs plus Basidiomycota EST sequences available from NCBI revealed 256 proteins. Fifty-nine of these proteins were not previously identified in a partial proteome of P. pachyrhizi germinated urediniospores. Genes and proteins identified fell into functional categories of metabolism, cell cycle and DNA processing, protein fate, cellular transport, cellular communication and signal transduction, and cell rescue. However, 38% of ESTs and 24% of proteins matched only to hypothetical proteins of unknown function, or showed no similarity to sequences in the current NCBI database. Three novel Phakopsora genes were identified from the cDNA library along with six potentially rust-specific genes. Protein analysis revealed eight proteins of unknown function, which possessed classic secretion signals. Two of the extracellular proteins are reported as potential effector proteins.

Conclusions: Several genes and proteins were identified that are expressed in P. pachyrhizi during appressoria formation. Understanding the role that these genes and proteins play in the molecular and biochemical processes in the infection process may provide insight for developing targeted control measures and novel methods of disease management.

Background

Asian soybean rust (ASR), caused by the fungal pathogen Phakopsora pachyrhizi Sydow & Sydow is an aggressive foliar pathogen of soybeans. Initially identified in Asia, it has since spread to all major soybean-growing regions of the world, including the United States [1,2]. The impact of disease on crop yields is influenced by temperature and humidity, spore load introduced into a field, and the growth stage of soybeans when first infection occurs. Field trials in Brazil found yield loss averaging 37% when infection began at R5 growth stage and 67% when infection started at R2 [3]. Observations in Asia have reported yield losses up to 80% under high disease pressure and favorable environmental conditions [4,5].

While fungicide applications are able to reduce yield loses, a limited number of fungicides are available for foliar application on soybeans. The cost of fungicide applications, efficacy of treatments on maturing plants and dense canopies, and environmental impact are all considerations for the soybean grower. Six resistance genes (Rpp1-Rpp6) have been identified that provide resistance
to *P. pachyrhizi* [6-11]. However, these genes display only race-specific resistance to selected isolates of *P. pachyrhizi*.

When a urediniospore makes contact with a soybean leaf, a single germ tube elongates across the leaf surface. At the tip of this germ tube a specialized infection structure, the appressorium, is formed. While most rust pathogens penetrate the host indirectly by entering through stomatal openings and then breaching the mesophyll cell wall, *P. pachyrhizi* is one of a few rusts that enter the host by direct penetration of the cuticle and epidermal cell wall [12]. Recent transmission electron microscopy confirmed that *P. pachyrhizi* uses mechanical force to penetrate the cuticle and digestive enzymes to penetrate the epidermal cell wall [13]. The fungus continues to grow invasively in the host forming haustoria, colonizing hyphae, and ultimately uredinia.

In a previous study, a cDNA library was utilized to evaluate gene expression during urediniospore germination of *P. pachyrhizi* [14]. A subsequent independent study examined the partial proteome of germinated urediniospores [15]. The next critical step in the infection cycle is appressorium formation. *P. pachyrhizi* infects over 90 species of legumes, and this broad host range is unique among the rusts [16]. Elucidating the events during appressorium formation could shed light on the mechanism allowing broad-spectrum interaction.

Obligate pathogens require a living host on which to survive and propagate, making it difficult to separate fungal-specific genes or proteins from those of the host. However, induction of appressoria by surface contact with artificial substrates is possible for some fungi, including *P. pachyrhizi* [17,18]. Despite the ability to induce appressorium formation *in vitro*, a review of the literature found appressoria-specific EST libraries generated only for the plant pathogens *Magnaporthe grisea*, *Puccinia triticina*, and *Colletotrichum higginsianum* [19-21].

Several studies have utilized bioinformatics to identify appressoria proteins from EST sequencing projects [21,22], but only a few have used a proteomics approach to identify differentially expressed proteins accumulated during appressorium formation. For example, comparison of protein expression patterns in germinating conidia to those observed in appressoria revealed five proteins that were up-regulated during appressorium formation in *M. grisea* [23]. Protein profiles of three developmental stages of *Phytophthora infestans* (cysts, germinated cysts, and appressoria-forming cysts) found 13 proteins to be up- or down-regulated during different developmental stages [24]. Likewise, another study of *P. infestans* identified four up-regulated genes and their protein products in cysts with appressoria [25].

This study identified ESTs and proteins present during, and possibly required for, appressorium formation in *P. pachyrhizi*. This is one of the few studies to evaluate genes or proteins present specifically during appressorium formation and the first to combine the two techniques. The comparison of identified transcripts to accumulated proteins allows for a more comprehensive analysis of the molecular and biochemical processes occurring during appressorium formation.

**Methods**

**Fungal strain and growth conditions**

*P. pachyrhizi* isolate Taiwan 72-1 was maintained at the USDA-ARS Foreign Disease-Weed Science Research Unit Plant Pathogen Containment Facility at Fort Detrick, MD [26] under Animal and Plant Health Inspection Service permit. Urediniospores were obtained and maintained as described previously [27].

Urediniospores were germinated by floating on the surface of sterile distilled water containing 50 μg/ml each of ampicillin and streptomycin in a 9” x 13” glass baking dish, for 6 h at room temperature in the dark. Germinated urediniospores were collected onto Whatman No. 1 filter paper (Whatman; Piscataway, NJ) and flash frozen in liquid nitrogen.

Appressoria were generated by application of 150 ml of a 100,000 urediniospore/ml suspension onto a 245 mm x 245 mm x 20 mm polystyrene dish, followed by incubation in the dark at room temperature for 6 h. Water was decanted from the plates and collected for protein extraction. This collected water will subsequently be referred to as the Appressoria Water Fraction (AWF). A sterile water wash was used to remove urediniospores that did not germinate from the plate. Plates were examined under a microscope before and after washing to confirm the presence of appressoria. Appressoria, germ tubes, and germinated urediniospores were collected by scraping the plate surface with a cell scraper (Nunc, Thermo Fisher Scientific; Rochester, NY) and immediately frozen and stored in liquid nitrogen.

**RNA isolation and library construction**

Germinated urediniospores and appressoria were each separately ground under liquid nitrogen with a mortar and pestle. Total RNA was isolated by phenol:chloroform extraction [28], and precipitated overnight with 12 M lithium chloride.

Purification of poly(A)* mRNA, synthesis of cDNA, and library construction was performed by SeqWright (Houston, TX) following standard protocols for library construction via suppression subtractive hybridization [29]. The driver for the subtracted library was generated from 1400 μg of total RNA extracted from germinated urediniospores, and 1400 μg of total RNA from appressoria was used as the tester. The subtracted-cDNA pool was cloned into pBluescript II KS(+) vector (Stratagene;
La Jolla, CA) and transformed into DH10B E. coli strain (Invitrogen; Carlsbad, CA). Average insert size for the library was determined by plasmid mini-prep, PCR using T3 and T7 primers, and gel electrophoresis of 15 randomly selected colonies from the library (performed by SeqWright).

Sequence analysis
Single-pass sequencing of clones from the cDNA library was performed on an Applied Biosystems 3700 DNA analyzer (Applied Biosystems; Framingham, MA) at the USDA-ARS Eastern Regional Research Center Nucleic Acids Facility (Wyndmoor, PA). Initial sequence analysis was performed at the Advanced Biomedical Computing Center at the National Cancer Institute-Frederick (ABCC/NCI-Frederick) (Frederick, MD). Sequenced clones were subjected to BLASTN analysis against all non-chordate ESTs in the GenBank database to identify clones with similarity to existing P. pachyrhizi ESTs [14, http://www.ncbi.nlm.nih.gov/nucest/?term=txid170000 [Organism:nonexp]]. Clones that did not share similarity to these P. pachyrhizi ESTs were subjected to additional bi-directional sequencing, using anchored poly(T)N primers to read through poly(A) tails. Additional primers, as needed, were designed using Primer3 (http://frodo.wi.mit.edu/primer3/) [30]. Subsequent sequence analysis and compilation of full-length sequences was performed using Chromas 2.33 (Techelysium Pty; Helensvale, Australia). Putative functions were assigned to clones by BLASTX analysis against the NCBI non-redundant protein database. Putative functional categories were determined using the Munich Information Center of Protein Sequences (MIPS) Functional Catalogue (http://mips.helmholtz-muenchen.de/proj/functatDB/search_main_frame.html).

Redundancy within the subtracted library was identified by BLASTN analysis of the library against itself. Redundant and overlapping ESTs were assembled into contigs using the program CAP3 (http://pbl.univ-lyon1.fr/cap3.php) [31].

Quantitative real-time RT-PCR analysis of transcript levels during the infection cycle
The soybean cultivar Williams 82 was inoculated with P. pachyrhizi isolate Taiwan 72-1, as described previously [27]. Leaves were collected from three biological replicates at 0, 6, 12, 24, 72, 168, and 336 h post inoculation (hpi). Freshly germinated urediniospores and appressoria, separate from that used to generate the cDNA library, were produced as described above. Total RNA was isolated from 100 mg of each sample and 100 mg of urediniospores using the RNeasy Mini Plant kit (Qiagen; Valencia, CA) following the manufacturer’s protocol. Six ESTs specific to the appressoria-enriched cDNA library and representing four putative functional categories and one unclassified protein were selected for transcript analysis. Three ESTs common to the appressoria-enriched cDNA library and the germinated urediniospore library [14] were also selected for analysis. Nucleotide sequences of each clone were compared to the Trace Archives for P. pachyrhizzi Whole Genome Shotgun (WGS) sequences by BLASTN to identify the positions of potential introns. Primers were designed to span putative introns where possible, using the primer design program Primer3 (http://frodo.wi.mit.edu/primer3/).

To assess transcript levels quantitative real-time RT-PCR (qRT-PCR) was performed using three biological replicates and two technical replicates for each template. First-strand cDNA synthesis was performed using the QuantiTect Reverse Transcription kit (Qiagen) following the manufacturer’s protocol. Real-time PCR reactions were performed on the SmartCycler System (Cepheid; Santa Clara, CA) using the QuantiTect SYBR Green PCR kit (Qiagen) following the manufacturer’s protocol. Sequences and annealing temperature for each primer set are listed in Table 1. Melt curve analysis was performed to verify specificity of the PCR products, and control reactions containing no template or DNA without reverse transcriptase were included. Absolute quantification of target molecules was conducted using the sigmoidal model, and lambda gDNA was used to generate the optical calibration factor (OCF) [32]. Primers for α-tubulin were included to assess RNA integrity and demonstrate functionality of the RT-PCR assay [33]. For each time point, the average number of target molecules and standard deviation was calculated using the three biological replicates. Data were analysed by analysis of variance.

Quantitative real-time RT-PCR analysis of transcript levels during urediniospore germination, germ tube elongation, and appressoria formation
Urediniospores were germinated as described above on the surface of water for 6 and 24 h, and on an appressoria-inductive surface for 6 and 24 h. Total RNA was isolated from 100 mg of each sample and from 100 mg of urediniospores using the RNeasy Mini Plant kit (Qiagen) following the manufacturer’s protocol. Twelve target genes common to the appressoria-enriched cDNA library and the germinated urediniospore cDNA library were selected for transcript analysis, and primers were designed as described above. Quantitative real-time RT-PCR was performed on three biological replicates and three technical replicates for each target using the SmartCycler System as described above for qRT-PCR during the infection cycle. Sequences and
| Clone/contig | Target gene product | Primer sequences | Amplicon size (bp) | Primer (nM) | Annealing temperature (°C) |
|-------------|---------------------|-----------------|-------------------|------------|--------------------------|
| Pp3004⁰ | serine/threonine-protein kinase | Forward: GGAACCAACGTCGACAAGAG <br> Reverse: CTGGTCCTCCTCAACCTCAG | 130 | 200 | 60 |
| Pp3243¹ | prefoldin subunit 5 | Forward: AACTGTACAACTGGCATTC <br> Reverse: CTTTAGTGCCCCAAACGAG | 81 | 200 | 59 |
| Pp3282¹ | ubiquitin-protein ligase | Forward: AAATTCTGGAGACAGATTGC <br> Reverse: TGGCACTACCTTGTGGGTGA | 150 | 300 | 60 |
| Pp3495¹ | G2/M phase checkpoint control protein Sum2 | Forward: CAGGCTGAACAGACGATCC <br> Reverse: GCCGGAAAGAACGATAAATTC | 105 | 200 | 58 |
| Pp3505¹ | hypothetical protein | Forward: GAGACAGGCCATGGAGAG <br> Reverse: GGTCAGTGGCCTCACTG | 104 | 200 | 60 |
| Pp3684¹ | HMG-CoA reductase | Forward: CGACAGCTTGCTCATGATTC <br> Reverse: CTTAACGAAGTTCGAGCTGC | 87 | 200 | 60 |
| contig2F¹ | S-aminolevulinate synthase | Forward: GATGAGGTTCACGCCATTG <br> Reverse: GTCCACCGATTCATACAC | 117 | 200 | 60 |
| Pp3186¹ | hypothetical protein | Forward: AAGGACTGGTGAGGTTGGTG <br> Reverse: TGCTCTACCACCGTAGGACC | 76 | 200 | 60 |
| contig2N¹ | P-type cation-transporting ATPase | Forward: GCCGGAAGGGAGGGTCGAG <br> Reverse: GCCGAAATGTGACATTACG | 124 | 400 | 55 |
| Pp3042² | GTPase-binding protein | Forward: TGCCAGAAAATGGTTCTCTC <br> Reverse: GCCAAGAATGTGACATTACG | 102 | 300 | 58 |
| Pp3205² | NADPH oxidase A | Forward: ACCGAAAGCTGCCATATTGA <br> Reverse: GGTCGGCTTCCATATAAAACAA | 134 | 300 | 56 |
| Pp322³ | septin/cell division control/GTP binding protein | Forward: TCCTCCCAAGAGAACCATCCAG <br> Reverse: AAGTCTCCATAGGCCGGAGT | 135 | 300 | 60 |
| Pp3409² | MFS sugar transporter | Forward: CCGATGTACATGGAGACCTG <br> Reverse: CTTGGAATGCCACCATGATG | 123 | 300 | 60 |
| Pp3717² | tripeptidyl-peptidase 1 precursor | Forward: GCGCGAGGGTTTTTCATAAT <br> Reverse: CTCTGGACCCATCGAACAT | 116 | 300 | 59 |
| Pp3888² | microtubule associated protein | Forward: GGGATCAGCTCTACGTGC <br> Reverse: AGATTGGCCACGACCTTTT | 150 | 300 | 60 |
| Pp3944² | autophagy-related protein 8 | Forward: CCGATATCTTGACTTGAGAG <br> Reverse: CATAGCAAGTGGACACTG | 107 | 300 | 60 |
| Pp3998² | delta (12) fatty acid desaturase | Forward: CCCATCTCCATCCATCCAC <br> Reverse: TTGGACACAGATGATG | 135 | 300 | 59 |
| Pp4000² | subtilase-type proteinase psp3 | Forward: TCATTTGACATTGGAAACGAG <br> Reverse: GCAACCCTAGGAGGGTCTT | 93 | 300 | 60 |
| contig2S² | acyl-CoA dehydrogenase | Forward: AATGCAACCCGCTCTATGC <br> Reverse: CTTGATTGACGATTACCAAGG <br> contig6B² | conidiation-related protein | Forward: AATGCAAGAAGTGCGAGAC <br> Reverse: TGGCAAGCTCCTTCTTTT | 107 | 300 | 60 |
| Pp481² | MAS3 protein | Forward: CGTATGAGTAGCTCAAGCAGAC <br> Reverse: CCTGAAACCTACGGTGCTTC | 68 | 200 | 62 |
annealing temperature for each primer set are listed in Table 1. Primers for α-tubulin were included to assess RNA integrity and demonstrate functionality of the assay [33]. For each time point, the average number of target molecules and standard deviation was calculated using the three biological replicates. Data were analysed by analysis of variance.

Protein extraction
Appressoria-enriched samples weighing 300 mg were ground to a fine powder in liquid nitrogen and suspended in 1 ml of isoelectric focusing (IEF) buffer containing 7 M Urea, 2 M Thiourea, 4% (3-[(3-Chloramidopropyl) dimethylammonio] propanesulfonate) (CHAPS) and 25 mM dithiothreitol (DTT). The suspension was mixed thoroughly and incubated at room temperature for 30 min on a shaker at 100 rpm. Samples were centrifuged for 15 min at 14,000 g and the supernatant was collected. This protein sample will be referred to as the appressoria-enriched fraction (AEF). The AWF was collected as described above to a final volume of 500 ml per biological replicate and applied to a 0.45 μm filter units (Millipore).

Table 1 Primer pairs used for real-time quantitative reverse transcription PCR analysis of expression patterns of Phakopsora pachyrhizi ESTs (Continued)

| NA  | P. pachyrhizi Forward  | CCAAGGCTTCTTCGTGTTTC  | 65  |
|-----|-------------------------|------------------------|-----|
| α-tubulin Forward  | CCAAGGCTTCTTCGTGTTTC  | 65                      |
|       Reverse       | CAAGAGAAAGAGCCCAAACC  | 200                     |
|       Reverse       | CAAGAGAAAGAGCCCAAACC  | 60                      |

1 Primer pairs used for quantification of mRNA transcripts throughout the infection cycle.
2 Primer pairs used for quantification of mRNA transcripts during germination, germ tube elongation, and appressoria formation.
3 Primer pairs used for both quantitative analyses.

Two-dimensional gel electrophoresis and sample preparation
Two-dimensional gel electrophoresis (2-DE) was carried out as previously described [15]. Prior to 2-DE, samples were diluted to the appropriate protein concentration in IEF buffer and 1.0% pH 3-10 carrier ampholytes (Invitrogen) were added. For first dimension 400 μg of AEF or 200 μg of AWF total protein was applied to a 13 cm pH 3-10NL Dry Gel Strip (G. E. Healthcare; Piscataway, NJ). For second dimension electrophoresis, strips were separated on 4-12% Bis-Tris gels (BioRad; Hercules, CA) and stained with Simply Blue SafeStain (Invitrogen). Gels were run from two biological replicates of AEF and 385 protein spots were picked. A total of 150 spots were picked from gels run from three biological replicates of AWF. Spots were excised from gels manually using sterile filtered pipette tips (Mettler-Toledo; Columbus, OH). Selected protein spots were destained for 30 min at 37°C in 200 μl of a solution containing 2 mg/ml NH4HCO3 in 50% ACN/H2O (v/v). This step was repeated as necessary until complete removal of visible coloration. Following destaining, samples were dehydrated in 50 μl of 100% ACN solution for 15 min at room temperature. The spots were then air-dried for 10 min at room temperature, followed by trypsin-digestion for 16 h at 37°C in 20 μl of a buffer containing 20 μg/μl activated trypsin (Promega; Madison, WI), 10% ACN and 40 mM NH4HCO3 for 16 h at 37°C.

Peptides were cleaned and spotted on Matrix Assisted Laser Desorption Ionization (MALDI) plates using the Digilab/Investigator ProMS MALDI Preparation Station (Genomics Solutions; Ann Arbor, MI) programmed for peptide cleanup using the C18 ZipTip procedure according to the manufacturers recommendations (Millipore).

Mass spectrometry
Trypsin-digested proteins were subjected to mass spectrometry analysis using a 4700 Proteomics Analyzer instrument (MALDI-Time of Flight (TOF)/TOF) (Applied Biosystems) in the positive reflectron mode as previously described [15]. Spectra between the range of 800 and 4000 Da in MS mode were acquired through the averaging of 1000 spectra, and 2000 spectra in the MS/MS mode. Up to 10 most intense ions were selected for MS/MS analysis using post-source decay (PSD) with 1 keV acceleration voltage. Criteria for ion selection were based in a signal-to-noise ratio cutoff of 20 and exclusion of all common trypsin autolysis peaks and common keratin contaminants. Instrument conversion of time-of-flight to mass (Da) for the monoisotopic ions was obtained with a tolerance of 50 ppm or better according to calibration with a peptide calibration mixture (Applied Biosystems). The MS/MS TOF calibration was optimized to 0.1 Da or better from the PSD of Glu1-fibrinopeptide B fragments.

Combined MS and MS/MS data were submitted for analysis using GPS Explorer version 3.6 software (Applied Biosystems) with Mascot version 2.3.02 search engine (Matrix Science; Boston, MA) against a custom database.
sequence database of EST sequences in FASTA format. The analyses criteria included the following variable modifications: methionine oxidation, formation of pyroglutamine from N-terminal glutamine, and carbamidomethylation of cysteine residues from the reduction and alkylation of proteins. Reported proteins from database searches of putative peptide/protein sequences are within ≥95% confidence interval.

The database was constructed from a subset of the NCBI EST database compiled in March 2010 using the keywords Rust or Basidiomycota. The dataset was combined with EST sequences from the appressoria-enriched SSH library described above for a total of 280,119 sequences. Identification of proteins were validated by reanalyzing all samples using a decoy database composed of randomized entries of the corresponding original database.

Database analysis
Protein scores are based on the sum of the ion scores for the peptide mass fingerprints and MS/MS of selected peptides. A match score greater than the protein score threshold of 75 is considered significant within a 0.05 probability. The full data set from these analyses is included in the Additional file 1: Table S1.

Putative protein identities were assigned to ESTs based on BLAST searches against the NCBI nr protein database using open reading frames (ORF) corresponding to the peptides identified in each accession. BLASTX was performed using EST accessions with the highest protein score to identify protein homologues. Proteins were categorized using Uni-Prot Protein Knowledge Database [35] and placed into functional categories using the Munich Information Center for Protein Sequences (MIPS) Functional Catalogue. Proteins with multiple functions were given secondary classifications. Peptide sequences, protein BLAST identities and functional categories for each protein are listed in Additional file 1: Table S1. A comparison of Additional file 1: Table S1 to a data set of P. pachyrhizi germinated urediniospores [15] identified a subset of different proteins for further analysis.

Proteins with putative signal peptides containing peptide cleavage sites were identified using SignalP 3.0 [36]. Target P [37] and PSORT [38] were used to predict intra- or extracellular localization of the proteins.

Results
Fungal strain and growth conditions
After 6 h on polystyrene plates, germination rates of urediniospores averaged 60%, with 77% of germinated urediniospores showing mature, melanized appressoria (Figure 1). Washing plates with distilled water raised the percent appressoria on each plate to an average of 86%.

Less than 1% of the urediniospores remaining on the plates following the wash step were ungerminated. The remainder of the urediniospores were germinated with either immature or no appressoria.

Sequence analysis and annotation of expressed genes
Single pass sequencing of 1133 cDNA clones led to the identification of 1029 ESTs. The sequences of the P. pachyrhizi EST clones were submitted to NCBI as GenBank Accession numbers JK649959 to JK650987. The remaining 104 clones sequenced contained either no inserts or consisted of poor quality sequence and were discarded. A total of 238 non-redundant ESTs were identified (Additional file 2: Table S2), of which 169 appeared only once and 69 were represented by multiple clones at frequencies ranging from 2 to 481. The frequency of redundant clones is shown in Figure 2. Assembled sequences of the redundant ESTs were submitted to NCBI as GenBank Accession numbers JQ863574 to JQ863646. Singleton sequences ranged from 129 to 1217 bp, and assembled contigs ranged from 424 to 1781 bp.

BLASTN analysis of single pass sequences identified 41 ESTs with no homology to P. pachyrhizi ESTs from germinated urediniospores and P. pachyrhizi-infected soybean leaves. Bi-directional sequencing was performed on these 41 ESTs, and the sequences were submitted to NCBI as GenBank Accession numbers JQ083238 to JQ083278. Of these, 29 non-redundant ESTs were identified with no homology to P. pachyrhizi. Twenty-seven ESTs were singletons, and two ESTs occurred twice.
Sequence similarity comparisons of the 29 non-redundant ESTs to the NCBI non-redundant protein database identified 26 ESTs, which fell into eight functional categories (Table 2). Twenty-four ESTs shared identity to proteins from members of the fungal phylum Basidiomycota, and one EST had similarity to a member of the Ascomycota. The remaining EST had identity to an IS10 transposase from tomato. Three ESTs (Pp3394, Pp3734, Pp3842) showed no significant similarity to any protein entries in GenBank.

Quantitative real-time RT-PCR analysis of transcripts during soybean infection

Six ESTs specific to the appressoria-enriched cDNA library were selected for analysis of transcription levels during the infection process on soybean. Transcripts were assessed via real-time RT-PCR as a measure of target molecule for each of these ESTs (genes): Pp3684 (hydroxymethylglutaryl coenzyme A (HMG-CoA) reductase), Pp3495 (G2/M phase checkpoint control protein Sum2), Pp3243 (prefoldin subunit 5), Pp3282 (ubiquitin-protein ligase), Pp3004 (serine/threonine-protein kinase), and Pp3505 (a hypothetical protein). The sequences represented four protein functional categories and one unclassified hypothetical protein. Three ESTs (Pp3004, Pp3243, Pp3842) showed no significant similarity to any protein entries in GenBank.

Quantitative real-time RT-PCR analysis of transcripts during urediniospore germination, germ tube elongation, and appressoria formation

Twelve ESTs common to both the appressoria-enriched cDNA library and the germinated urediniospore cDNA library were selected for transcript analysis during urediniospore germination, germ tube elongation, and appressoria formation to test if genes expressed in germinated urediniospores were expressed at higher levels in appressoria. The ESTs (genes) evaluated were: Pp3042 (small GTPase-binding protein), Pp3222 (septin; cell division control/GTP binding protein), Pp3717 (tripeptidyl-peptidase 1 precursor), Pp3944 (autophagy-related protein 8), Pp4000 (subtilase-type proteinase psp3), Pp3205 (NADPH oxidase A), contig2S (acyl-CoA dehydrogenase), contig481 (Magnaporthe appressoria specific 3 (MAS3) protein), Pp3409 (major facilitator superfamily (MFS) sugar transporter), Pp3998 (delta 12 fatty acid desaturase), Pp3888 (microtubule associated protein), and contig6B (conidiation-related protein). All of the genes were expressed in the five samples, and the expression patterns fell into four groups (Figure 4).

The largest group consisted of the ESTs Pp3042, Pp3205, Pp3222, Pp3717, Pp3944, Pp4000, and contig2S had lowest transcript levels in urediniospores (Figure 4A).
| Clone/contig | Accession No. | Description | Species | E value |
|-------------|--------------|-------------|---------|---------|
| Pp3070      | EGG05957     | family 1 Carbohydrate esterase | Melampsora larici-populina | 5.00E-9  |
| Pp3292      | EFP76188     | FLU1-II (Glutamine synthetase) | Puccinia graminis f. sp. tritici | 8.00e-56 |
| Pp3391      | XP_001833926 | Palmityltransferase AKR1 | Coprinopsis cinereaOkayama | 9.00E-141 |
| Pp3684      | XP_571450    | Hydroxymethylglutaryl-CoA reductase (NADPH) | Cryptococcus neoformans var. neoformans | 3.00E-65 |
| Pp3741      | EGG05960     | family 4 carbohydrate esterase | M. larici-populina | 8.00E-14 |
| Pp3942      | EFP86206     | mannosyl phosphorylinositol ceramide synthase SUR1 | P. graminis f. sp. tritici | 1.00E-128 |
| Pp3291      | XP_001834782 | DNA replication complex GINS protein | C. cinereaOkayama | 1.00E-08 |
| Pp3495      | EGD98306     | G2/M phase checkpoint control protein SUM2 | Trichophyton tonsurans | 6.00E-26 |
| Pp3590      | EFP88202     | DNA repair protein RAD51 | P. graminis f. sp. tritici | 3.00E-161 |
| Pp3976      | XP_003195178 | Chromatin remodeling-related protein | Cryptococcus gattii | 3.00E-45 |
| Pp3243      | XP_001838238 | Prefoldin subunit 5 | C. cinereaOkayama | 3.00E-26 |
| Pp3282      | XP_567169    | Ubiquitin-protein ligase | C. neoformans var. neoformans | 2.00E-26 |
| Pp3060      | XP_001883758 | Cytoplasmic dynein intermediate chain | Laccaria bicolor | 2.00E-128 |
| Pp3772      | XP_571408    | Membrane transporter | C. neoformans var. neoformans | 3.00E-20 |
| Pp3004      | XP_571731    | Serine/threonine-protein kinase orb6 | C. neoformans var. neoformans | 0.00 |
| Pp3435      | EFP87863     | Inositol hexakisphosphate kinase 2 | P. graminis f. sp. tritici | 8.00E-96 |
| Pp3508      | XP_003192457 | Heat shock transcription factor 2 | C. gattii | 1.00E-19 |
| Pp3161      | EGG09840     | hypothetical protein | M. larici-populina | 6.00E-96 |
| Pp3267      | EFP87812     | hypothetical protein | P. graminis f. sp. tritici | 1.00E-6 |
| Pp3464      | EFP91488     | hypothetical protein | P. graminis f. sp. tritici | 1.00E-19 |
| Pp3502      | XP_569331    | hypothetical protein | M. larici-populina | 4.00E-115 |
| Pp3505      | XP_002396863 | hypothetical protein | M. larici-populina | 7.00E-37 |
| Pp3853      | EGG01299     | hypothetical protein | M. larici-populina | 5.00E-25 |
| Pp3884      | EFP87187     | hypothetical protein | P. graminis f. sp. tritici | 1.00E-24 |
| contig2bb   | EFP88656     | hypothetical protein | P. graminis f. sp. tritici | 3.00E-73 |
Both germinated urediniospore and appressoria samples collected at 6 h yielded high transcript levels, with germinated urediniospores tending to be higher than appressoria. The transcript levels in germinated urediniospore and appressoria samples collected at 24 h were lower than transcript levels from germinated urediniospore and appressoria samples collected at 6 h.

The second group, ESTs Pp3409 and Pp3998, had lowest transcript levels in appressoria samples collected at 6 and 24 h (Figure 4B). For Pp3409 the highest transcript level was from germinated urediniospores collected at 6 h, with the corresponding 24 h samples showing 33% fewer transcripts. Appressoria samples collected at 6 h and 24 h showed 75% and 85% fewer transcripts than germinated urediniospores at 24 h, while urediniospores showed 47% fewer transcripts. For Pp3998, the highest transcript level was from germinated urediniospores collected at 24 h, while the corresponding 6 h samples had 38% fewer transcripts. Appressoria collected at 6 h and 24 h showed 67% and 50% fewer transcripts than germinated urediniospores at 6 h. Transcript levels from urediniospores were 27% less than that found in 24 h germinated urediniospores.

The third group, consisting of ESTs Pp3888 and contig481, and α-tubulin, had less change in transcript levels across all samples than any of the other groups (Figure 4C). The highest transcript levels were observed in germinated urediniospores collected at 6 h.

The EST contig6B comprised the fourth group. The highest transcript levels were in urediniospores and lowest transcript levels occurred in appressoria (Figure 4D). Transcript levels in appressoria at 6 h and 24 h were only 9% and 7% that of urediniospores. Transcript levels in germinated urediniospores at 6 h were 66% that of urediniospores, and decreased to 35% by 24 h.

**Similarity to *Magnaporthe grisea* appressoria specific protein 3 (MAS3) and fungal proteins with predicted signal peptide sequences**

BLASTX analysis revealed five ESTs with similarity to MAS3, which was the third most redundant EST from an appressoria cDNA library of *Magnaporthe grisea* [39]. SignalP predicted all five ESTs contain putative signal peptide sequences. Contig481 had the highest identity to MAS3 at 46% (Figure 5). Among the ESTs, contig481 and contig2Y share 100% identity at the amino acid level,
and contig19 has 70% identity to contig481. In addition, contig19 showed the highest identity (51%) to gEgh16 of Blumeria graminis. These ESTs also share identity to three secreted proteins from Melampsora larici-populina and to a putative secreted protein from Puccinia graminis f. sp. tritici (Figure 5). BLASTP analysis of the open reading frames of the ESTs revealed that all have similarity to the protein superfamily DUF3129, which is a eukaryotic family of proteins with no known function.

MALDI-TOF/TOF analysis

A total of 535 spots were excised from five 2-DE gels and analyzed in triplicate by MALDI-TOF/TOF. Data from these analyses are included as Additional file 1: Table S1. Protein identifications were validated when re-analysis of the dataset against a decoy database resulted in zero false positives. Combined MS and MS/MS analyses resulted in 480 putative identities within the established threshold limits (Additional file 1: Table S1). A
Figure 5  Alignment of predicted amino acid sequences of *Phakopsora pachyrhizi* ESTs with similarity to MAS3 and other putative secreted fungal proteins. The predicted amino acid translation of five EST contigs from the *P. pachyrhizi* appressoria-enriched cDNA library aligned with MAS3 of *Magnaporthe grisea* (AAK52794), GEgh16 from *Blumeria graminis* (AAB05211), and putative secreted proteins from *Puccinia graminis* f. sp. *tritici* (XP_003331958) and *Melampsora larici-populina* (EGF99900, EGF99871, EGG08588). Asterisks indicate amino acid identities, and dots denote conserved amino acid substitutions. Putative signal peptide sequences are in bold.
total of 256 protein spots were identified in replicate gels of the AEF and AWF. Eliminating redundancy within each sample type yielded 140 proteins: 115 and 25 proteins were identified from the AEF and AWF, respectively (Additional file 3: Table S3). Four of the proteins were unique to AWF, and 55 were found only in the AEF. A total of 119 different proteins were identified in this study, 59 of which were not found in the previous analysis of proteins from germinated urediniospores of *P. pachyrhizi* [15] (Table 3). These proteins have been designated as predicted *Phakopsora pachyrhizi* proteins (PHAP).

A search of the custom EST database found that 110, or 92%, of the 119 proteins shared similarity to putative proteins from rusts. Of these 110 proteins, 90 had identity to *P. pachyrhizi* ESTs, 15 of which were from the appressoria-enriched cDNA library. The remaining 20 proteins had similarity to ESTs from *Melampsora*, *Uromyces* and *Puccinia* spp. Of the nine proteins that were not similar to any rust sequences, four had similarity to *Listilago maydis* (cytochrome C peroxidase, GTP binding protein, a hypothetical protein, and nucleosome assembly protein), two to *Sporobolomyces roseus* (ATPase delta subunit and ubiquinol-cytochrome c reductase), one to *Leucosporidium scottii* (methylene-tetrahydrofolate dehydrogenase), and two to *Microbotrium violaceum* (serine-threonine phosphatase and ubiquitin). BLASTN analysis of these nine proteins against the *P. pachyrhizi* trace archive sequences in GenBank revealed accessions with high identity to all nine proteins. The *P. pachyrhizi* genomic traces for all nine proteins shared significant amino acid similarity to *Melampsora* ESTs (gi|120507777) were also found to share significant amino acid similarity to PHAP0129. Alignment of the predicted proteins of these six ESTs revealed two conserved regions (Figure 6).

Eight proteins listed in Table 3 were predicted to contain classical secretion signals [36]. Three proteins, PHAP0038, PHAP0073, and PHAP0113, shared similarity to well-characterized intracellular proteins related to trafficking and proteolysis. PHAP0038 was identified as a glucose-regulated protein that is secreted into the ER and is involved in the assembly of protein complexes for secretion or translocation to membranes [40]. PHAP0113 was identified as a vacuolar protease A with aminopeptidase activity [41], and PHAP0073 was identified as a neddylin/ubiquitin which is required for protein assembly in the ubiquitination pathway [42].

The remaining five proteins containing a predicted secretion signal were proteins of unknown function. BLAST searches of the custom EST database identified *P. pachyrhizi* ESTs containing full-length open reading frames for four of the proteins, and a *U. maydis* EST with an ORF for the other protein. Subsequent BLASTN analysis of the *U. maydis* EST sequence identified homologous *P. pachyrhizi* genomic sequences. BLASTX analysis using *P. pachyrhizi* EST sequences corresponding to PHAP0052 and PHAP0059 found significant similarity to a *Melampsora larici-populina* nuclear membrane protein with a conserved putative stress response domain (EGG05479.1) and hypothetical protein containing a conserved fasciclin domain (EGG10923.1), respectively. BLASTX analysis using the *P. pachyrhizi* EST corresponding to PHAP0054 revealed similarity to a conserved hypothetical protein of *U. maydis* (XP756219.1).

Two of the *P. pachyrhizi* ESTs, gi|120521555 and gi|120521631, encode for the small molecular weight proteins PHAP0129 and PHAP0055 with 132 and 71 amino acids, respectively. Both proteins possess secretion signals with no motifs for localization to organelles, suggesting they may function as extracellular fungal effectors [43]. EST gi|120521631 did not share similarity to any DNA or amino acid sequences currently in GenBank and is a unique *P. pachyrhizi* extracellular protein (PHAP0055). BLASTX analysis using the EST for PHAP0129 identified sequence similarity to ESTs from *Puccinia triticina* (gi|282831716) and *Uromyces viciae-fabae* (gi|164246325). The translated ORFs of three *P. pachyrhizi* ESTs (gi|120499561, gi|120520239, gi|120507777) were also found to share significant amino acid similarity to PHAP0129. Alignment of the predicted proteins of these six ESTs revealed two conserved regions (Figure 6).
Table 3  Putative proteins identified during *Phakopsora pachyrhizi* appressoria formation grouped into functional categories using the Munich Information Center of Protein Sequences (MIPS) Functional Catalogue

| Protein ID | GeneInfo Identifier No. | Description | Species | Peptides | Score |
|------------|-------------------------|-------------|---------|----------|-------|
| **01. Metabolism** | | | | | |
| PHAP0002  | gi|254121616 | 2-isopropylmalate synthase | *Melampsora larici-populina* | 4 | 151 |
| PHAP0008  | gi|120528021 | 5-methyltetrahydropteroyltriglutamate-homocysteine methyltransferase | *Phakopsora pachyrhizi* | 3 | 116 |
| PHAP0010  | gi|120509127 | acetyl-CoA C-acyltransferase | *P. pachyrhizi* | 11 | 649 |
| PHAP0031  | gi|120522633 | deoxyuridine S’-triphosphate nucleotidohydrolase | *P. pachyrhizi* | 6 | 416 |
| PHAP0032  | gi|120523312 | dienelactone hydrolase | *P. pachyrhizi* | 10 | 243 |
| PHAP0043  | gi|120509948 | glycine dehydrogenase | *P. pachyrhizi* | 3 | 115 |
| PHAP0064  | gi|120519389 | isocitrate dehydrogenase | *P. pachyrhizi* | 5 | 102 |
| PHAP0065  | gi|120515452 | isocitrate lyase | *P. pachyrhizi* | 4 | 147 |
| PHAP0071  | gi|290897165 | NADP-dependent mannitol dehydrogenase | *M. larici-populina* | 3 | 83 |
| PHAP0103  | gi|00563667 | methylene-tetrahydrofolate dehydrogenase | *Leucosporidium scottii* | 3 | 89 |
| PHAP0109  | gi|254151530 | UTP-glucose-1-phosphate uridylyltransferase | *M. larici-populina* | 7 | 129 |
| **02. Energy** | | | | | |
| PHAP0027  | gi|120521145 | cytochrome C | *P. pachyrhizi* | 10 | 428 |
| PHAP0089  | gi|120515891 | pyruvate dehydrogenase beta | *P. pachyrhizi* | 12 | 339 |
| PHAP0101  | gi|120500717 | succinate-CoA ligase beta | *P. pachyrhizi* | 6 | 171 |
| PHAP0107  | gi|161646583 | ubiquinol-cytochrome C reductase | *Sporobolomyces roseus* | 6 | 78 |
| **10. Cell cycle and DNA processing** | | | | | |
| PHAP0024  | gi|209002725 | cell division cycle protein | *Melampsora medusae* | 8 | 214 |
| PHAP0044  | gi|118187570 | GTP binding protein/GTPinase | *Ustilago maydis* | 5 | 91 |
| PHAP0074  | gi|254164953 | nuclear segregation protein | *M. larici-populina* | 3 | 66 |
| PHAP0076  | gi|66651924 | nucleoside diphosphate kinase | *Uromyces viciae-fabae* | 5 | 173 |
| PHAP0096  | gi|120423194 | septin | *Uromyces appendiculatus* | 1 | 52 |
| **11. Transcription** | | | | | |
| PHAP0084  | gi|120503605 | polyadenylate binding protein | *P. pachyrhizi* | 9 | 379 |
| **12. Protein Synthesis** | | | | | |
| PHAP0117  | gi|120521780 | 40 S ribosomal protein S16 | *P. pachyrhizi* | 5 | 89 |
| PHAP0006  | gi|282815640 | 40 S ribosomal protein S19 | *Puccinia tricina* | 10 | 368 |
| PHAP0118  | gi|120498576 | 60 S acidic ribosomal protein | *P. pachyrhizi* | 5 | 88 |
| PHAP0009  | gi|120522205 | 60 S ribosomal protein L10 | *P. pachyrhizi* | 13 | 732 |
| **14. Protein Fate (folding, modification, destination)** | | | | | |
| PHAP0026  | gi|169738476 | cyclophilin | *U. appendiculatus* | 6 | 118 |
| PHAP0038  | gi|120515849 | glucose-regulated protein | *P. pachyrhizi* | 21 | 379 |
| PHAP0050  | gi|290908126 | hsp-10 protein | *Melampsora occidentalis* | 6 | 198 |
| PHAP0073  | gi|120425527 | neddylin/ubiquitin | *U. appendiculatus* | 3 | 129 |
Table 3 Putative proteins identified during *Phakopsora pachyrhizi* appressoria formation grouped into functional categories using the Munich Information Center of Protein Sequences (MIPS) Functional Catalogue (Continued)

| Accession | Description                        | Species          | Gene Number | E-Value | Description  |
|-----------|------------------------------------|------------------|-------------|---------|--------------|
| g|120523430 Proteasome subunit beta P. pachyrhizi 9 343 |
| g|120515648 Vacuolar protease A P. pachyrhizi 7 153 |
| 16. Structural & Protein Binding PHAP013 Actin lateral binding protein P. pachyrhizi 19 494 |
| g|156597317 Serine-threonine phosphatase Microbotryum violaceum 3 60 |
| g|120527960 Sterol binding protein P. pachyrhizi 11 113 |
| 20. Cellular transport, transport facilitation and transport routes PHAP01202 ADP, ATP carrier protein P. pachyrhizi 6 96 |
| g|120523715 Electron transport flavoprotein alpha P. pachyrhizi 5 200 |
| g|169735819 Mitochondrial import receptor TOM40 P. pachyrhizi 14 223 |
| g|254113321 Succinate dehydrogenase M. lanici-populina 8 302 |
| 32. Cell Rescue, Defense and Virulence PHAP0029 Cytochrome C peroxidase U. maydis 6 195 |
| g|120497372 Glutaredoxin-1 P. pachyrhizi 9 323 |
| g|120503688 Heat shock HSS1 P. pachyrhizi 7 376 |
| g|120509299 NADH-cytochrome b5 reductase P. pachyrhizi 5 321 |
| g|120514870 Polyubiquitin P. pachyrhizi 12 410 |
| 38. Transposable elements PHAP0057 Hypothetical protein U. maydis 3 65 |
| 99 Unclassified PHAP0030 Cytoplasmic protein P. pachyrhizi 5 313 |
| g|120521555 Hypothetical protein P. pachyrhizi 3 111 |
| g|120523258 Hypothetical protein P. pachyrhizi 2 191 |
| g|120510614 Hypothetical protein P. pachyrhizi 12 289 |
| g|120519941 Hypothetical protein P. pachyrhizi 15 512 |
| g|120534989 Hypothetical protein P. pachyrhizi 5 554 |
| g|103096425 Hypothetical protein U. maydis 3 65 |
| g|120521631 Hypothetical protein P. pachyrhizi 5 172 |
| g|120524713 Hypothetical protein P. pachyrhizi 5 141 |
| g|120506400 Hypothetical protein P. pachyrhizi 10 190 |
| g|120508009 Hypothetical protein P. pachyrhizi 3 119 |
| g|120518154 Hypothetical protein P. pachyrhizi 15 402 |
| g|120519874 Hypothetical protein P. pachyrhizi 4 98 |
| g|120519876 Hypothetical protein P. pachyrhizi 6 194 |

1Predicted *Phakopsora pachyrhizin* protein (PHAP).

2Proteins identified from the appressoria water fraction.

3Proteins containing a secretion signal.
Discussion

This study identified ESTs and proteins present during, and possibly required for, appressoria formation in *P. pachyrizi*. Spore germination, germ tube elongation, and appressoria formation are all part of a single complex physiological process from which individual steps cannot be readily uncoupled. Therefore, the collection of data specific to appressoria formation presents a technical challenge. Suppression subtractive hybridization (SSH) was utilized to generate a cDNA library enriched for transcripts specifically involved in appressoria formation, while reducing the occurrence of transcripts also associated with urediniospore germination and germ tube elongation. Generation of a cDNA library also allowed for the possible detection of transcripts that may be present in relatively low copy number. Alternatively, 2-DE and MS analysis of proteins extracted from appressoria offered a profile of high abundance proteins, which may not be reflective of corresponding transcript levels. Combining these two techniques provided a clearer image of the molecular and biochemical processes occurring during germination, germ tube elongation, and appressoria formation than either method viewed alone.
Comparison of the appressoria-enriched cDNA library to existing ESTs from germinated urediniospores of *P. pachyrhizi* revealed 29 non-redundant ESTs specific to the appressoria-enriched cDNA library. Ten of the 29 transcripts (35%) have roles in metabolism or cell cycling. This is to be expected given the importance of both autophagy and mitosis in appressoria formation. Autophagy is widely conserved among eukaryotes and is responsible for the degradation and recycling of proteins, organelles and cytoplasm in response to stress conditions that allow the cell to adapt to environmental or developmental changes [44]. In the absence of exogenous nutrients before host colonization, spore germination, germ tube elongation and appressoria formation must be fuelled by the breakdown of spore contents. Additionally, the metabolism of lipids, glycerol and sugars from the spore are utilized for the biosynthesis of glycerol in the appressorium, thereby generating the necessary turgor pressure to allow for penetration of the host cuticle. This breakdown and recycling of old cellular components has been shown to play a key role in pathogenicity of fungal pathogens, including *M. grisea* (*M. oryzae*), *Colletotrichum orbiculare*, and *Fusarium graminearum* [45-47]. The *P. pachyrhizi* appressoria-enriched ESTs (Table 2) include genes that encode proteins involved in fungal metabolism and autophagy, such as HMG-CoA reductase, a key enzyme for cholesterol biosynthesis, and glutamine synthetase, an important enzyme in amino acid metabolism. Expression analysis of HMG-CoA reductase indicates that this transcript is present in high levels in both germinated urediniospores and appressoria, but in relative low levels in urediniospores prior to germination (Figure 3, EST Pp3684). HMG-CoA reductase has also been shown to be up-regulated in appressoria of *M. grisea* [48]. Similarly, glutamine synthetase is highly expressed during pathogenesis of *Colletotrichum gloeosporioides* on *Stylosanthes guianensis* [49].

Cell cycling is pivotal for multicellular eukaryotes to coordinate the differentiation of tissues and organs. Appressoria formation requires switching from polarized hyphal growth, to expansion and cellular differentiation of appressoria, to ultimate resumption of polarized growth during penetration peg formation. Entry into mitosis, specifically the S phase, is required to regulate initiation of appressoria morphogenesis and conidial cell death in *M. grisea* [45]. Blockage of cell cycling at later stages of mitosis did not affect appressoria formation, suggesting that the checkpoint regulating cellular differentiation operates at the G2-M boundary [50]. A transcript for the G2/M phase checkpoint control protein SUM2 was identified among the *P. pachyrhizi* appressoria-enriched ESTs (Table 2). Interestingly, qRT-PCR analysis showed transcript levels in germinated urediniospores to be nearly twice that in appressoria. However, transcript levels in appressoria were more than four times that in urediniospores prior to germination (Figure 3, EST Pp3495).

Cyclic AMP-, MAP kinase-, and calcium/calmodulin-dependent signaling pathways are involved in the induction and development of appressoria [51-53]. In *P. pachyrhizi*, a putative serine/threonine protein kinase was identified and expressed in germinated urediniospores and appressoria (Figure 3, EST Pp3004). Serine/threonine kinases play a role in autophagy and fungal morphogenesis. Autophagy is blocked in mutants of the *MgATG1* gene in *M. grisea*, resulting in reduced lipid turnover, inadequate appressorial turgor, reduced ability to penetrate and infect a host, and decreased conidiation [54].

An EST from the *P. pachyrhizi* appressoria-enriched cDNA library was found with 98% similarity to autophagy-related protein 8 (Atg8) from *Moniliophthora perniciosa* (GenBank accession ACD93204) [55]. During autophagy, membrane-bound autophagosomes are formed, delivered to and fused with lysosomes or vacuoles where their contents are degraded. Atg8 is one of two ubiquitin-like proteins required for autophagosome formation [56]. Targeted mutation of Atg8 in *M. grisea* arrested conidial cell death via autophagy and prevented production of penetration hyphae, thus, preventing appressoria-mediated penetration of the host cuticle [45]. The role of Atg8 during germination and appressoria formation in *P. pachyrhizi* was supported by qRT-PCR analysis, which found increased transcript levels in germinated urediniospores and appressoria relative to urediniospores (Figure 4, EST Pp3944).

BLASTX analysis of appressoria-enriched ESTs revealed eight with similarity to hypothetical proteins of unknown function from *P. graminis f. sp. tritici* or *M. laricis-populina*. These two fungi, along with *P. pachyrhizi*, are members of the Order Puccinales. ESTs Pp3502 and contig2bb also showed similarity to other members of the Basidiomycota, while the other six ESTs may represent rust-specific transcripts. Gene expression analysis of EST Pp3505 revealed the highest transcript levels occur in appressoria relative to urediniospores and germinated urediniospores (Figure 3, EST Pp3505), supporting a role for this transcript in appressoria formation of *P. pachyrhizi*. BLASTX analysis of three additional ESTs, Pp3394, Pp3734, and Pp3842, did not find any significant similarity to any entries in the NBCI non-redundant protein database, suggesting that these transcripts represent genes specific to *P. pachyrhizi*.

Of the 238 non-redundant ESTs identified, 209 (88%) were found in common with previously sequenced ESTs from germinated urediniospores and infected soybean leaves. It is reasonable to expect that a portion of the
genes involved in the cascade of events required for appressoria morphogenesis may be triggered early during spore germination and germ tube elongation. Additionally, some of the genes required for appressoria formation may be the same genes necessary for spore germination and germ tube elongation. Identification of genes known to play a role in appressoria formation, penetration peg formation, and early infection in both the appressoria-enriched ESTs and the germinated urediniospore ESTs reinforces this expectation. Analysis of transcripts in other plant pathogenic fungi have shown greater changes in gene expression occur during spore germination, with fewer changes subsequently during germ tube elongation and appressoria formation [48,57].

A putative NADPH oxidase was identified among the ESTs common to both P. pachyrhizi cDNA libraries. NADPH oxidases generate reactive oxygen species (ROS) that are involved in various physiological processes and cellular differentiation in fungi [58]. Deletions of NADPH oxidase (Nox) genes block differentiation of sexual fruiting bodies in Aspergillus nidulans [59] and differentiation of ascogonia to perithecia in Podospora anserina [60]. In M. grisea Nox1 and Nox2 mutants formed normal looking appressoria but failed to penetrate the host, suggesting a role for Nox-derived ROS in penetration peg formation [61]. In P. pachyrhizi, qRT-PCR analysis revealed little to no expression of NOX in urediniospores, while expression in both germinated urediniospores and appressoria were equally high at 6 h, and reduced by more than half at 24 h. This profile matches the proposed role of ROS as a signalling pathway in cellular differentiation. Such signalling is unnecessary in dormant spores, but in high demand during germination and appressoria formation. It has been shown in M. grisea that the generation of ROS occurs during conidial germination, appressoria development, and during hyphal tip growth [61].

Another EST common to both P. pachyrhizi EST libraries, was a putative subtilase-type proteinase. The transcript levels of a subtilase-type proteinase in appressoria is nearly five times that observed in urediniospores, and the expression levels in germinated urediniospores is nearly six times that of urediniospores. Subtilisin-like serine proteases were abundant in both a cDNA library and a serial analysis of gene expression (SAGE) of appressoria of M. grisea [62,63]. Targeted deletion of the vacuolar serine protease, SPM1, resulted in decreased sporulation and appressoria development, and attenuated infection [64].

P-type ATPases are integral membrane proteins required for the maintenance of phospholipid asymmetry in biological membranes, and they are important for infection-related morphogenesis, such as penetration peg formation [65]. P-type ATPase ESTs were identified in both P. pachyrhizi cDNA libraries. The penetration defective mutant, PDE1, of M. grisea exhibited reduced appressoria-mediated penetration and reduced disease symptoms on a susceptible host [65]. They may also be significant for the delivery of virulence-associated proteins. A mutation of MgApt2 in M. grisea inhibited the hypersensitive response in resistant rice cultivars, suggesting that the secretion of fungal proteins perceived by the host during a resistance response might also require MgApt2 [66]. The qRT-PCR analysis of the P-type ATPase in P. pachyrhizi showed increased transcript levels in germinated urediniospores and appressoria, compared with low levels in urediniospores, suggesting a possible role in germination and appressoria formation.

Expression levels of genes analyzed by qRT-PCR found most transcripts to be present in urediniospores, usually at low levels relative to germinated urediniospores and appressoria (Figure 3 and Figure 4). The presence of most transcripts at low levels in dormant spores indicates that stabilized transcripts may be stored in the spore for translation during, or immediately following, germination [67]. Two P. pachyrhizi transcripts did not fit this profile. Transcript levels of EST Pp3205 (NADPH oxidase) in urediniospores were negative in two of three replicate runs. Contig6B had high transcript levels in urediniospores, with decreased levels in germinated urediniospores, and low levels in appressoria. This transcription pattern is consistent with contig6B’s putative identification as a conidiation-related protein 6 (CON6). The gene con-6 is expressed during conidiation in Neurospora crassa, but is not expressed in mycelium. It was shown that shortly after spore germination con-6 mRNA disappears and the CON6 polypeptide is rapidly degraded [68].

Of the 119 proteins identified from appressoria-enriched preparations, 59 (49.6%) proteins were not identified in the previous study of proteins from germinated urediniospores [15]. The differential protein profiles may be indicative of metabolic and physiological differences between germination on water versus a solid substrate. Differential expression was not as great between the two cDNA libraries as it was between the two proteomes, suggesting a possible greater influence of surface contact on translation than on transcription. As with the ESTs, the majority of proteins with an identified putative function play a role in metabolism. Additionally, proteins involved in cell cycling, protein fate, and cellular transport are well represented. Several of these, neddylin/ubiquitin, septin, GTP binding protein/GTPase are discussed below for their potential role in appressoria formation and early infection.

Isocitrate lyase was one of the proteins identified in this study, and is necessary for the utilization of fatty acids and is required for pathogenicity in Leptosphaeria
urediniospores and appressoria at 6 hpi relative to urediniospore penetration of the host cuticle [72]. Transcript necessary to differentiate appressoria and mediate mechanical support for generation of turgor pressure is critical for appressoria formation and likely function as an intermediate step through a covalent bond that links ubiquitin (an E1 enzyme) and ubiquitin-protein ligase in appressoria and germination, germ tube elongation, transcripts are accumulating without activation of subsequent translation. Alternatively, transcripts may be translated, but the protein turnover rate may be such that it precludes identification by 2-DE. The nature of the two techniques used may also explain some of the variation. Generation of a SSH cDNA library allows for the detection of transcripts that may be present in relatively low copy number, while 2-DE detects proteins of relative high abundance.

Fifteen proteins identified in the appressoria-enriched preparations were also found as ESTs in the appressoria-enriched cDNA library. All fifteen of these represented common, well-characterized proteins such as actin, catalase, α-tubulin, and aldehyde reductase, and all were also present in both the germinated urediniospore cDNA library and the partial proteome of germinated urediniospores. It is possible that during germination and germ tube elongation, transcripts are accumulating without activation of subsequent translation. Alternatively, transcripts may be translated, but the protein turnover rate may be such that it precludes identification by 2-DE. The nature of the two techniques used may also explain some of the variation. Generation of a SSH cDNA library allows for the detection of transcripts that may be present in relatively low copy number, while 2-DE detects proteins of relative high abundance.

While 59 proteins from the appressoria-enriched preparations did not share any sequence similarity with the 29 appressoria-enriched ESTs, commonality of function was identified. Neddylin/ubiquitin was identified among the proteins (Table 3, PHAP0073), while ubiquitin-protein ligase was found among the ESTs (Table 2, EST Pp3282). Both of these proteins are involved in the ubiquination of proteins targeted for degradation in the proteasome. In conjunction with E1 (ubiquitin-activating enzyme) and E2 (ubiquitin-conjugating enzyme), neddylin acts as an intermediate step through a covalent bond with the target protein and forms a bridge between E2 and E3 (ubiquitin-protein ligase) [42]. Expression levels of the ubiquitin-protein ligase in appressoria and germinated urediniospores were more than double that of urediniospores (Figure 3, EST Pp3282).

A GTP binding protein (GTPase) was identified among the proteins (Table 3, PHAP0044) and ESTs (Table 1, EST Pp3042). Transcript analysis by qRT-PCR indicates that EST Pp3042 was highest during urediniospore germination and early germ tube elongation (Figure 4). GTP binding proteins have been shown to affect the formation of septa in infectious hyphae of U. maydis [72]. Septa are critical for appressoria formation and likely function as mechanical support for generation of turgor pressure necessary to differentiate appressoria and mediate mechanical penetration of the host cuticle [72]. Transcript levels of a putative septin were elevated in germinated urediniospores and appressoria at 6 hpi relative to urediniospores (Figure 4, EST Pp3222). Septin was also identified among the appressoria proteins (Table 3). Septins are conserved cytoskeletal GTPases with multiple functions, including organizational markers during cell division and polarized growth. In filamentous fungi, septins assemble into a wide variety of complexes, including those that form at growing hyphal tips and at the site of future septum formation. In M. oryzae deposition of the septin ring defines the position of the appressorium septum prior to mitosis in the germ tube, and septin ring formation appears to be regulated by the DNA replication checkpoint that initiates appressorium morphogenesis [73]. In U. maydis septin mutants have reduced symptom development on maize and produce fewer mature teliospores than wild-type [74], which is consistent with the requirement for septin during morphogenesis and cellular division.

A putative acyl-CoA dehydrogenase EST was found in both germinated urediniospore and appressoria cDNA libraries (Figure 4, EST contig2S), and acetyl-CoA acyltransferase was identified among the high abundance proteins (Table 3). These two enzymes are involved in the first and last steps of fatty acid β-oxidation, which is essential for the development and melanization of appressoria in M. grisea [75].

Five non-redundant EST contigs were identified with partial homology to one another and shared similarity to MAS3 of M. grisea and gEgh16 of Blumeria graminis f. sp. hordei. MAS3 and gEgh16 are both members of multigene families and are expressed during early infection [39,76,77]. Similarly, the five contigs may comprise a multigene family in P. pachyrhizi. Expression analysis of the most redundant transcript, contig481, revealed that this putative MAS3 homolog was highly expressed in germinated urediniospores and present in urediniospores at levels similar to those in appressoria (Figure 4, EST contig481).

Prior to removing redundancy, nearly half (49.6%) of the clones sequenced from the appressoria-enriched cDNA library showed similarity to MAS3 of M. grisea and gEgh16 of Blumeria graminis f. sp. hordei. MAS3 and gEgh16 are both members of multigene families and are expressed during early infection [39,76,77]. Similarly, the five contigs may comprise a multigene family in P. pachyrhizi. Expression analysis of the most redundant transcript, contig481, revealed that this putative MAS3 homolog was highly expressed in germinated urediniospores and present in urediniospores at levels similar to those in appressoria (Figure 4, EST contig481).

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While transcripts are present, it is not until the infection process reaches the stage of penetration peg formation or actual penetration that MAS3 plays a role during infection. There may be a signal from the mature appressoria or the host plant that triggers translation at the critical point between appressoria formation and penetration.

Eight putative extracellular proteins were identified in AEF and AWF (Table 3). Three of these are well-characterized intracellular proteins that share similarity to proteins in other fungi: glucose-regulated protein, neddylin/ubiquitin, and vacuolar protease A. Four proteins share similarity to hypothetical proteins found in other fungi. The remaining putative extracellular protein, PHAP0055, did not have any similarity to proteins in the NCBI non-redundant protein database, suggesting it is unique to *P. pachyrhizi*.

The *P. pachyrhizi* protein PHAP0059 contains a conserved fasciclin domain which has been demonstrated to be involved in cell adhesion, appressoria turgor, and pathogenicity in *M. oryzae* [78,79]. Interestingly PHAP0059 was only found in the AEF and not among the germinated urediniospore proteins. The AEF was generated on a hard surface, while urediniospores were germinated by floating on the surface of water. This suggests that contact with a hard surface may be necessary to induce signals that regulate transcription or protein processing for translation of cell adhesion proteins.

Two *P. pachyrhizi* proteins, PHAP0055 and PHAP0129, exhibit characteristics similar to those of putative effector proteins identified from a haustorium-enriched proteome of *P. triticina* [43]. Both are small molecular weight proteins, with no known function or conserved domains, and contain a predicted signal peptide cleavage site at the N-terminus. PHAP0129 also contains an even number of cysteine residues which is characteristic of some plant-targeted fungal effector proteins [21]. In addition, PHAP0129 was found only in the AEF, further supporting its putative identification as an extracellular protein. Proteins with high identity to PHAP0129 were found in *P. pachyrhizi*, *U. viciae-fabae* and *P. triticina*, suggesting a putative effector protein family exists in rusts.

**Conclusions**

This study utilized SSH and 2-DE and MALDI-TOF/TOF to identify genes and proteins expressed during appressoria formation. Genes and proteins involved in early infection by *P. pachyrhizi* are candidates for targeted control measures. Several genes and proteins were identified as unique to *P. pachyrhizi* and/or other rusts. One of these unique proteins is a putative extracellular effector protein. A potential rust-specific effector protein family was also identified. Determining the role of these effector proteins will be an important step in understanding the mechanisms of pathogenesis.

**Additional files**

**Additional file 1**: Table S1. Proteins identified using the custom EST database.

**Additional file 2**: Table S2. Non-redundant *Phakopsora pachyrhizi* clones identified from the appressoria-enriched cDNA library.

**Additional file 3**: Table S3. Non-redundant proteins identified using the custom EST database.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

CLS participated in the design and execution of the experiments. She performed the soybean inoculations, developed the method for generating appressoria on polystyrene plates, extracted RNA, analyzed ESTs, performed qRT-PCR, and drafted the manuscript. MBM conducted the protein extractions and 2-DE, analyzed the MALDI-TOF/TOF data, and assisted in preparing the manuscript. LLF constructed the custom EST database, conducted the MALDI-TOF/TOF, and assisted in the proteomic analysis. AN oversaw the MALDI-TOF/TOF and assisted in the proteomic analysis. GWS developed computer algorithms for EST analysis. DGL and RDF conceived of the project, wrote the plan for implementation, oversaw the execution of the experiments and data analysis, and revised and edited the manuscript. All authors read and approved the final manuscript.

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