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Abstract

Rust fungi are devastating pathogens for several important crop plants. The biotrophic lifestyle of rust fungi requires that they influence their host plants to create a favorable environment for growth and reproduction. Rust fungi secrete a variety of effector proteins that manipulate host target proteins to alter plant metabolism and suppress defense responses. Because of the obligate biotrophic lifestyle of rust fungi, direct evidence for effector function is difficult to obtain, and so suites of experiments utilizing expression in heterologous systems are necessary. Here, we present results from a yeast cell death suppression assay and assays for suppression of PAMP-triggered immunity (PTI) and effector triggered immunity (ETI) based on delivery of effectors through the bacterial type III secretion system. In addition, subcellular localization was tested using transient expression of GFP fusion proteins in Nicotiana benthamiana through Agrobacterium infiltration. We tested 31 representative effector candidates from the devastating common bean rust pathogen Uromyces appendiculatus. These effector candidates were selected based on features of their gene families, most important lineage specificity. We show that several of our effector candidates suppress plant defense. Some of them also belong to families of effector candidates that are present in multiple rust species where their homologs probably also have effector functions. In our analysis of candidate effector mRNA expression, some of those effector candidates that gave positive results in the other assays were not up-regulated during plant infection, indicating that either these proteins have functions at multiple life stages or that strong up-regulation of RNA level in planta may not be as important a criterion for identifying effectors as previously thought. Overall, our pipeline for selecting effector candidates based on sequence features followed by screening assays using heterologous expression systems was successful in discriminating effector candidates. This work lays the foundation for functional characterization of U. appendiculatus effectors, the identification of effector targets, and identification of novel sources for resistance in common bean.

Keywords

Uromyces appendiculatus, effectors, PTI suppression, ETI suppression, bax cell death suppression, localization, expression patterns

Disciplines

Agricultural Science | Agriculture | Plant Breeding and Genetics | Plant Pathology

Comments


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Authors
Candidate Effectors From *Uromyces appendiculatus*, the Causal Agent of Rust on Common Bean, Can Be Discriminated Based on Suppression of Immune Responses

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Rust fungi are devastating pathogens for several important crop plants. The biotrophic lifestyle of rust fungi requires that they influence their host plants to create a favorable environment for growth and reproduction. Rust fungi secrete a variety of effector proteins that manipulate host target proteins to alter plant metabolism and suppress defense responses. Because of the obligate biotrophic lifestyle of rust fungi, direct evidence for effector function is difficult to obtain, and so suites of experiments utilizing expression in heterologous systems are necessary. Here, we present results from a yeast cell death suppression assay and assays for suppression of PAMP-triggered immunity (PTI) and effector triggered immunity (ETI) based on delivery of effectors through the bacterial type III secretion system. In addition, subcellular localization was tested using transient expression of GFP fusion proteins in *Nicotiana benthamiana* through *Agrobacterium* infiltration. We tested 31 representative effector candidates from the devastating common bean rust pathogen *Uromyces appendiculatus*. These effector candidates were selected based on features of their gene families, most important lineage specificity. We show that several of our effector candidates suppress plant defense. Some of them also belong to families of effector candidates that are present in multiple rust species where their homologs probably also have effector functions. In our analysis of candidate effector mRNA expression, some of those effector candidates that gave positive results in the other assays were not up-regulated during plant infection, indicating that either these proteins have functions at multiple life stages or that strong up-regulation of RNA level *in planta* may not be as important a criterion for identifying effectors as previously thought. Overall, our pipeline for selecting effector candidates based on sequence features followed by screening assays using heterologous expression systems was successful in discriminating effector candidates. This work lays the foundation for functional characterization of *U. appendiculatus* effectors, the identification of effector targets, and identification of novel sources for resistance in common bean.

**Keywords:** *Uromyces appendiculatus*, effectors, PTI suppression, ETI suppression, bax cell death suppression, localization, expression patterns
INTRODUCTION

Fungal pathogens use effectors to influence their host plants. These effectors may function in weakening or killing the infected tissue as is the case for necrotrophic pathogens. Alternatively, effectors produced by biotrophic pathogens may influence the host into transferring nutrients or forming structures to accommodate the pathogen. Perhaps, the most important among the effectors are those that suppress plant defense responses. Suppression of plant defense is especially important for biotrophic pathogens since they can be effectively killed by hypersensitive plant cell death responses.

Recently, we described a comprehensive pipeline of assays in heterologous systems that aimed to identify true effectors from collections of candidate effectors (Qi et al., 2018). This pipeline is centered on identifying candidate effectors that suppress plant defense responses. Due to their biotrophic lifestyle, it is still impossible to genetically transform rust fungi with a single exception that is based on a cloned avirulence gene (Lawrence et al., 2010), which makes it necessary to utilize heterologous organisms to test for defense suppression. The bacterial type III secretion system (T3SS), which has the natural purpose of transferring effector proteins into host cells, is a well-established approach for testing the ability of fungal effectors to suppress defense responses (Sohn et al., 2007). To induce transfer of a fungal protein via the T3SS, a plasmid is introduced that fuses the T3SS signal peptide to the N-terminus of the fungal candidate effector replacing its own signal peptide. In our assays, we used *Pseudomonas syringae pv. tomato* strain DC3000 (*Pst* DC3000) or *Pseudomonas fluorescens* effector to host analysis (EtHAn) (a non-pathogen with artificially added T3SS) (Thomas et al., 2009) to deliver candidate effectors into *Nicotiana benthamiana*. *Pst* DC3000 causes HR when it is infiltrated to *N. benthamiana*—except if it carries an effector that can suppress HR. On the other hand, *P. fluorescens* triggers basal defense. Evidence for the basal defense reaction is provided by infiltration with *Pst* DC3000 several hours later; where basal defense is already established, *Pst* DC3000 cannot trigger a subsequent HR (Oh and Collmer, 2005). Therefore, if leaf tissue, which was first infiltrated with *P. fluorescens* EtHAn carrying an effector candidate and later with *Pst* DC3000, shows HR, this can be considered evidence that the effector can suppress basal defense.

Because suppression of HR can be connected to suppression of cell death, we complemented our defense suppression assays with a cell death suppression assay in yeast. For this, we used a yeast system with galactose inducible BAX-induced cell death. If yeast cells expressing a candidate effector grow well on a medium containing galactose the effector suppresses cell death in this assay. Subcellular localization can also aid at effector function (Petre et al., 2015; Petre et al., 2016). We used agroinfiltration to transiently express GFP fusion proteins of our effector candidates in *N. benthamiana*. Another indicator of effector function is expression of the effector candidate that coincides with the potential function. For those candidate effectors that yielded interesting results in the screens described above, we also elucidated the expression pattern throughout the infection process starting with the urediospore.

In previous work (Link et al., 2014; Qi et al., 2016; Qi et al., 2018), we aimed to identify effectors of *Phakopsora pachyrhizi*, which causes Asian soybean rust. Here, we used a similar pipeline to assess a collection of effector candidates from *Uromyces appendiculatus*. The effector candidates were selected from haustorial-expressed proteins that were predicted to encode a signal peptide (Link et al., 2014). *U. appendiculatus* is the causal agent of common bean rust, which is one of the most devastating diseases of common bean (*Phaseolus vulgaris*). Common bean rust leads to severe yield losses through reduced photosynthesis in the infected leaves and the carbohydrate sink formed by the fungus resulting in reduced numbers of pods, reduced pod fill, and therefore, fewer seeds (Liebenberg and Pretorius, 2010). Common bean is an important protein source for millions of people in Mesoamerica and Africa where it is cultivated in small plots for self-supply. Although resistance genes are available to be deployed against *U. appendiculatus* (Souza et al., 2008), gene-for-gene resistance has often been overcome by rust pathogens, so it cannot be considered as durable. Novel approaches to achieve resistance with improved durability may revolve around effectors. For example, the expression of effectors could be silenced using host-induced gene silencing (HIGS), effector targets in the host plant could be deleted or modified to render effectors ineffective or artificial resistance proteins interacting with the effectors could be introduced into plants. In our study on haustorial transcriptomes (Link et al., 2014), we explored *P. pachyrhizi* and *U. appendiculatus* together, expecting to find differences and commonalities in the effector repertoire. The same expectation was yet another reason for us to continue research on the *U. appendiculatus* effector candidates. In the meantime, additional information on potential *U. appendiculatus* effectors has been obtained using proteomics and HIGS approaches (Cooper et al., 2016; Cooper and Campbell, 2017). Some of these candidates were investigated by us as well, and where possible, a comparison of results is presented.

MATERIALS AND METHODS

Bacterial Strains and Plasmids Mobilizations

Bacterial strains used in this study are listed in Table 1. *E. coli* and *Agrobacterium tumefaciens* were grown in Lysogeny broth (LB). *Pseudomonas* strains were grown in King's B (KB) medium at 28°C. Standard triparental mating with *E. coli* HB101 (pRK2013) as a helper strain was used to mobilize plasmids from *E. coli* to *Pseudomonas* strains.

Plant Material

*P. vulgaris* plants (cultivar “Primel”) were grown in greenhouse chambers with 16-h light/8-h dark and 22°C/18°C. *N. benthamiana* plants were grown in a growth room at an average temperature of 24°C (range, 20°C–26°C), with 45% to 65% relative humidity under long-day conditions (16 h light).
**Candidate Effectors From** *Uromyces appendiculatus*

**Fungal Isolate, in vitro Structures of** *U. appendiculatus*, and Inoculation of *P. vulgaris*

Urediospores of *U. appendiculatus* (SWBRI, laboratory collection, Phytopathology, University of Hohenheim, Stuttgart, Germany) were finely dispersed on polyethylene (PE) sheets using fine gauze material for sifting. The PE sheets were then sprayed with water using a chromatography vaporizer and afterward kept at 20°C, 95% humidity in the dark. For germ tubes (gt), the structures were harvested from the PE sheets after 4 h and for appressoria (ap) after 9 h. Also, for gt, smooth PE sheets were used, whereas for ap formation, the PE sheets were scratched using a brass brush. Formation of both gt and ap was checked microscopically. The structures were scraped from the PE sheets, dried by vacuum filtration, and stored in 2-ml tubes at −70°C after freezing in liquid nitrogen. For plant inoculation, a watery suspension with 0.05% spores, 0.08% milk powder, and 0.01% Tween20 (w/v) was produced and stirred for 30 min at room temperature in the dark. The suspension was then sprayed on 21 days old *P. vulgaris* plants using a chromatography vaporizer. The inoculated plants were kept at 95% humidity in the dark at 15°C for 24 h before putting them back to greenhouse conditions with 16-h light/8-h dark and 15°C/13°C.

**RNA Preparation and cDNA Synthesis**

*U. appendiculatus in vitro* structures were harvested as described above. Pieces from leaves of infected plants at different stages and uninfected plants were cut out with a cork borer (12 mm diameter, four pieces from different leaves per sample, roughly 100 mg), put into plastic tubes, frozen in liquid nitrogen, and kept at −70°C until RNA preparation. Frozen samples were homogenized using FastPrep®-24 MP (Biomedicals GmbH, Eschwege, Germany) at 4.5 m/s for three times 20 s with two 2-mm steel beads and with cooling the samples in liquid nitrogen between homogenization rounds. RNA purification was done using the Agilent Plant RNA Isolation Mini Kit (Agilent Technologies, Santa Clara, CA, USA) with adding an additional centrifugation step (2 min, 16,000 rpm, room temperature) between adding the extraction buffer to the homogenized tissue and using the mini prefiltration column. RNA concentration, quality, and integrity were checked measuring OD260/280 and running 1% denaturing agarose gels. Storage of the isolated RNA was at −70°C.

Before reverse transcription, RNA was treated with DNasel (Thermo Fischer Scientific, Schwerte, Germany) using 1-µg RNA in a 10-µl reaction. For cDNA synthesis, the RevertAid RT Reverse Transcription Kit (Thermo Fischer Scientific) was used. Twenty-microliter reactions starting with the DNasel-treated RNA from above were performed following instructions from the manufacturer and using random hexamer primers. cDNA was stored at −20°C.

**Cloning**

Inserts for cloning were produced by PCR on cDNA from urediospore RNA using nested PCR. First-round PCR was in 10-µl reactions with 1 µl 10× buffer, 1 µl 10 mM dNTP mix, 0.5 µl 0.1 µM f1 primers each (see Table S1), 1-µl cDNA, and 0.1 µl Long PCR Enzyme Mix (Thermo Fischer Scientific). Second round PCR was in 50-µl reactions with 5 µl 10× buffer, 5 µl 10 mM dNTP mix, 2.5 µl f2 primers each, 2 µl PCR product from the first round reaction, and 0.5 µl Long PCR Enzyme Mix. Cycling conditions were the same for both PCRs: initial denaturation 3 min 94°C, then 35 cycles with 20 s 95°C, 30 s 55°C, and 60 s 68°C; final elongation 7 min 68°C. Second round PCR products were purified using the peqGOLD Cycle-Pure Kit (PEQLAB Biotechnologie GmbH, Erlangen, Germany) and inserted into the pCR™8/GW/TOPO® plasmid using the pCR™8/GW/TOPO TA Cloning® Kit (Invitrogen, Life Sciences, Carlsbad, CA, USA), following the instructions of the manufacturer and using 4 µl of the purified PCR product. All constructs were verified by sequencing. For production of constructs without the signal peptide, the inserts were PCR amplified again using a third set of primers and again inserted into the pCR™8/GW/TOPO® plasmid. All three sets of primers are supplied in Table S1.

<table>
<thead>
<tr>
<th>Strain or plasmid</th>
<th>Genotype or relevant phenotype</th>
<th>Source or reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>E. coli DH5α</td>
<td>F- endA1 glnV44 thi-1 recA1 relA1 gyrA96 deoR supE44 lacZΔM15 lacYΔF75 endA1-lacZYA-argF-U169 hisD2ΔR17trp1-Δ629 leu2-3,112thi1-1  rpsL15 indB  proA937  lacI977lacZ15 lacYΔ7 ΔproA</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>TOP10</td>
<td>F- mcrA Δmrr-hsdRMS-mcrBC 80lacZAM15 lacX74 80lacZM15mcrB1 80lacZAM15mcrB1 80lacZM15mcrB1</td>
<td>Invitrogen</td>
</tr>
<tr>
<td><em>P. syringae pv. tomato</em> DC3000</td>
<td>Wild type, Rif®</td>
<td>(Foine et al., 1997)</td>
</tr>
<tr>
<td><em>P. fluorescens</em> EiHAn</td>
<td>P. fluorescens Pf0-1 carrying a working TTSS transfer machinery, Rif®, Gm®</td>
<td>(Thomas et al., 2009)</td>
</tr>
<tr>
<td>Agrobacterium tumefaciens GV3101</td>
<td>Carries Vir plasmid encoding T-DNA transfer machinery, Rif®, Gm®</td>
<td>(Koncz and Schell, 1986)</td>
</tr>
<tr>
<td><em>Saccharomyces cerevisiae</em> BF264-15Dau</td>
<td>MATα ade1 his2 leu2-3,112trp1-1α Ura3</td>
<td>Melissa G. Mitchum Lab</td>
</tr>
<tr>
<td>Plasmids</td>
<td>Gateway-compatible entry vector, Spr</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>pCR™8/GW/TOPO®</td>
<td>Gateway-compatible version of pEDV3, Gmr</td>
<td>(Sohn et al., 2007)</td>
</tr>
<tr>
<td>pEDV6</td>
<td>Gateway-compatible version of pGBKT7, Kmr</td>
<td>Melissa G. Mitchum Lab</td>
</tr>
<tr>
<td>pGBKT7-GW</td>
<td>Gateway-compatible version of pGBKT7, Kmr</td>
<td>Melissa G. Mitchum Lab</td>
</tr>
<tr>
<td>pYEps1-bax</td>
<td>Expression controlled by GAL7 promoter, Amp, LEU2</td>
<td>Martin et al., 2009</td>
</tr>
<tr>
<td>pSITEII-3C1</td>
<td>Gateway-compatible binary vector for transiently over-expression of EGFP-fused protein in planta, Sp</td>
<td></td>
</tr>
</tbody>
</table>

*Antibiotics were used at the following concentrations (µg/ml): Rifampicin (Rif) 100, Gentamycin (Gn) 50, Kanamycin (Km) 100, Spectinomycin (Sp) 50, Chloramphenicol (Cm) 30, and Ampicillin (Amp) 100.*
**Bacterial Inoculation and Growth in planta**

*N. benthamiana* plants used in this study were between 5 and 6 weeks old. All plant assays were performed by infiltrating a bacterial suspension into plant leaves with a needleless syringe. *Agrobacterium* strains were re-suspended in the induction buffer (100 μM acetosyringone, 10 mM MES, pH 5.6, and 10 mM MgCl₂) and kept at room temperature for 3 h before infiltration, all other strains were re-suspended in inoculation buffer (10 mM MgCl₂). The areas of bacterial infiltration were marked lightly with a Sharpie® permanent marker. Levels of bacterial inoculum used in experiments are noted in the figures and legends.

**Agroinfiltration-Mediated Transient Expression in N. benthamiana**

To create GFP-Uaca_N constructs, the Uaca_N was PCR amplified and cloned into the Gateway entry vector pCR®-8GW/TOPO® (Invitrogen) and then recombined into the Gateway binary destination vector pSITEII-3C1 (Table 1). The resulting binary constructs were introduced into *A. tumefaciens* GV3101 using the freeze-thaw method (An et al., 1988). GFP was detected in agroinfiltrated *N. benthamiana* leaves using a Zeiss Axio Imager epi-fluorescence microscope at 48 h after infiltration.

**Yeast Cell Death Assays**

Yeast cell death assays were performed following the procedure previously described (Laloux et al., 2010) with some modifications. Yeast strain BF264-15Dau carrying the plasmid Yep51-bax, which is inducible by galactose to express BAX protein, and Gateway-compatible pGBK7-GW containing Uaca_N, was cultured in liquid SD/-Leu/-Trp media at 30°C for overnight before the yeast cells were resuspended in liquid YNB/Gal/-Leu/-Trp medium (Yeast Nitrogen Base 6.7 g/L, galactose 2%, -Leu/-Trp DO Supplement 0.64 g/L) for six-hour induction. Yeast cells were collected by centrifugation, washed in liquid YNB/Gal/-Leu/-Trp medium for at least four times, and adjusted to OD₆₀₀ nm = 1. The resulting yeast inoculum was serially diluted and placed on YNB/Gal/-Leu/-Trp plates. The plates were photographed at the fourth day after culturing at 30°C. Nematode effector 4P01, which consistently and strongly suppresses BAX cell death [Melissa G. Mitchum, personal communication and Qi et al. (2018)], and empty vector were included in each plate as positive and negative control, respectively.

**RT-qPCR Measurements**

Real-time PCR reactions were run on a CFX96™ Real-Time-PCR System (Bio-Rad Laboratories, Hercules, CA, USA). The SensiFAST™ SYBR® No-ROX mix (Bioline Reagents Ltd., London, UK) was used with primers at 0.4-μM final concentrations and 2-μl cDNA in 20-μl reactions. We used a two-step PCR protocol with an initial denaturation step of 95°C for 5 min, and then 5 s at 95°C, and 15 s at 60°C. Forty cycles were completed; then a melt curve analysis was run spanning 65 to 95°C with 0.5°C temperature increase per 5 s. Data logging and the determination of Cq values were done using Bio-Rad CFX Manager 2.1 (Bio-Rad Laboratories). We used a threshold of 2,500 RFU, the estimated mean of thresholds set by automatic threshold determination. Means of the two technical replicates were calculated using the GenEx software package: GenEx 6.0.1.612 (MultiD Analyses AB, Göteborg, Sweden). Using GenEx, we also performed efficiency correction, normalization to the two reference genes, *Act* and *CytB* [determined as stable comparing several candidate reference genes; publication in preparation; strategy similar to the one described in Hirschburger et al. (2015) including all stages also used here] and calculated the expression relative to the gt sample. The geometric means of the three biological replicates were calculated in MS Excel.

We used Primer3 (Koressaar and Remm, 2007; Untergasser et al., 2012) for primer design and GeneRunner for additional checking and optimizing the primers. Primers were chosen to obtain amplicons between 150 and 250 bp in length. All primer pairs were tested for efficiency using dilution series with four tenfold dilutions prepared from cDNA of gt or 7 dpi stages. Primer data are summarized in Table S2.

**RESULTS**

**Pre-Selection of Candidate Effectors**

Our study on the haustorial transcriptomes of *P. pachyrhizi* and *U. appendiculatus* resulted in a much higher number of predicted secreted proteins for *U. appendiculatus* (395) than for *P. pachyrhizzi* (156) (Link et al., 2014). This prompted us to use a different cloning strategy. Instead of aiming at cloning all and starting with the largest, we focused on features that were in part already described in Link et al. (2014) to select among the predicted secreted proteins with the goal of cloning 40 to 50 secreted proteins. Faced with a similar problem, Petre et al. (2015) used the following criteria to select candidate effectors: haustorial expression, in planta-specific expression, no protein domain, Pucciniales specific, positive selection, induction during biotrophic growth, and homology to rust effectors. Due to the absence of genome sequence and time course transcriptome data, we chose specificity to Pucciniales as our primary criterion to select candidates. We preferentially selected candidates with homology to known rust effectors and excluded those with annotated functional domains. These criteria were applied to the protein families resulting from clustering analysis performed by Link et al. (2014) rather than to the single proteins. We then chose representative proteins from these families as candidates to be cloned based on whether they were the only *U. appendiculatus* protein in the family or the *U. appendiculatus* protein that fit the Hidden Markov model to the family best. We expected that this family-wise approach would provide us with a manageable number of candidates, and the ability to make general conclusions based on the protein representing each family. The *U. appendiculatus* candidates were termed Uaca_N, with the number N, indicating the order in which the candidates were selected and cloned.

For each Uaca_N, we first cloned the full ORF into the pCR®8 Gateway entry vector, and then we also produced a corresponding clone in pCR®8 minus the coding sequence.
for the predicted signal peptides, because our screens were dependent on intracellular expression or expression from the bacterial T3SS. The latter set of clones with no signal peptide were subsequently designated Uaca_N_s. The Uaca_N were cloned from cDNA derived from RNA that was extracted from U. appendiculatus SWBR1 urediospores. Because single-step PCR on this kind of cDNA proved unsatisfactory in earlier experiments, we used nested PCR for all Uaca_N. The resulting pCr™8 plasmids were tested for inserts by colony PCR and sequenced. The Uaca_N were cloned from the pCr™8 plasmids containing the Uaca_N. All primers used for the cloning procedure are presented in Table S1. Only 31 of the 48 selected Uaca_N were successfully cloned and used in subsequent assays for cell death suppression in yeast and plants, PTI suppression, localization, and time course expression analysis. The description of the gene families and rationale for selection is provided in Table 2 along with a summary of the results from each of the assays. Information, including the contig names that can be found in public databases, cloned ORF sequence, the protein sequence, protein length, cysteine content, and the predicted signal peptide that was removed is provided in Data Sheet 1.

**Suppression of HR and Basal Defense by Uaca_Ns**

The 31 Uaca_N were transferred into the bacterial T3SS vector pEDV6 (effector detector vector 6), which previously has been used to deliver non-bacterial proteins into plants (Fabro et al., 2011; Upadhyaya et al., 2014; Qi et al., 2018). These 31 plasmids plus the empty pEDV6 were introduced into Pst DC3000 and P. fluorescens strain EtHAn (a non-pathogen engineered to carry the T3SS) (Thomas et al., 2009). Pst DC3000 induces HR in N. benthamiana due to recognition of the effector HopQ1-1 (Wei et al., 2007), which provides a straightforward system for screening Uaca_N for the ability to suppress ETI when expressed from pEDV6 (Qi et al., 2016; Qi et al., 2018). To test for suppression of HR, Pst DC3000 strains carrying pEDV6 expressing the 31 Uaca_N were blunt syringe-inoculated into leaves of N. benthamiana. Among the 31 strains, Uaca_9ns, Uaca_12ns, Uaca_14ns, and Uaca_22ns were able to reproducibly suppress the HR caused by Pst DC3000 to different extents (Figure 1). The other 27 strains were able to cause HR in N. benthamiana leaves that was comparable to the pEDV6 empty vector control. These data demonstrate that four Uaca_Ns can suppress ETI induced by HopQ1-1 in N. benthamiana. To test for suppression of basal defense, we used the assay established by Oh and Collmer (2005). The principle of this test is that non-pathogenic P. fluorescens EtHAn induces basal defense in N. benthamiana. Pst DC3000, on the other hand, triggers hypersensitive cell death. When Pst DC3000 is infiltrated into tissue that previously was infiltrated with EtHAn and, therefore, exhibits basal defense, no hypersensitive response is observed. The assay was successfully used to screen for Phakopsora pachyrhizi effectors (Qi et al., 2018). EtHAn strains carrying the 31 Uaca_Ns, together with empty vector control, were blunt syringe-inoculated into leaves of N. benthamiana, and 7 h later overlapping sites were inoculated with Pst DC3000. Of the 31 Uaca_Ns, six (Uaca_4, Uaca_5, Uaca_7, Uaca_9, Uaca_28, and Uaca_44) enabled Pst DC3000-induced HR, indicating that these six effector candidates are able to suppress the basal defense induced by EtHAn (Figure 2).

**Suppression of BAX Induced Cell Death in Yeast by Uaca_Ns**

All 31 Uaca_Ns were transferred to plasmid pGBKKT7-GW, and the resulting constructs were transformed into yeast strain BF264-15Dau, which carries the plasmid YEp51-bax BF264-15Dau that confers bax gene expression under galactose induction (Zha et al., 1996). Bax induces cell death through effects like production of reactive oxygen species. We found that 25 of the 31 Uaca_Ns could suppress BAX cell death to various extents, and we categorized the suppression as strong, medium, weak, and negative (see Figure 3 for representative examples). Into these categories fell 12, 10, 3, and 6 proteins, respectively. We also checked for tendencies using the same plots as for HR and PTI. Here, again, a tendency toward high cysteine content but no clear groupings could be seen in our plots.

**Subcellular Localization of Uaca_Ns**

To determine the subcellular localization of Uaca_Ns in planta, 31 Uaca_Ns were cloned into pSITEII-3C1 (Martin et al., 2009) to fuse the GFP reporter to their amino termini. The GFP-Uaca_N fusion proteins were transiently expressed in N. benthamiana by agroinfiltration. GFP fluorescence was visualized by epifluorescence microscopy in N. benthamiana epidermal cells. There were three basic patterns of GFP distribution in the cells: cytoplasmic, nuclear, and cytoplasmic + nuclear (Figure 4, Supplementary Figure 2, and Table 2). Of the 31 GFP-Uaca_N fusion proteins, 18 were distributed in both the nucleus and the cytoplasm (e.g. Figure 4B). Another 12 were observed only in the cytoplasm and excluded from the nucleus (e.g. Figure 4D), 2 of which appear to be located mainly in small bodies, which could be organelles, other subcellular compartments, or artifacts of protein aggregation (Figure 4C). Only one, GFP-Uaca_1, was mostly located in the nucleus (Figure 4A). These findings indicate that U. appendiculatus, similar to other rust fungi, may deliver effector proteins to a variety of distinct host cell compartments (Petre et al., 2015).
Cluster 112 has most members \[1\] and contains

- Strong
- Medium
- N+C

ΔBAX

Cluster 2768 is lineage specific to Pucciniaceae, has six conserved cysteines and

-−−−

Subcellular

Cluster 874 has only few homologs per species, seven or three conserved cysteines

Cluster 2565 is lineage specific to Pucciniaceae, maybe even Uromyces. Uaca_16 is

C

ΔII

Cluster 112 has the most members \[1\] and contains

−

Cluster 824 is not very conserved; N-terminus differs strongly but conserved cysteines

−

Cluster 1_0_152 is mostly lineage specific for

Cluster 1_63 is specific to rust fungi. Uaca_25 has the longest sequence of the family.

+++

Negative

0

−

Cluster 2456, lineage specific to Pucciniaceae \[1\]. Fits the pattern best.

C, aggregation

Medium

Cluster 1293, only one homolog per species.

Strong

C

Cluster 398 \[1\]. Proteins in the cluster have 12 conserved cysteines and some

−

Strong

N+C

C

Cluster 2917, basically just one gene with one homolog in

C

Cluster 145 has three conserved cysteines, and conserved aromatic residues.

N+C

Weak

−

Cluster 398 \[1\]. Proteins in the cluster have 12 conserved cysteines, and some are

−

Strong

N+C

−

Cluster 1_0_162 appears specific to

+(−1 or 16% # or 27% wt).

U. appendiculatus

ΔII

ΔBAX

Subcellular localization

GE

Negative

0

−

N+C

nt

TABLE 2 | Effector candidates and results of the experimental assays. ΔHR, suppression of the hypersensitive response; ΔII, suppression of innate immunity; ΔBAX, suppression of yeast cell death. Subcellular localization and predicted subcellular localization (Pred); C, cytoplasm; N, nucleus; GE, gene expression; 0, no regulation, expression levels vary less than tenfold; − in planta down-regulation; + in planta up regulation (−/+ > tenfold, −−/++ > hundredfold, −−−/+++ > thousandfold; nt, not tested).

<table>
<thead>
<tr>
<th>Name</th>
<th>Description of the gene families that the effector candidates belong to and where applicable the actual protein*</th>
<th>ΔHR</th>
<th>ΔII</th>
<th>ΔBAX</th>
<th>Subcellular localization</th>
<th>GE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uaca_1</td>
<td>Family of rust transferred proteins [Kemen et al., 2005; Puthoff et al., 2009]. Likely functions are formation of fibrils [Kemen et al., 2013] and inhibition of proteases [Pretsch et al., 2013].</td>
<td>–</td>
<td>–</td>
<td>Medium</td>
<td>All N, some C</td>
<td>nt</td>
</tr>
<tr>
<td>Ua_RTP1</td>
<td>Expressed in planta (cell wall fraction) [2].</td>
<td>–</td>
<td>–</td>
<td>Strong</td>
<td>N+C</td>
<td>0</td>
</tr>
<tr>
<td>Uaca_2</td>
<td>Family of rust transferred proteins [Kemen et al., 2005; Puthoff et al., 2009]. Likely functions are formation of fibrils [Kemen et al., 2013] and inhibition of proteases [Pretsch et al., 2013].</td>
<td>–</td>
<td>–</td>
<td>Medium</td>
<td>N+C</td>
<td>−</td>
</tr>
<tr>
<td>Ua_RTP2</td>
<td>Expressed in planta (P30, cell membrane and organelle fraction) [2].</td>
<td>–</td>
<td>+</td>
<td>Strong</td>
<td>N+C</td>
<td>+</td>
</tr>
<tr>
<td>Uaca_3</td>
<td>Cluster 1.0.152 is mostly lineage specific for U. appendiculatus; also has a motif similar to RxLR [1].</td>
<td>–</td>
<td>+</td>
<td>Strong</td>
<td>C, aggregation</td>
<td>++</td>
</tr>
<tr>
<td>Uaca_4</td>
<td>Cluster 398 [1]. Proteins in the cluster have 12 conserved cysteines, and some are highly expressed or up-regulated in planta. Uaca_4 itself is abundantly expressed and closest homolog to UHSP42, highly expressed in the haustorium [Link and Voegele, 2008].</td>
<td>–</td>
<td>+</td>
<td>Strong</td>
<td>N+C</td>
<td>++</td>
</tr>
<tr>
<td>Uaca_5</td>
<td>Cluster 20 is specific to Pucciniales when only Basidiomycetes are concerned but contains members from other plant pathogens also. Horizontal gene transfer? [1]</td>
<td>–</td>
<td>+</td>
<td>Strong</td>
<td>C, aggregation</td>
<td>++</td>
</tr>
<tr>
<td>Uaca_7</td>
<td>Cluster 398 [1]. Proteins in the cluster have 12 conserved cysteines and some are highly expressed or up-regulated in planta. Uaca_7 itself is most abundantly expressed.</td>
<td>–</td>
<td>+</td>
<td>Medium</td>
<td>N+C</td>
<td>++</td>
</tr>
<tr>
<td>Uaca_9</td>
<td>Cluster 112 has most members [1] and contains P. pachyrhizi proteins (PpEC_23, de_novo_3939, de_novo_7164, de_novo_1784) [Qi et al., 2016; De Carvalho et al., 2017; Qi et al., 2018] showing defense suppression; of the Ua homologs Uaca_9 fits the HMM second best. Proteomics showed expression of the protein both during spore germination and in planta (mostly cytoplasmic fraction) [2].</td>
<td>+</td>
<td>+</td>
<td>Strong</td>
<td>N+C</td>
<td>0</td>
</tr>
<tr>
<td>Uaca_10</td>
<td>Cluster 112 has the most members [1] and contains P. pachyrhizi proteins (PpEC_23, de_novo_3939, de_novo_7164, de_novo_1784) [Qi et al., 2016; De Carvalho et al., 2017; Qi et al., 2018] showing defense suppression; of the Ua homologs Uaca_10 fits the HMM third best.</td>
<td>–</td>
<td>–</td>
<td>Medium</td>
<td>C, aggregation</td>
<td>0</td>
</tr>
<tr>
<td>Uaca_11</td>
<td>Cluster 2456, lineage specific to Pucciniaceae [1]. Fits the pattern best.</td>
<td>–</td>
<td>–</td>
<td>Strong</td>
<td>C</td>
<td>++</td>
</tr>
<tr>
<td>Uaca_12</td>
<td>Cluster 2456, lineage specific to Pucciniaceae [1]. Fits the pattern second best.</td>
<td>+</td>
<td>–</td>
<td>Medium</td>
<td>C</td>
<td>+++</td>
</tr>
<tr>
<td>Uaca_14</td>
<td>Cluster 1.2 is lineage specific to Pucciniales and has eight conserved cysteines and two additional motives [1]. Uaca_14 fits the pattern second best. Proteomics also found the protein expressed in planta in the cell wall fraction, as well as other proteins from the family: UAHYP_19A_R_H01, Ua_contig00019 [2].</td>
<td>+</td>
<td>–</td>
<td>Medium</td>
<td>C</td>
<td>++</td>
</tr>
<tr>
<td>Uaca_16</td>
<td>Cluster 2568 is lineage specific to Pucciniaceae, maybe even Uromyces. Uaca_16 is one of the few proteins with haustorial expression in the family. Proteins in the family have highly similar sequences, there are no cysteines, very high content of tyrosine (17 or 16% # or 27% wt).</td>
<td>–</td>
<td>–</td>
<td>Strong</td>
<td>−</td>
<td>−−−</td>
</tr>
<tr>
<td>Uaca_20</td>
<td>Cluster 145 has three conserved cysteines, and conserved aromatic residues.</td>
<td>–</td>
<td>–</td>
<td>Medium</td>
<td>N+C</td>
<td>+</td>
</tr>
<tr>
<td>Uaca_22</td>
<td>Cluster 974 has only few homologs per species, seven or three conserved cysteines plus one not conserved cysteine. Serine- and threonine-rich, also many aromatic aa.</td>
<td>–</td>
<td>+</td>
<td>Medium</td>
<td>C</td>
<td>0</td>
</tr>
<tr>
<td>Uaca_23</td>
<td>Cluster 2768 is lineage specific to Pucciniaceae, has six conserved cysteines and consists of very short proteins. Uaca_23 is the longest protein of the cluster. Proteomics found the protein both in germlings and in planta (cell wall fraction) [2].</td>
<td>–</td>
<td>–</td>
<td>Medium</td>
<td>C</td>
<td>0</td>
</tr>
<tr>
<td>Uaca_24</td>
<td>Cluster 1.0.162 appears specific to U. appendiculatus. It is a very small family, has very short proteins (100 \text{ aa}) and four conserved cysteines.</td>
<td>–</td>
<td>–</td>
<td>Negative</td>
<td>C</td>
<td>++</td>
</tr>
<tr>
<td>Uaca_25</td>
<td>Cluster 1_63 is specific to rust fungi. Uaca_25 has the longest sequence of the family.</td>
<td>–</td>
<td>–</td>
<td>Negative</td>
<td>N+C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_27</td>
<td>Cluster 2822 is specific to U. appendiculatus. It is a small family of short secreted proteins with cysteines.</td>
<td>–</td>
<td>–</td>
<td>Strong</td>
<td>N+C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_28</td>
<td>Cluster 2917, basically just one gene with one homolog in U. fabae; specific to Uromyces.</td>
<td>–</td>
<td>+</td>
<td>Negative</td>
<td>N+C</td>
<td>−−</td>
</tr>
<tr>
<td>Uaca_31</td>
<td>Cluster 824 is not very conserved; N-terminus differs strongly but conserved cysteines are present. Lineage specific to Pucciniales.</td>
<td>–</td>
<td>–</td>
<td>Negative</td>
<td>N+C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_32</td>
<td>Cluster 1293, only one homolog per species. By proteomics found in planta, cell wall fraction [2].</td>
<td>–</td>
<td>–</td>
<td>Strong</td>
<td>C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_34</td>
<td>Cluster 2484 consists of only two very similar homologs. Very short proteins, two conserved cysteines. Lineage specific to Pucciniales.</td>
<td>–</td>
<td>–</td>
<td>Strong</td>
<td>N+C</td>
<td>nt</td>
</tr>
</tbody>
</table>

(Continued)
**TABLE 2 | Continued**

<table>
<thead>
<tr>
<th>Name</th>
<th>Description of the gene families that the effector candidates belong to and where applicable the actual protein</th>
<th>ΔHR</th>
<th>ΔII</th>
<th>ΔBAX</th>
<th>Subcellular localization</th>
<th>GE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uaca_36</td>
<td>Cluster 1_229, only two homologs. Lineage specific for <em>U. appendiculatus</em>, 10 conserved cysteines.</td>
<td>–</td>
<td>–</td>
<td>Strong</td>
<td>N+C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_37</td>
<td>Cluster 464 is lineage specific to Pucciniales; eight conserved cysteines.</td>
<td>–</td>
<td>–</td>
<td>Negative</td>
<td>N+C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_38</td>
<td>Cluster 1206 is lineage specific to Pucciniales; conserved cysteines.</td>
<td>–</td>
<td>–</td>
<td>Weak</td>
<td>N+C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_40</td>
<td>Cluster 2240 is specific to Pucciniales; twelve conserved cysteines with a similar pattern to that of cluster 112.</td>
<td>–</td>
<td>–</td>
<td>Strong</td>
<td>N+C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_41</td>
<td>Cluster 2826, specific to Pucciniaceae.</td>
<td>–</td>
<td>–</td>
<td>Medium</td>
<td>C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_42</td>
<td>Cluster 3063 is lineage specific to <em>U. appendiculatus</em> and consists of only two similar proteins with twelve or eighteen cysteines respectively, which might be splicing variants. Uaca_43 is the shorter protein.</td>
<td>–</td>
<td>–</td>
<td>Strong</td>
<td>N+C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_43</td>
<td>Cluster 3112 is specific to Puccinia; only one homolog per species</td>
<td>–</td>
<td>+</td>
<td>Weak</td>
<td>N+C</td>
<td>++</td>
</tr>
<tr>
<td>Uaca_44</td>
<td>Cluster 3113, specific to <em>U. appendiculatus</em>, six conserved cysteines, of the two proteins of the family Uaca_45 is slightly shorter and has more serines.</td>
<td>–</td>
<td>–</td>
<td>Medium</td>
<td>N+C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_45</td>
<td>Cluster 3121, lineage specific to <em>Uromyces</em></td>
<td>–</td>
<td>–</td>
<td>Negative</td>
<td>C</td>
<td>nt</td>
</tr>
<tr>
<td>Uaca_46</td>
<td>Cluster 3121, lineage specific to <em>U. appendiculatus</em></td>
<td>–</td>
<td>–</td>
<td>Strong</td>
<td>N+C</td>
<td>nt</td>
</tr>
</tbody>
</table>

*Here we explain why each Uaca_N was chosen. Clustering and motif results refer to Link et al. (2014) [1]; proteomics results (Cooper et al., 2016) [2] are also mentioned. Additional literature has full citation in the table. No overlap between our candidates and silenced effector candidates (Cooper and Campbell, 2017) was found.

**Expression Patterns of Uaca_Ns**

For 17 Uaca_Ns that were assumed to have effector function after the assays described above, we also tested the expression pattern (Figure 5, Supplementary Figure 3). Using RT-qPCR, we tested Uaca_N expression in three stages produced *in vitro* (ungerminated urediospore, germ tube, and appressorial stage) and five *in planta* stages (3, 5, 7, 10, and 14 dpi). The first three *in planta* stages are meant to represent infection and haustoria formation, and the latter two time points encompass sporulation and senescence. *In planta* stages earlier than 3 dpi were not tested because of the low proportion of fungal RNA in total RNA prepared from inoculated leaves, which causes highly variable results. Based on the fact that the sequences were initially found in the haustorial transcriptome, we expected the genes to be up-regulated *in planta*. This was the case for nine of the 17 effector candidates tested for gene expression. Two Uaca_Ns were up-regulated >10-fold (Uaca_4, Uaca_20), six >100-fold (Uaca_5, Uaca_7, Uaca_11, Uaca_14, Uaca_24, Uaca_44), and one >1000-fold (Uaca_12) (Figure 5A, Supplementary Figure 3). On the other hand five genes showed no or little differential expression, including Uaca_9 and Uaca_10 (Figure 5B, Supplementary Figure 3). Three of the tested Uaca_Ns, including Ua_RTP2, were even down-regulated *in planta*, Ua_RTP2 >10-fold, the others >100-fold. All three genes showed maximal expression in the appressorial stage (Figure 5C,
**FIGURE 2** | Suppression of basal defense by Uaca_Ns. The white dashed lines show an outline of the patches infiltrated with *P. fluorescens* strain EtHAn with empty pEDV6 vector control or expressing Uacas from pEDV6, and the red dashed lines show the patches infiltrated with *Pst* DC3000 seven hours later. (A) left upper patch, EtHAn with empty vector control; left lower patch, EtHAn with Uaca_4; right upper patch, EtHAn with Uaca_5; right lower patch, EtHAn with Uaca_7; (B, C, D) upper patch, EtHAn with empty vector control; lower patch, EtHAn with Uaca_9 (B), Uaca_28 (C), and Uaca_44 (D). The inoculum density of EtHAn strains and *Pst* DC3000 strains was adjusted to OD$_{600}$nm = 0.2 and 0.02, respectively. Images were taken 48 hpi of EtHAn strains. At least three independent repeats of this assay were performed for each Uaca_N.

**FIGURE 3** | Examples of phenotypes of the yeast cell death suppression assay. Growth phenotypes of serial dilutions of yeast strains grown on medium that induced the expression of *bax*. Uaca_Ns with representative phenotypes were chosen for display. Strong suppression was defined by growth detected at the $10^{-4}$ dilution, Medium suppression was defined by growth detected at the $10^{-3}$ dilution, Weak suppression was defined by growth at the $10^{-1}$ or $10^{-2}$ dilution, and Non-suppression by no growth at any dilution. This assay was performed two independent times.
Supplementary Figure 3). The high expression of some candidate effectors at the appressorial stage suggests that they have functions that are needed during early infection processes.

DISCUSSION

Relatively little is known about the functions of effectors produced by rust fungi. There are very few examples of bona fide effectors known for the rust fungi (Petre et al., 2014). These few were either identified as avirulence (Avr) proteins (Ellis et al., 2007; Upadhyaya et al., 2014) or directly localized inside plant cells by immunofluorescence microscopy (Kemen et al., 2005). Some of these effectors have been analyzed down to the crystal structure (Wang et al., 2007; Ve et al., 2013) and functions were inferred (Wan et al., 2019) but even for most of those effectors, little is known about their biochemical functions and their roles in virulence (Petre et al., 2014). In contrast to this, it is hypothesized that there are many effectors and that they have important functions, especially in suppression of host resistance. This hypothesis is supported by results with other plant pathogens, such as Oomycetes like Phytophthora infestans, where hundreds of effectors were found (Tyler et al., 2006). Results from genome or transcriptome sequencing of different rust species are consistent with oomycete data in that large numbers of secreted proteins have been predicted, and most of these are of unknown function, opening the possibility that they could have effector function (Duplessis et al., 2011; Cantu et al., 2013; Nemri et al., 2014).

In comparison to bacteria and oomycetes, research on rust effectors is hampered by the fact that so far no common sequence feature for effectors has been identified. To overcome this limitation, other criteria were used to determine whether secreted proteins are likely effectors. Saunders et al. (2012) for example built an effector prediction pipeline based on the following criteria: expression induced in planta, similarity to HESPs or Avrs, known effector motif or NLS, small and cysteine-rich, repeat containing, long intergenic region, or no PFAM domain. The central element of this pipeline is clustering of the proteins into families. In our study on the haustorial transcriptomes of P. pachyrhizi and U. appendiculatus (Link et al., 2014), we also used clustering to identify families of secreted proteins, and then established whether the family was specific to the rusts or lineages within the rusts, which was a paramount criterion for classifying a secreted protein as an effector candidate.

The other approaches to identify effectors are experimental screens. Secreted proteins can be screened against cultivars with known resistance genes to find Avrs, which yielded an effector candidate. 

FIGURE 4 | Representative examples of Uaca_N subcellular localization. (A) Uaca_1, localized in the nucleus; (B) Uaca_2, localized in both nucleus and cytoplasm; (C) Uaca_5, localized in the cytoplasm and forming aggregates; (D) Uaca_11, localized in the cytoplasm. The scale bar is equal to 50 µm. Images were taken 48 hpi. Two replications of the imaging were performed, and at least four representative images were taken each time.
in coffee rust (Maia et al., 2017), or against non-host plants to identify effector-specific recognition (Dagvadorj et al., 2017). Very recently, a screen for effectors suppressing RNA silencing was successful (Yin et al., 2019). Most important are screens based on subcellular localization (Petre et al., 2015; Petre et al., 2016) or suppression of defense responses (Sohn et al., 2007). It has been discussed that the protein fusion has an effect on the outcome of such assays, and that results could differ between experiments with N-terminal fusions and C-terminal fusions. We recently implemented a combination of different screens to identify secreted proteins of *P. pachyrhizi* with effector-like functions (Qi et al., 2018). Here we used N-terminal fusions based on the rationale that the tags are replacing the signal peptide. For the first effector candidate that we identified this way, PpEC23 (Qi et al., 2016), we found that the C-terminus is important for its interactions with a host-transcription factor. This finding reassured us that N-terminal fusions can be functional. We cannot exclude that for some effector candidates C-terminal fusions would be the better alternative but it was not feasible for us to do the screens with both N- and C-terminal fusions.

For *U. appendiculatus*, we combined our theoretical predictions with our experimental screens. The selective criteria we used to identify Uaca_Ns yielded a higher proportion of secreted proteins with positive results in the *in planta* assays than
we previously observed with the *P. pachyrhizi* candidate effectors. Six effector candidates or 19% could suppress basal defense compared to 14%. With only six negatives, the vast majority of our candidates could suppress BAX-induced cell death in yeast. Most striking was that we could find four Uaca_Ns that can suppress the hypersensitive response whereas only one such protein could be found for *P. pachyrhizi*. Therefore, we conclude that a pre-selection of candidate effectors based on sequence features can help reduce experimental efforts.

Also, by covering many different protein families our results can point to other proteins, which may be valuable targets for testing. Because many of the families of effector candidates extend between species of the rust fungi, researchers interested in different species may profit from our results. As a caution, it should be mentioned that it is highly likely that expression patterns, targets, and in consequence, functions of effectors in the same family are different. Therefore, the results, even though they do not constitute functional characterization, should not be directly transferred to other proteins of the family—rather, our results are meant to point at interesting gene families. Several studies have used clustering to group families of secreted proteins, but it is challenging to make comparisons among the families from these different studies. So far, matching our gene families to the clusters found by other groups, e.g. Saunders et al. (2012), Cantu et al. (2013), Nemri et al. (2014), or De Carvalho et al. (2017) remains a laborious exercise. A second challenge is that transcriptome studies are inherently incomplete. We expect that as more genomes become available more comprehensive comparisons can be performed. This is an especially exciting potential outcome of the rust pangenomics project, “Reference Genomes for 50 Rust Fungi” (https://jgi.doe.gov/csp-2018-duplessis-reference-genomes-50-rust-fungi/).

A few genes and gene families that merit special attention are discussed below.

The first two candidates were chosen to obtain further information on proteins of the RTP family (Putthoff et al., 2008; Pretsch et al., 2013). *Uf*-RTP1p has been shown to induce fibril formation inside host cells, and it also has protease inhibitor activity (Kemen et al., 2013; Pretsch et al., 2013). These two functions indicate a possible structural role for *Uf*-RTP1p as well as a role in protecting the fungus against an active defense response mediated by proteases. Here, we found that both *Ua*RTP1 (Uaca_1) and *Ua*RTP2 (Uaca_2) did not suppress plant defenses in the assays performed. Therefore, no additional functions can be added to the RTP portfolio. At this time, we are not sure if the suppression of BAX induced cell death that was observed (medium for both *Ua*RTP1 and *Ua*RTP2) has relevance. The localization of our GFP fusions nicely coincide with the immunolocalization of the native protein *Uf*-RTP1p and also *Us*RTP1p (Kemen et al., 2005). Kemen et al. (2005) could observe that *Uf*-RTP1p first accumulates around the haustorium or more specifically in the extrahaustorial matrix (Kemen et al., 2013). In cells infected with older haustoria, the protein was found in the cytoplasm and also in the nucleus. Also, an NLS was found in the *Uf*-RTP1 sequence that could account for the nuclear localization. The GFP fusion proteins are also localized in the cytoplasm and the nucleus inside *N. benthamiana* cells. This coincidence indicates that heterologous expression of GFP fusions of effector candidates in *N. benthamiana* can correctly predict the targeting of the native effector protein. On the other hand, the study of Cooper et al. (2016) found *Ua*RTP1 preferentially in cell walls. The latter proteomic result can be explained by the finding that *Uf*-RTP1p first accumulates in the extrahaustorial matrix before also being imported into the plant cytoplasm. The result for gene expression for Uaca_2 was surprising. The gene is most highly expressed in the appressorial stage but down-regulated in planta. This seems to be in contradiction to the results from Cooper et al. (2016). This discrepancy could be explained by the different experimental approaches targeting proteins and transcripts respectively, however. Also, the fungal isolates and the plant genotypes used in the respective experiments were different. Our results here indicate that proteins from the RTP family may have different expression patterns and since they end up in different fungal structures also may have different functions.

Uaca_9 and Uaca_10 were chosen as members of cluster 112, which is lineage specific to Pucciniales. The predominant feature of members of this cluster is a motif of 10 conserved cysteines. *PpEC_23*, the first effector candidate from *P. pachyrhizi* for which we could show suppression of plant immunity (Qi et al., 2016; Qi et al., 2018) belongs to this cluster. Three more *P. pachyrhizi* proteins from this family were also tested (De Carvalho et al., 2017), and these also could suppress immunity in *N. benthamiana*. All the GFP fusion proteins were localized in the cytoplasm and nucleus with *PpEC_23* exhibiting strong aggregation. It is intriguing that of the two *U. appendiculatus* proteins tested here, Uaca_9 showed clear suppression of plant immunity and also strong suppression of BAX-induced cell death, whereas Uaca_10 did not suppress either basal defense or the hypersensitive response. The localization of both proteins was consistent with other members of the family from *P. pachyrhizi* with Uaca_10 also aggregating. The mRNA expression of Uaca_9 was induced by less than 10-fold in planta and Uaca_10 mRNA transcripts did not significantly change. The lack of dynamic changes in the expression of these two Uaca_Ns is consistent with *PpEC_23*, which had the strongest gene expression in the appressorial stage. These observations suggest that the effectors from this gene family may be needed throughout the infection process. Overall our results provide further evidence that the gene family represented by cluster 112 is a highly interesting family of effector candidates in *U. appendiculatus* as well as in *P. pachyrhizi*. In particular, the domain structure of these proteins and how the domains contribute to defense suppression through interactions with host proteins is of interest.

Seven more Uaca_Ns stand out as highly likely effectors, especially because they either suppress HR or PTI. Since we based the choice of effectors on interesting features of the corresponding protein families, these are highlighted here. To our knowledge, no other proteins from these families were experimentally validated so far. Uaca_4 and Uaca_7 belong to cluster 398, a cluster made up of proteins highly expressed or up-regulated in the haustorium (Link et al., 2014). These are small, cysteine-rich proteins with a pattern of 12 cysteines. Cluster 398 corresponds to tribe 110, which was identified...
by Saunders et al. (2012). Our RT-qPCR results showed that Uaca_4 and Uaca_7 are also strongly up-regulated in planta by 3 dpi and, therefore, are expressed in a manner consistent with previous results for this family. Both proteins were able to suppress BAX-induced cell death in yeast and innate immunity, and they were localized to the cytoplasm and the nucleus when expressed in N. benthamiana. Most importantly, these Uaca_Ns suppressed PTI in the overlapping infiltration assay (Figure 2). These data coupled with the small, cysteine-rich nature of these proteins provides strong indicators that members of cluster 398 should be investigated further to understand the mechanisms by which they function in rust–host interactions. Uaca_5 belongs to cluster 20 (Saunders et al. (2012): tribe76) for which our phylogenetic analysis indicates that it could have been acquired by the Pucciniales by horizontal gene transfer relatively early, before most of the Pucciniales species were separated (Link et al., 2014). Another explanation for this phenomenon might be that all other (non biotrophic) basidiomycete species lost the gene because it is only useful in pathogenesis. All our data support the assumption that this protein is an effector: it suppresses innate immunity and yeast cell death, and it is strongly up-regulated in planta. Uaca_12 and Uaca_11 belong to cluster 2465 (Saunders et al. (2012): tribe213). Although both proteins are strongly up-regulated in planta and suppress yeast cell death, Uaca_12 can also suppress HR, which makes it a highly interesting effector candidate. Uaca_14 also suppresses the HR as well as yeast cell death. Cluster 1_2 that it belongs to (Saunders et al. (2012): tribe38), has conserved cysteines. The expression pattern also fits to what is expected for effectors. Finally, Uaca_22 belongs to cluster 874 that has members in several rust species but only few homologs per species. There is suppression of the HR but no strong regulation of the gene.

Our study provides experimental evidence for effector-like functions of proteins from several families of U. appendiculatus effector candidates. The data presented here lays a strong foundation for further work to characterize U. appendiculatus effectors, which may lead to identification of effector targets and novel sources of resistance in common bean. In addition, our study also led to some general findings or conclusions. To begin with, it became clear that selecting candidate effectors based on Pucciniales specificity of the gene family was useful as noted from the higher ratio of positive results in our U. appendiculatus screens versus previous screens of P. pachyrhizi effector candidates (Qi et al., 2018). EffectorP (Sperschneider et al., 2016; Sperschneider et al., 2018) that is trained to predict effectors from sequence features of proteins secreted by pathogenic fungi was not yet available when we did our candidate selection. Recently, we have run the predictions and found that EffectorP1.0 predicted 16 of our candidates as effectors, EffectorP2.0 15 (Data Sheet 1). The predictions differed between the two algorithms for three effector candidates. Among the candidates predicted as effectors are both Ua_RTPs and also the homologs of PpEC23, which is not surprising since both Uf_RTP1 and PpEC23 belong to the positive training set for EffectorP2.0. In this case the prediction of EffectorP is exactly the same as with our approach. A true test of the algorithms is not possible with our data, mostly because our results do not yet identify bona fide effectors. In our opinion, EffectorP would have been useful in conjunction with lineage specificity and gene family when selecting effector candidates for this study.

We also identified slight trends regarding size and cysteine content of the effector candidates in suppression of yeast cell death and plant defenses. Our RT-qPCR measurements of gene expression indicate that effector expression can be highly diverse even though all effector candidates in this study were originally identified as being expressed in haustoria. Three candidates, for which there is strong evidence that they are effectors, showed relatively small changes in mRNA expression or they were even down-regulated in planta. These results indicate that up-regulation in planta may not pose as strong evidence for effector function as previously thought, or that mechanisms of posttranscriptional regulation should also be considered as an important mechanism for controlling effector expression. On the other end of our candidate spectrum, we also found one gene (Uaca_24) that was highly up-regulated in planta, but it was not able to suppress immune responses in the assays performed. This could be an effector with a different function, such as inducing the release of nutrients to the pathogen or other changes in infected cells or tissues. Additional assays for defense suppression and nutrient release are needed to further explore the functions of this and other candidate effectors.

DATA AVAILABILITY STATEMENT

All datasets generated for this study are included in the manuscript/Supplementary Files.

AUTHOR CONTRIBUTIONS

MQ, RV, SW, and TL planned and designed the study; TL chose and cloned the candidate genes; YM recloned Uaca coding sequences into vectors for use in all the expression assays; MQ and LD performed assays for PTI and ETI suppression; JS performed bax cell death suppression experiments, JG performed the epi-fluorescence microscopy, MR did the RT-qPCR measurements; MQ, SW, and TL wrote the manuscript. All authors read and approved the submitted manuscript version.

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SUPPLEMENTARY MATERIAL

The Supplemental Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2019.01182/supplementary-material

DATA SHEET 1 (.xlsx) Microsoft excel file containing sequence IDs that can be found in the public databases (alternative name), the open reading frame sequences together with the peptide sequences and information about the predicted signal peptide that was removed in cloning. The length of the full peptides is also included as well as the number of cysteines contained in the proteins (% C). Prediction results of EffectorP1 and EffectorP2 are incorporated in this table to complement information on effector candidate prediction. EffectorP1 and EffectorP2 were not available when this study was initiated, so their output could not be used for candidate effector selection criteria.

REFERENCES

We, T., Williams, S. J., Catanzariti, A. M., Rafiqi, M., Rahman, M., Ellis, J. G., et al. (2019). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.