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Fast Evolution and Lineage-Specific Gene Family Expansions of Aphid Salivary Effectors Driven by Interactions with Host-Plants

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Abstract

Effector proteins play crucial roles in plant-parasite interactions by suppressing plant defenses and hijacking plant physiological responses to facilitate parasite invasion and propagation. Although effector proteins have been characterized in many microbial plant pathogens, their nature and role in adaptation to host plants are largely unknown in insect herbivores. Aphids rely on salivary effector proteins injected into the host plants to promote phloem sap uptake. Therefore, gaining insight into the repertoire and evolution of aphid effectors is key to unveiling the mechanisms responsible for aphid virulence and host plant specialization. With this aim in mind, we assembled catalogues of putative effectors in the legume specialist aphid, Acyrthosiphon pisum, using transcriptomics and proteomics approaches. We identified 3,603 candidate effector genes predicted to be expressed in A. pisum salivary glands (SGs), and 740 of which displayed up-regulated expression in SGs in comparison to the alimentary tract. A search for orthologs in 17 arthropod genomes revealed that SG-up-regulated effector candidates of A. pisum are enriched in aphid-specific genes and tend to evolve faster compared with the whole gene set. We also found that a large fraction of proteins detected in the A. pisum saliva belonged to three gene families, of which certain members show evidence consistent with positive selection. Overall, this comprehensive analysis suggests that the large repertoire of effector candidates in A. pisum constitutes a source of novelties promoting plant adaptation to legumes.

Key words: Acyrthosiphon pisum, salivary proteins, host adaptation, positive selection, pest evolution, plant defenses.

Introduction

Insects comprise the most diverse group of metazoans, and evidence indicates that the evolution of herbivory has played a fundamental role in promoting their species richness and diversification (Wiens et al. 2015). Almost half of the currently known insect species feed on plants (Wu and Baldwin 2010),
and herbivorous insect groups exhibit faster rates of diversification compared with nonherbivorous species (Wiens et al. 2015). A central hypothesis accounting for higher species richness in herbivorous insects proposes an evolutionary interaction between plant defense mechanisms and plant exploitation strategies of insects (Janz 2011). Furthermore, continuous interactions between host plants and herbivorous insects are predicted to make herbivore generalism difficult and constrain a given insect species to one or a few host species (Forister et al. 2015). Since plants provide not only food resources, but also habitats and mating sites to many herbivorous insects, plant specialization may induce divergent selection in insect populations at a range of traits that can lead to reproductive isolation and speciation (Peccoud et al. 2010; Mullen and Shaw 2014). Attempting to unveil the basic mechanisms of insect herbivory provides opportunities to understand the evolutionary and mechanistic basis of plant specialization by herbivorous insects, in particular, and the diversification of metazoan life, in general.

Aphids (Insecta: Aphidomorpha) are pests of wild and cultivated plants that directly reduce plant nutritients by ingesting phloem sap and indirectly cause diseases by transmitting plant pathogens (Blackman and Eastop 2000; Harris and Maramorosch 2014). Aphids are also excellent subjects for host specialization studies. Their clade is composed of approximately 5,000 species (Blackman and Eastop 2000), and most are considered to be plant specialists (Peccoud et al. 2010). Aphid mouthparts are modified into a rostrum or beak with the mandibles and maxillae forming needle-like styles. Aphid secrete gelling saliva during the early stages of feeding to form a feeding sheath surrounding the styles, and then secrete watery saliva into various plant cells (Moreno et al. 2011). Saliva contains effectors that modulate physiological responses to herbivory and permit feeding (Rodriguez et al. 2017). These effectors are likely exposed to natural selection, in particular by plant surveillance systems and defense mechanisms (Will et al. 2013). Interference with plant defenses through various mechanisms has been demonstrated for several effectors secreted by microbial plant pathogens, which ultimately promotes persistence and even spread of these pathogens (Varden et al. 2017). A subset of effectors, the so-called avirulence proteins, are detected by plant surveillance systems and trigger strong immunity in specific plants, determining the incompatibility (Bent and Mackey 2007).

Effectors are diverse, making prediction of effector functions often difficult from the amino acid sequences. As a result, relatively few salivary effectors from aphids have characterized interactions with active host defense responses. In planta expression of salivary effectors C002, Mp1, and Mp2 from the green peach aphid, Myzus persicae, increases M. persicae fecundity on the host plants Arabidopsis thaliana and Nicotiana benthamiana, whereas expression of orthologous genes from another aphid species (the pea aphid, Acyrthosiphon pisum) in these plants has no effect on M. persicae growth (Pitino and Hogenhout 2013). This observation supports specialization of orthologous effectors to distinct plant species during aphid divergence. Some salivary proteins that are known to contribute to aphid plant exploitation are expressed in salivary glands (Wang et al. 2015a, 2015b) and a few showed sites under positive selection (Pitino and Hogenhout 2013; Thorpe et al. 2016). However, a global and comprehensive evolutionary analysis of aphid salivary genes has yet to be reported.

A catalogue of putative salivary effectors was created for A. pisum upon completion of the genome sequence (The International Aphid Genomics Consortium 2010). Combined transcriptomics and proteomics produced a catalog of 324 secreted proteins (Carolan et al. 2011). A small number of effectors predicted in this catalogue were functionally characterized and have been shown to be involved in plant-aphid interactions (Mutti et al. 2006, 2008; Wang et al. 2015a, 2015b). Significant developments in RNA-seq technology and high-resolution mass-spectrometry (MS) provide new impetus to revise the salivary gene catalogue and to define a new and expanded set of candidate salivary effector genes for further analyses. In addition, the genome sequences of two specialist aphids, the Russian wheat aphid, Diuraphis noxia (Nicholson et al. 2015) and the soybean aphid, Aphis glycines (Wenger et al. 2017), together with that of the generalist green peach, M. persicae, (Mathers et al. 2017) have been recently published. These data offer the opportunity to better understand the evolutionary dynamics of salivary effector candidates, in particular, their suspected role in the adaptation of aphid lineages to their host plants (Pitino and Hogenhout 2013; Rodriguez et al. 2017). The critical effectors that drive aphid–plant interactions may display peculiar gene expression and evolutionary patterns, which we search by combining transcriptomics and proteomics in A. pisum and by conducting a comparative analysis of aphid genomes.

Materials and Methods

Aphids, Plants, and Growth Conditions

Acyrthosiphon pisum lineage LSR1 (used for whole-genome sequencing; The International Aphid Genomics Consortium 2010) was maintained in a growth chamber at 18 °C with a 16 h day/8 h night photoperiod on broad bean, Vicia faba (Castel cultivar), at low density to avoid the production of winged individuals. Vicia faba was grown in a growth chamber at 18 °C with a 16 h day/8 h night photoperiod for 10 days before installation of the aphids.

RNA Sequencing

To prepare RNA samples from aphid salivary glands and alimentary tracts, 9-day-old individuals reared at a density of 10–15 aphids per V. faba plant were rapidly dissected with fine forceps in saline solution. The dissected organs were
Soaked in RNA later (QIAGEN) immediately after dissection to avoid RNA degradation. The dissected tissues were pooled in several batches, and RNA was extracted by Nucleospin RNA XS (Macherey-Nagel) and quantified. On average, RNA samples from 200 pairs of salivary glands or 20 alimentary tracts that were dissected on the same day were pooled for one replicate of RNA-seq experiment. Four replicates were prepared by 4 days of dissection with 5–6 persons.

RNA depletion, single stranded-RNA library preparation, multiplexing, and sequencing were performed by Genevizia (New Jersey, USA). Sequencing was performed on the Illumina HiSeq2500 platform, in a 2 x 125 bp paired-end (PE) configuration in High Output mode (V4 chemistry). Each sample was sequenced on four different flowcell lanes to avoid lane effect. In total, 471,933,074 reads were obtained for eight samples. Raw data is available in NCBI Sequence Read Archive (https://trace.ncbi.nlm.nih.gov/Traces/sra/) with last accessed May 22, 2018 with reference number SRP141110.

Mapping and Differential Expression Analysis
Gene expression of A. pisum salivary glands and alimentary tracts was analyzed using the Acyr_2.0 (GCF_000142985.2) reference genome assembly and the NCBI Acrystosiphon pisum Annotation Release 102, both available at ftp://ftp.ncbi.nlm.nih.gov/ncstl/gest domestica/Acrystosiphon_pisum. The paired-end libraries were mapped on the reference genome using STAR v2.5.2 (Dobin et al. 2013) with the following parameters: outFilterMultimapNmax = 5, outFilterMismatchNmax = 3, alignIntronMin = 10, alignIntronMax = 50,000, alignMatesGapMax = 50,000. Fragment counts per genes were estimated by Subread featureCounts (Liao et al. 2014) using default parameters. Differential expression analysis between SGs and AT was then conducted following the workflow proposed by Law et al. (2016). The raw fragment counts were converted to counts per million (CPM) using the edgeR (Robinson et al. 2010) R-implemented package (R-Core Team 2017). Expressed genes were filtered based on a CPM > 1 in at least three libraries among the eight analyzed libraries and CPMs were normalized by the edgeR TMM method for Normalization Factor calculation (Robinson and Oshlack 2010). The mean–variance relationship of the log-CPM was estimated by the voom function (Law et al. 2014) and incorporated in the empirical Bayes analysis from the limma package (Ritchie et al. 2015) to fit linear models and compare SG vs. AT tissues. Validation of the described differential expression analysis is presented in supplementary fig. S1, Supplementary Material online. In addition, normalized fragments per kilobase million (FPKM) were calculated with edgeR on the four salivary glands samples and the four alimentary tracts samples separately to obtain whole transcriptomes of each organ. Genes with FPKM > 0.5 in at least three libraries per tissue were considered as expressed.

Saliva and Salivary Gland Proteomics
LSR1 aphids of mixed ages were reared on V. faba and approximately 2,000 aphids were installed on 12 perspex rings (radius 4.5 cm, height 5 cm), each containing 5 ml of a chemically defined diet formulation AP3 (Febvay et al. 1988) held between two stretched sheets of Parafilm™. The aphids were reared at 18 °C with 16 h day/8 h night photoperiod and the diets were collected and replaced every 24 h. New V. faba-reared aphids were added to the diets to maintain aphid numbers. The daily collected diets were pooled and stored at −80 °C for later use. Three independent replicates were produced by pooling the collected diet from three daily collections (approximately 150 ml). Pooled diets were concentrated at 4 ºC in a Vivascell 250 Pressure Concentrator using a 5,000 molecular weight cut-off (MWCO) polyethersulfone (PES) membrane. The volume of concentrate was further reduced by centrifugation at 3,400 × g in a Vivaspin 6 with a 5000 MWCO. Proteins from this final concentrate (300 µl) were purified using a 2D Clean-up Kit (GE Healthcare) following the manufacturer’s instructions and the resulting protein pellet was suspended in 6 M urea, 2 M thiourea, 0.1 M Tris–HCl, pH 8.0, and quantified using the Qubit™ protein quantification system (Invitrogen). Ten micrograms were removed from each sample for protein digestion.

Adult LSR1 aphids (14–16 days old, reared on V. faba) were dissected in ice-cold saline and dissected SGs were immediately transferred to 60 µl PBS supplemented with Roche Complete™ protease inhibitor cocktail (PIC; final EDTA concentration: 0.2 mM). Thirty pairs of SGs were pooled per replicate and homogenized with a disposable pestle. Sixty microliter of 12 M urea, 4 M thiourea, and PIC was added and samples were homogenized further, centrifuged at 9,000 × g for 5 min to pellet cellular debris and the supernatant was removed and quantified. One hundred microgram of protein was removed and purified using the 2D Clean-up Kit. The solubilized protein lysates were resuspended in 6 M urea, 2 M thiourea, 0.1 M Tris–HCl, pH 8.0, and requantified. Twenty micrograms were removed from each sample for protein digestion.

Sample Preparation for Mass Spectrometry
For mass spectrometry, three independent biological replicates were analyzed. Fifty-mM ammonium bicarbonate was added to each sample, and proteins were reduced with 0.5 M dithiothreitol (DTT) at 56 °C for 20 min. The proteins were then alkylated with 0.55 M iodoacetamide (IAA) at room temperature for 15 min, in the dark. One microliter of 12 M urea, 4 M thiourea, and PIC was added and samples were homogenized further, centrifuged at 9,000 × g for 5 min to pellet cellular debris and the supernatant was removed and quantified. One hundred microgram of protein was removed and purified using the 2D Clean-up Kit. The solubilized protein lysates were resuspended in 6 M urea, 2 M thiourea, 0.1 M Tris–HCl, pH 8.0, and requantified. Twenty micrograms were removed from each sample for protein digestion.
terminated by adding 1 µl of 100% trifluoroacetic acid (Sigma Aldrich) and incubation at room temperature for 5 min. Samples were centrifuged for 10 min at 13,000 x g and purified for mass spectrometry using either the ZipTip pipette procedure (Millipore) for saliva and C18 Spin Columns (Pierce) for salivary glands, following the manufacturer’s instructions. The eluted peptides were dried using a SpeedyVac concentrator (Thermo Scientific Savant DNA120) and resuspended in 2% v/v acetonitrile and 0.05% v/v trifluoroacetic acid (TFA). Samples were sonicated for 5 min to aid peptide resuspension followed by centrifugation for 5 min at 13,000x g. The supernatant was removed and used for mass spectrometry.

Mass Spectrometry

One microgram of each digested sample was loaded onto a QExactive (ThermoFisher Scientific) high-resolution accurate mass spectrometer connected to a Dionex Ultimate 3000 (RSLCnano) chromatography system. The peptides were separated by a 2–40% gradient of acetonitrile on a Biobasic C18 Picofrit™ column (100 mm length, 75 mm ID), using a 120-min reverse-phase gradient at a flow rate of 250 nl min⁻¹ with a runtime of 50 and 130 min for saliva and salivary glands, respectively. All data were acquired with the mass spectrometer operating in automatic data dependent switching mode. A full MS scan at 70,000 resolution and a range of 400–1,600 m/z was followed by an MS/MS scan at resolution 17,500 and a range of 200–2,000 m/z, selecting the 15 most intense ions prior to MS/MS.

Protein identification of MS/MS data was performed using MaxQuant v1.5.6.5 (www.maxquant.org; last accessed May 22, 2018) following the general procedures and settings outlined in Hubner et al. (2010). The Andromeda search algorithm (Cox et al. 2011) implemented in the MaxQuant software was used to correlate MS/MS data against the protein reference sequence set of A. pisum obtained from the NCBI (27,984 entries, May 2016) and a contaminant sequence set provided by MaxQuant. The following search parameters were used: first search peptide tolerance of 20 ppm, second search peptide tolerance 4.5 ppm with cysteine carbamidomethylation as a fixed modification and N-acetylation of protein and oxidation of methionine as variable modifications and a maximum of two missed cleavage sites allowed. False Discovery Rates (FDR) were set to 1% for both peptides and proteins and the FDR was estimated following searches against a target-decoy database. Peptides with minimum length of seven amino acids were considered for identification.

Data Analysis

Perseus v.1.5.5.3 (www.maxquant.org, last accessed May 22, 2018) was used for data analysis, processing and visualization. The data matrix was first filtered for the removal of contaminants and peptides identified by site. Label Free Quantitation (LFQ) intensity values were log2 transformed and proteins not found in all three replicates in at least one group were omitted from the analysis. A data-imputation step was conducted to replace missing values with values that simulate signals of low abundant proteins chosen randomly from a distribution specified by a downshift of 1.8 times the mean standard deviation (SD) of all measured values and a width of 0.3 times this SD.

Gene Ontology and Secretion Prediction

The GO annotation was performed on the whole A. pisum proteome available on NCBI using Blast2GO v2.5.0 (Götz et al. 2008). Associations were realized using a blastp [BLAST+ v2.5.0 (Camacho et al. 2009)] search against the nonredundant protein database (release 2017-2-4) with the following parameters: e-value = 1e−8, max_target_seqs = 20, soft_making = false, and Interproscan v5.13.52.0 (Jones et al. 2014). Assigned GO terms for genes of interest groups were categorized by molecular function (MF), biological process (BP) and cellular component (CC) and GO enrichment was investigated using hypergeometric tests in R. The P-values were adjusted for multiple comparisons using Bonferroni correction. The whole A. pisum proteome was analyzed with SignalP v3.0 and v4.1 (Dyrøv Bendtsen et al. 2004; Petersen et al. 2011) and SecretomeP v2.0 (Bendtsen et al. 2004) to characterize the presence of signal peptide or nonclassical secretion signal, respectively. Then, membrane inserted domains were predicted using TMHMM v2.0 (Krogh et al. 2001) for transmembrane domains and PredGPI (Pierleoni et al. 2008) for GPI anchors. Finally, the results were combined to define the list of A. pisum secreted proteins that have secretion signals and no membrane insertion domains.

Orthology Analysis

To determine groups of orthologs among the 17 genomes (supplementary table S1, Supplementary Material online), we first kept the longest protein isofoms from each species with an home-made perl script and then ran the OrthoDB_soft_1.6 (Kriventseva et al. 2015) using standard parameters. To establish the species phylogeny, 478 groups of conserved single-copy orthologs were extracted and their protein sequences aligned with MAFFT (Katoh et al. 2005). Alignments were concatenated and used to generate a maximum likelihood tree with RAxML (Stamatakis 2006) with default parameters and 1,000 replicates, considering Tetranychus urticae as an outgroup. The phylogeny was then used to establish the level of orthology of each A. pisum gene. The enrichments of orthologous categories among data set were analyzed using hypergeometric test implemented in R.
Evolution Analyses of Salivary Genes

Single Copy Genes

Ortholog groups that were represented by a single gene in the A. pisum genome and in at least one of the other Aphididae genomes were extracted. If alternative transcripts of a gene were present, the longest coding sequence (CDS) for each species was used. Pairs of orthologous CDS (one CDS per species: A. pisum, A. glycines, A. pisi MD. noxia, A. pisum/ M. persicae) were aligned using MAFFT (Katoh et al. 2005) within the Hot algorithm from GUIDANCE2 (Sela et al. 2015) to produce reliable codon-based alignments without poorly aligned regions. From these codon alignments, pairwise dN/dS values were calculated with PAML v4.8 using the YN00 program (Yang and Nielsen 2000; Yang 2007). We removed ortholog pairs with dS > 2 or dN > 0.5 to avoid mutational saturation, as well as pairs with dS = 0. dN/dS of different gene categories within each species pair were compared using a Kruskal–Wallis test, and Nemenyi-Tests for multiple gene categories within each species pairs were compared as described in the supplementary material and methods S1, Supplementary Material online, in order to compute more accurate evolutionary rates and to test selection models. For this task, reciprocal blast searches were run with blastn (BLAST+ v2.5.0) using A. pisum, M. persicae, A. glycines and D. noxia CDS sequences against the different databases (e-value < 10^-10). Then, for each obtained ortholog group, a codon-based alignment was generated with PRANK (Löytynoja and Goldman 2005) and cleaned from poorly aligned regions using GUIDANCE2, specifying a minimum alignment quality threshold of 0.93. The cleaned alignment was used to generate a maximum likelihood phylogenetic tree with RAxML and both alignment and tree served to estimate dN/dS with codeml [implemented in PAML v4.8 (Yang 2007)] under different models (Yang et al. 2000; Yang and Nielsen 2000). The null models (M0 = one ratio, M1 = neutral, M7 = β) were compared with alternative models (M2 = selection, M8 = β + ω) using the likelihood ratio test (LRT), which compares twice the difference in log likelihood to a χ^2 distribution. Finally, if selection models were found more likely, the Bayes Empirical Bayes (BEB) procedure was used to compute the posterior probability of evolution under positive selection for each site (Yang et al. 2005).

Gene Families

Families containing genes expressed in A. pisum salivary glands and their orthologs in arthropod species were extracted from OrthoDB groups. Then, we examined if the two salivary gene catalogs contain more members of multigene families than expected by chance. For each catalog, a significant effect is demonstrated if the number of genes that belong to multigene families that are represented by two or more copies in the catalogue lies above the 95% confidence interval (CI) of the expectation. 95% CI was computed by randomly sampling the number of genes contained in a catalog (740 and 3,603 genes) from the list of 18,603 genes and counting the number of gene-family members (that belong to a multigene family represented by two more copies in the sample) in this random sample. This step was repeated 10,000 times.

To test for positive selection on certain sites and branches (selection acting on foreground branch compared with background branches) of the three selected gene families (Cysteine-Rich Protein, Angiotensin-Converting-Enzyme-like and Aminopeptidase-N families), a cleaned alignment and a maximum-likelihood tree were generated as described above, and were fed to the codeml branch-site (BS) model (Yang and Nielsen 2002). The BS model classified the sites in four categories: class 0 where sites were under negative selection (ω = 1) on both foreground and background branches, class 1 where sites evolved under neutral evolution (ω = 1) on both foreground and background branches, class 2a where the sites were under positive selection (ω ≥ 1) on the foreground branch and under negative selection (ω < 1) on background branches, and class 2b where the sites evolved under positive selection (ω ≥ 1) on the foreground branch and under neutral evolution (ω = 1) on background branches. The null model in which the foreground branch may have different proportions of sites under neutral evolution (dN/dS of 1), and the alternative model, in which the foreground branch may have sites under positive selection, were applied to all branches of each gene family. The likelihood of the selection model was computed as described in the previous section and P-values were corrected for multiple testing comparisons as described by Anisimova and Yang (2007). In case the selection model was retained, the posterior probability of particular sites evolving under positive selection was computed.

All phylogenetic trees shown in figures were designed in iTOL (Letunic and Bork 2007).

Results

Generation of the A. pisum Salivary Effector Candidate Gene Sets

We conducted transcriptome analyses of A. pisum salivary glands (SGs) to generate two catalogues of salivary effector candidates: one that considers up-regulation of their expression in SGs and another that does not. This approach assumes that the vast majority of salivary proteins secreted into plants are expressed in aphid SGs (Mutt et al. 2008; Carolan et al. 2011; Naessens et al. 2015) and that most salivary effectors
are highly expressed in SGs compared with other organs (Wang et al. 2015a, 2015b). Up-regulation of expression in SGs was determined in comparison to expression levels in the alimentary tract (AT), which we chose due to ease of isolation and RNA extraction. RNA-seq by Illumina technology was conducted on dissected SG and AT tissues of *A. pisum*. In SGs, 12,040 genes passed the cut-off value for gene expression (fig. 1A) and encode proteins that may or may not be secreted in saliva. A protein secretion prediction pipeline was applied to the global gene set of *A. pisum* (N = 18,601) using SignalP v3 or v4.1 (Dyrløv Bendtsen et al. 2004; Petersen et al. 2011), as well as SecretomeP (Bendtsen et al. 2004), which predicts noncanonical secretion signals. Genes encoding transmembrane domains [predicted by TMHMM 2.0 (Krogh et al. 2001)] or GPI-anchors [predicted with PredGPI (Pierleoni et al. 2008)] were considered as not secreted, leaving 3,603 encoded proteins predicted to be secreted. These constituted the candidate SG-expressed effector set (fig. 1A and supplementary table S2, Supplementary Material online).

Identification of Aphid Proteins in Artificial Diets and Salivary Glands

Proteins from artificial diets fed upon by aphids were analyzed by proteomics-based mass spectrometry (MS). Fifty-one proteins were supported by more than one peptide detected by MS in at least two out of the three replicates or by one peptide in all three replicates of saliva samples from artificial diets.

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**Fig. 1.**—Pipelines used to establish sets of candidate *Acyrthosiphon pisum* salivary effector genes expressed and up-regulated in salivary glands (A). Pipeline used to identify proteins from saliva injected in artificial diet and composition of the secreted proteins (B).
Aphididae Specific Genes Are Enriched in the SG-Up-Regulated Effector Set

To assess the phylogenetic distribution of genes constituting the SG-up and SG-expressed effector sets, an analysis of orthology was conducted between A. pisum and 16 other arthropods whose genome sequences were available, and which cover a wide range of divergence from A. pisum (fig. 2A). Relative to the whole set of genes annotated in the A. pisum genome, the SG-up effector set was significantly enriched in highly conserved genes found across the arthropod or insect clades (hypermorphic test, \( P < 0.01 \)). The set was also slightly enriched in genes found only in Aphidomorpha, whereas the proportion of A. pisum-specific genes was significantly reduced (hypermorphic test, \( P < 0.001 \)). In contrast, in the SG-up effector set, the proportion of genes found only in Aphididae was significantly higher than in the whole gene set, and highly conserved genes found across arthropods were less frequent (hypermorphic test, \( P < 0.001 \)) (fig. 2B). In our comparison, this pattern was specific to the SG-up effector set as the genes expressed or up-regulated in ATs showed significant enrichment of highly conserved genes found across arthropod and insect clades (supplementary fig. S3A, Supplementary Material online).

Evolutionary rates vary among of A. pisum Salivary Candidate Effectors

The enrichment of Aphididae-specific genes in the SG-up effector set may be associated with rapid molecular evolution related to aphid-specific gene functions. This hypothesis was tested for the SG-up effector set by estimating the ratio of nonsynonymous to synonymous substitutions (\( dN/dS \) or \( \omega \)) in three categories: SG-up effector set, SG-expressed effector set and the remaining genes (fig. 3). Only single-copy gene pairs of orthologs between A. pisum and the three other Aphididae species were included in the orthology search, and pairwise evolutionary rates were analyzed (A. pisum/A. gossypii, A. pisum/D. noxia, A. pisum/M. persicaria). On average, genes of SG-up effector set showed higher \( dN/dS \) ratios than the genes of other sets (fig. 3). The same analysis comparing AT-up, AT-expressed and remaining genes revealed that the observed overall higher \( dN/dS \) ratio was specific to the genes of SG-up effector set (supplementary fig. S3B, Supplementary Material online).

The 11 single-copy salivary effector genes that were previously functionally characterized and shown to be involved in plant–aphid interactions were examined (table 1). All these single copy salivary effectors or their A. pisum orthologs were found in the SG-up effector set with the exception of Mif1 (Naessens et al. 2015), which was not up-regulated in SGs and not predicted to be secreted in our study (table 1). Most of these genes were highly expressed, and genes C002, M1, Ap25, Me23, Me10, Mp55, Shp, and M2p were Aphididae- or Aphidomorpha-specific.

To examine the evolutionary rates of these 11 effectors, we retrieved the orthologs from SG transcripts of two additional aphid species, Schizaphis graminum and Rhopalosiphum padi (supplementary material and methods S1 and table S4, Supplementary Material online). In addition, orthologs from other aphids were retrieved from public databases (supplementary table S4, Supplementary Material online) and used to estimate \( dN/dS \) ratios under different site-models of selection. The global \( dN/dS \) for each gene showed a range from 0.06298 (Arnet) to 0.64653 (Mpo) (table 1 and supplementary table S5, Supplementary Material online). Differences were found between insect- or arthropod-conserved effector genes like Arnet, Mif1 and Mpo10, which seem to have evolved under strong negative selection (\( dN/dS < 0.12 \)), and the aphid-specific, fast-evolving effector genes (\( dN/dS > 0.40 \), in the top 5% of values obtained from pairwise comparisons). Despite these faster evolutionary rates, the examination of null and alternative models of codon substitutions [M1 vs. M2 and M7 vs. M8] revealed signatures of positive selection only for genes Mpo1 and Me10 (supplementary table S5, Supplementary Material online). In Mpo1 and Me10, sites detected as under positive selection were mainly found in the region coding for the mature protein injected into the host plant (supplementary fig. S4, Supplementary Material online).

Episodic positive selection occurred in salivary expanded gene families in the A. pisum Lineage

Evolutionary novelties are usually brought by gene duplication followed by diversification in functions (Conant and Wolfe
Interestingly, duplicated genes are more frequently observed in SG-up and SG-expressed effector sets than expected. Indeed, the observed number of genes that are represented by two or more members of gene family in the two catalogs always lies above the 95% confidence interval (CI) (SG-up: 80 genes, 95% CI = [23, 55] and SG-expressed: 654 genes, 95% CI = [442, 540]). Three multi-gene families in particular, a cystein-rich protein family (CRP), the Angiotensin-Converting-Enzyme-like family (ACE) and the Aminopeptidase-N (apN) family, represented 46.9% of the proteins detected in *A. pisum* saliva (supplementary table S6, Supplementary Material online). Gene ontology analysis showed that the *ACE* and *apN* gene families belong to the metallopeptidase family and are involved in proteolysis, two functions for which the SG-up effector set was enriched (supplementary fig. S5, Supplementary Material online).

**The Cysteine-Rich Protein (CRP) Family**

The CRP Aphididae specific gene family was originally described as a family of 12 genes in *A. pisum* (Guo et al. 2014). Here, 15 gene members were identified from genomic data, and encode proteins of <200 amino acids with 14 highly conserved cysteine residues. The family is expanded in *A. pisum* with 15 copies compared with the other aphid species *D. noxia, A. glycines* and *M. persicae*, which have 6, 4, and 1 copies, respectively. Among the 15 *A. pisum* gene
copies, three were not detected in either SG or AT tissues by RNAseq or protein mass spectrometry. Eleven genes were up-regulated in SGs, nine encode proteins that were predicted to be secreted (supplementary table S6, Supplementary Material online), two of which (CRP-12 and 14) were found in saliva. The branch-site (BS) model of codon substitutions (Yang and Nielsen 2002) detected positive selection in five branches of the Aphididae CRP family tree (fig. 4A and B): the branch leading to D. noxia gene CRP-5 (branch #1), the ancestral branches #2, #4, and those leading to A. pisum CRP-2 (branch #3) and CRP-12 (branch #5). The Bayes Empirical Bayes (BEB) posterior probability (Yang et al. 2005) revealed sites under positive selection (BEB above 0.75) in each aforementioned branch (supplementary fig. S6A, Supplementary Material online), which are scattered across the protein (supplementary fig. S6B, Supplementary Material online). The functional importance of these sites is unknown, due to the absence of known domains within the proteins. Although positive selection seems to occur in some A. pisum CRP copies, no site under positive selection was detected in CRP-13, the most highly expressed A. pisum CRP.

Table 1  
Aphid Single-Copy Salivary Effector Genes Characterized for their Role in the Interaction with Host Plants

<table>
<thead>
<tr>
<th>Gene</th>
<th>Acyrthosiphon pisum Gene</th>
<th>Effector set</th>
<th>Expression Rank</th>
<th>Orthology Level</th>
<th>dN/dS</th>
<th>Aphid Phenotype</th>
<th>References*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mp1</td>
<td>LOC100165393 ACYPI006346</td>
<td>SG-up</td>
<td>19/740</td>
<td>Aphididae</td>
<td>0.6453</td>
<td>Expression promotes fecundity</td>
<td>[3, 4, 6, 10, 15]</td>
</tr>
<tr>
<td>C002</td>
<td>LOC100167863 ACYPI008617</td>
<td>SG-up</td>
<td>12/740</td>
<td>Aphididae</td>
<td>0.62307</td>
<td>Expression promotes fecundity</td>
<td>[1, 2, 3, 4, 6, 13]</td>
</tr>
<tr>
<td>Ap25</td>
<td>LOC100169287 ACYPI009919</td>
<td>SG-up</td>
<td>57/740</td>
<td>Aphididae</td>
<td>0.56219</td>
<td>Expression promotes fecundity</td>
<td>[14]</td>
</tr>
<tr>
<td>Me23</td>
<td>LOC100161198 ACYPI002439</td>
<td>SG-up</td>
<td>109/740</td>
<td>Aphididae</td>
<td>0.53025</td>
<td>Expression promotes fecundity</td>
<td>[5]</td>
</tr>
<tr>
<td>Me10</td>
<td>LOC100167427 ACYPI008224</td>
<td>SG-up</td>
<td>5/740</td>
<td>Aphididae</td>
<td>0.51787</td>
<td>Expression promotes fecundity</td>
<td>[5]</td>
</tr>
<tr>
<td>Shp</td>
<td>LOC100169243 ACYPI009881</td>
<td>SG-up</td>
<td>1/740</td>
<td>Aphidomorpha</td>
<td>0.48863</td>
<td>Silencing reduces survival/fecundity</td>
<td>[8, 12]</td>
</tr>
<tr>
<td>Mp55</td>
<td>LOC100569515 ACYPI33755</td>
<td>SG-up</td>
<td>18/740</td>
<td>Aphididae</td>
<td>0.42972</td>
<td>Expression promotes fecundity</td>
<td>[7]</td>
</tr>
<tr>
<td>Mp2</td>
<td>LOC100160479 ACYPI001774</td>
<td>SG-up</td>
<td>107/740</td>
<td>Aphididae</td>
<td>0.40948</td>
<td>Expression promotes fecundity</td>
<td>[7]</td>
</tr>
<tr>
<td>Mp10</td>
<td>LOC100145855 ACYPI000997</td>
<td>SG-up</td>
<td>303/740</td>
<td>Insect</td>
<td>0.12212</td>
<td>Expression reduces fecundity</td>
<td>[3]</td>
</tr>
<tr>
<td>Mif1</td>
<td>LOC100161225 ACYPI002465</td>
<td>None</td>
<td>4700/12040</td>
<td>Arthropod</td>
<td>0.09454</td>
<td>Silencing reduces fecundity</td>
<td>[9]</td>
</tr>
<tr>
<td>Armet</td>
<td>LOC100167188 ACYPI008001</td>
<td>SG-up</td>
<td>122/740</td>
<td>Arthropod</td>
<td>0.06298</td>
<td>Silencing reduces survival</td>
<td>[11]</td>
</tr>
</tbody>
</table>

*Expression ranks show the rank of effector SG expression level (based on Log[FPKM]) within the SG-up effector set except for Mif1 which does not belong to any effector sets (global SG expression scale).

**x Values represent the evolutionary rates within closely related Aphididae species.

The Angiotensin-Converting-Enzyme-like (ACE) Gene Family

In contrast to the Aphididae-specific CRP gene family, the ACE gene family is found among arthropods (Fig. 5 and supplementary table S6, Supplementary Material online). A search for the ortholog groups yielded ten members in the A. pisum genome [three were previously identified by (Wang et al. 2015b)], which are distributed in four clades along with orthologs from other insect species (Fig. 5 and supplementary table S6, Supplementary Material online). Seven of these genes were up-regulated in A. pisum SG tissue, including three members (ACE1, ACE2, and ACE5) of clade 4 encoding predicted secreted ACEs. ACE1, 5 and 10 were detected in saliva, although the latter lacks an encoded signal peptide, indicating errors in gene annotation or secretion prediction (supplementary table S6, Supplementary Material online). It is notable that the other aphid genomes comprise just one ACE of clade 4 (Fig. 6 A).

**Fig. 4.**—Evolutionary analysis of the Aphididae CRP gene family. (A) Codon-based maximum-likelihood tree used to compute dN/dS under the branch-site (BS) model of codon substitution. Numbers below branches indicate bootstrap support (percentage of the 1,000 replicates) when >40. DNOX, MPER, AGLY, and APIS indicate sequences from *Diuraphis noxia*, *Myzus persicae*, *Aphis glycines*, and *Acyrthosiphon pisum*, respectively. Branches where positive selection affected certain sites are in bold. (B) Estimated evolutionary parameters for these branches (numbered as on the tree). \( \omega \), \( \omega_1 \), and \( \omega_2 \) indicate the average dN/dS ratio for sites assigned to class 0 (\( \omega < 1 \), negative selection), class 1 (\( \omega = 1 \), neutral evolution) and class 2 (\( \omega > 1 \), positive selection), respectively, in the branch, and n.c. indicates insufficient dS to compute \( \omega \). p0, p2a, and p2b show proportions of sites in classes 0, 2a, and 2b, respectively, for each branch (Yang and Nielsen 2002).
this clade (fig. 6B and C): the ancestral branch (#1) of *A. pisum* ACE 1 and ACE5, the *A. pisum* ACE1 gene (#2), and the ancestral branch #3. Sites detected to evolve under positive selection are scattered across the protein (supplementary fig. S7A and B, Supplementary Material online).

**The Aminopeptidase-N (apN) Gene Family**

The aminopeptidase-N (apN) gene family was the most represented in *A. pisum* saliva proteome with 18 apN proteins detected, accounting for 36.7% of the 51 saliva proteins.
Another apN protein not detected here was identified in saliva by Carolan et al. (2009) (supplementary table S6, Supplementary Material online). Forty-seven apN genes were identified in the A. pisum genome, 27 of them belonged to ortholog groups present among insects, and the remaining 20 did not show orthologs in other species (supplementary table S6, Supplementary Material online). All 20 A. pisum-specific apN genes were up-regulated in SG tissue, ten encode proteins with predicted secretion, four of which were detected in saliva. Four of the ten proteins not predicted as secreted were also detected in saliva, indicating incorrect gene annotation and/or secretion prediction. The 27 apN genes with orthologs in other species were clustered in eight distinct clades (fig. 7). The A. pisum genes from clades 1, 2, 3, 4, and 7 were up-regulated in SGs with the exception of only three members. Genes from clades 5, 6, and 8 were not up-regulated in SGs. Apart from the one A. pisum gene that belongs to clade 1, all the other SG-up candidate effector genes were found in clade 4 (fig. 7).

Clade 4 is remarkable for the presence of 14 closely related gene copies in A. pisum and only a few copies from other insect species, including aphids (fig. 7). Eleven A. pisum copies were SG-up-regulated and detected in saliva, but two encode proteins that were not predicted as secreted (fig. 7). Genes of this clade generally encode proteins with a signal peptide followed by a M1 peptidase domain (IPR014782) and an ERAP1-C domain (IPR024571). However, some have lost one of the two domains, and the vast majority (apart from D. noxia apN-3, A. pisum apN-1 and A. pisum apN-5) possesses the HEXXH active site ensuring the peptidase function (fig. 8A).

Traces of positive selection were detected by the branch-site model in seven branches of apN clade 4 (fig. 8B and C): the branch leading to D. noxia gene apN-1 (branch #1), the ancestral branches #3, #5, #7, and those leading to A. pisum apN-3 (#2), apN-8 (#4), and apN-12 (#6) genes. Along these branches, sites under positive selection (with BEB > 0.75) were mainly located in the M1 peptidase and ERAP1-C domains, and in the uncharacterized part between the signal peptide sequence and M1 peptidase domain for branches #1, #2, and #3 (supplementary figs. S8 and S9, Supplementary Material online).

**Discussion**

**New Salivary Candidate Effector Gene Sets**

As evidence accumulates that aphid salivary effectors play important roles in plant-aphid interactions (Elzinga and Jander 2013), a comprehensive identification of aphid salivary
genes and analysis of their evolutionary patterns are essential to shed light on the process of aphid adaptation to specific host plants. Here, we have assembled new catalogues of candidate *A. pisum* salivary effector genes based primarily on comparative transcriptional analysis of SG and AT genes. We identified 3,603 genes expressed in SGs and potentially secreted in saliva. A subset of 740 genes have their expression up-regulated in SGs compared with AT. These sets combined are substantially larger than the previously published candidate effector catalogue (Carolan et al. 2011), reflecting the greater sensitivity of Illumina’s RNA-seq compared with EST sequencing. The salivary gland secretome established for *A. pisum* by Carolan et al. (2011) contains 324 genes, of which 95 are found in our SG-up effector set and 93 others are contained in the SG-expressed effector set. The remaining 120 encompass 111 SG-expressed genes encoding proteins considered as not secreted in our analysis, and 9 genes that were absent from our source transcriptome. Finally, 16 genes did not exist in the updated gene annotation of the *A. pisum* genome used here (NCBI Acrithosiphon pisum Annotation Release 102). The aphid line used here (LSR1) was collected from *Medicago sativa* (The International
Aphid Genomics Consortium (2010) and feeds well on the universal host plant of A. pisum, Vicia faba (our unpublished data). We used V. faba to rear the aphids for RNA-seq experiment because of the ease of producing large quantity of synchronized aphids on this plant. It was also because previous study revealed that A. pisum reared on two suitable legume plants show very similar expression patterns of salivary effector candidates (Eyres et al. 2016). There is a possibility that our catalogues missed the salivary genes that were expressed exclusively when the aphid was feeding on M. sativa, but we assume that such genes are rare.

Comparing expression levels between SGs and AT was fundamental in restricting the list of candidate effectors, which may be refined by sequencing RNAs from other tissues. Nonetheless, salivary genes not up-regulated in SGs should not be ignored. Some may indeed encode salivary effectors like Mf1, which suppresses induction of plant defense responses against aphids (Naessens et al. 2015). In addition, some effector proteins could be produced in tissues other than SGs and injected into plants. Proteins can move from the hemocoel of aphids to SGs to be secreted through saliva. However, only the chaperonin GroEL, which is produced by the aphid obligate endosymbiont in bacteriocytes, is known to be injected into plant tissue through this pathway (Chaudhary et al. 2014).

As 46 proteins out of the 51 (90%) detected in saliva through proteomics were encoded by SG-up genes, most A. pisum salivary proteins seem encoded by genes whose expression is up-regulated in SGs. The lack of significant correlation between the signal intensities of the proteins secreted in saliva and their gene expression level in SGs may result from insufficient statistical power and/or regulation of salivary protein secretion. Plant cues may be required to trigger certain aphid responses, including protein
secretion (Powell et al. 2006), and this may explain why some proteins encoded by highly transcribed salivary genes were not detected in the artificial diet. For example, C002 is reported to be secreted into the plant and required for aphid feeding (Mutti et al. 2008), but was not detected by our saliva proteomics despite its high transcription level. The comparison of transcriptomic and proteomic results revealed limits in the secretion prediction pipeline as 14 proteins detected in saliva were not predicted to be secreted. Five of them may be miss-annotated, but the rest might have been secreted by pathways that were not considered for the secretion prediction.

Fast Evolutionary Rates of Candidate Effectors Genes

SG-up genes showed lower degree of gene conservation than SG-expressed genes. Indeed, 50% of the A. pism SG-up effector set had either no ortholog in other genomes or only in Aphidomorpha and Aphididae, whereas 64.2% of the SG-expressed effector set had orthologs found among arthropods. Furthermore, single-copy SG-up genes presented higher dN/dS ratio than SG-expressed genes on average. This indicates that the evolution of SG-up genes tends to be less constrained (relaxed selection) or accelerated (positive selection). We did not observe such contrast in between AT-up and AT-expressed gene sets, highlighting the peculiar evolutionary history of genes constituting the SG-up effector set.

Interestingly, the analysis of the functionally characterized single-copy genes revealed different selective regimes. The more conserved effectors (e.g., Mif1, Armet, and Mp10) with low dN/dS ratio (<0.12) could be involved in fundamental functions required for plant feeding, for example, prevention of phloem clogging, repression/manipulation of general plant defenses, such that orthologous proteins of similar sequences may be effective on numerous host species (Ros et al. 2010; Furch et al. 2015; Naessens et al. 2015). In contrast, 11 effectors with dN/dS ratios above 0.40 have been characterized as modulators of plant-aphid interactions in functional studies, and their non-synonymous divergence may reflect adaptation to specific host plants. Some of them (C002, Mp1 and Mp2) were indeed shown to act in a species-specific manner to promote aphid performances on host plants (Pitino and Hogenhout 2013). Moreover, sites detected as under positive selection in Mp1 (Pitino and Hogenhout 2013; this study) and Me10 salivary effector genes may be involved in adaptation of their respective aphid lineages.

Although we cannot globally evaluate the relative contributions of relaxed and positive selection regimes on faster gene evolution, several evidences underline the importance of positive selection in shaping the evolution of some candidate effectors identified in our study. It would be unintuitive to hypothesize that SG-up candidate effector genes are more prone to relaxed selection than SG-expressed ones, especially when considering that 90% of the proteins detected in saliva in this study were encoded by SG-up genes. These SG-up effector candidates are more likely secreted by the aphid and thus exposed to selective pressures exerted by the plant surveillance and defense systems. Moreover, we did find traces of positive selection acting on certain sites and branches of SG-up candidate effector genes. Based on these observations and previous functional studies, we argue that the evolutionary history of some salivary effectors has been conditioned by the specialization of SG tissue during establishment of aphid-plant interactions.

Salivary Gene Family Expansions

In addition to single-copy genes, evolutionary histories of three multigenic families were examined. Remarkably, these three gene families encoded nearly half of the proteins detected in saliva and showed gene expansion in the A. pism lineage. Irrespective of their specificity to aphid lineages (CRP family) or conservation among insects (ACE and apN), these families show the highest number of members in A. pism, with 15, 10, and 47 genes, respectively. Gene duplication and diversification therefore appear to have largely contributed to the battery of A. pism salivary proteins. Accordingly, SG-up and SG-expressed effector sets were enriched in duplicated genes. Aphididae-specific duplications were also present, but in lower numbers, in the D. noxia genome for CRP and apN families, as well as in A. glycines for the CRP. Interestingly, clade 4 of the apN family, which contains a majority of candidate A. pism SG-up effector genes, did not comprise any A. glycines members. Further analyses may indicate whether the loss happened in an ancestor of the Aphidini subtribe or only in A. glycines (clade 4 is present in Aphidomorpha) or resulted from technical artifacts. Positive selection was detected on specific duplicated copies in D. noxia, but a tissue specific expression analysis or saliva proteomics is required to check whether they are SG specialized potential effectors.

Effector genes are often aggregated in large clusters in the genome of plant pathogens (e.g., CRN and RXLR effectors of the oomycete Phytophthora infestans, Haas et al. 2009). These clusters are thought to result from non-allelic homologous recombination and tandem duplications (facilitated in repeat-rich genomic regions), generally associated with rapid birth and death evolution (Jiang et al. 2008; Haas et al. 2009). Effector gene clusters have also been observed in a few herbivore insects, including the gall midge Mayetiola destructor, which shows a massive expansion of effectors organized in clusters (Zhao et al. 2015). Although some members of each of the three A. pism gene families are clustered on the same genomic scaffolds (Supplementary table S6, Supplementary Material online), we were not able to identify clear gene clusters resulting from tandem duplication events, possibly due to
Evolution and Gene Family Expansions of Aphid Salivary Effectors

The functions of the CRP, ACE and apN gene families in insects, including aphids, are largely unknown. The Aphididae-specific CRP gene family was first described by Guo et al. (2014), who demonstrated that CRP-13 expression in A. pisum was induced by feeding on plant in comparison to feeding on an artificial diet. However, silencing of CRP-13 did not affect aphid survival on host plant. Proteins with numerous cysteine residues, like those of the CRP family, were previously characterized for their antifungal activities in different insect species (Banzet et al. 2002). Interestingly, some small secreted cysteine-rich proteins act as effectors in the interaction between the Asian soybean rust fungus, Phakopsora pachyrhizi, and its host plant (Qi et al. 2016). One of them was shown to suppress plant immunity by interacting with a soybean transcription factor essential for negative regulation of immunity. Thus, the various cysteine-rich proteins appear to be active in different systems including plant-pathogen interactions, so the aphid-specific CRP family may play a role in aphid nutrition once injected into the host plant.

The A. pisum ACE1 and ACE2 genes were previously reported to contribute to aphid growth on plants, but not on artificial diet (Wang et al. 2015). As the encoded proteins are predicted to have protease functions, their involvement in cleavage of certain plant defense proteins or signaling components is speculated (Wang et al. 2015). Aminopeptidases have been found in aphid guts and reported as involved in digestion (Rahbé et al. 1995; Cristofoletti et al. 2003) or in virus binding (Linz et al. 2015), but little is known about their function in aphid saliva. Furch et al. (2015) showed that A. pisum saliva was able to degrade a major phloem protein 1 (PP1) in vitro. Because PP1 is involved in protein deposition on sieve plates after severe metabolic disturbance (Gaupels et al. 2008), its degradation by aphid saliva suggests that proteases in aphid saliva degrade PP1 to prevent sieve-element occlusion triggered by aphid feeding (Furch et al. 2015). In all cases, protease function relies on peptide processing activity ensured by the M2 peptidase domain. Despite the detection of positive selection in several A. pisum M1 and M2 peptidase genes, the active site HEXXH is conserved in most of the members, indicating a functional cleaving activity. Some copies may have lost active sites or domains, particularly in the A. pisum apN family members that were too divergent to be clustered, reflecting possible functional changes after high diversification.

Gene family expansion was previously described in A.pisum, particularly in cathepsin B gut proteases (Rispe et al. 2007), chemoreceptors (Smadja et al. 2009), small RNAs machinery (Jaubert-Possamai et al. 2010), amino acid transporters (Price et al. 2011) and cuticular proteins (The International Aphid Genomics Consortium 2010). In some cases, fast evolution and positive selection occurred on recently duplicated genes (Rispe et al. 2007; Smadja et al. 2009). The expansion of salivary gene families reported here in A. pisum and subsequent putative positive selection on some gene copies highlight the importance of gene duplication in adaptation in the A. pisum ancestry. Notably, A. pisum constitutes a complex that comes from the recent adaptive radiation of least 15 biotypes (races or cryptic species), each specialized to one or few related plant species within the Fabaceae (legume) family (Peccoud et al. 2009a, 2009b; Nouhaud et al. 2014; Peccoud et al. 2015). Population genomics analyses on A. pisum biotypes have highlighted salivary and chemosensory genes as likely contributing to host-specific adaptation in the A. pisum complex (Jaquiéry et al. 2012; Smadja et al. 2012; Nouhaud et al. 2014; Duvaux et al. 2015; Eyres et al. 2017). We may relate these results to the existence of many SG-up effector candidates that have undergone duplications and episodic positive selection in the A. pisum lineage. A large repertoire of effector genes might have increased the chance of adaptive mutations selected during biotype formation, reflecting evolutionary patterns seen in other parasites’ effectors (Stergiopoulos et al. 2012; Zhao et al. 2015).

Strikingly, we found only one member of each CRP and apN (clade 4) family in the genome of M. persicae, consistent with the overall lower rate of gene expansion compared with the A. pisum lineage (Mathers et al. 2017). M. persicae is perhaps the most generalist aphid known, being able to colonize hundreds of plant species across 40 families (Blackman and Eastop 2000). Monitoring of gene expression during host switches of M. persicae laboratory lineages have suggested that acclimation to different plant species was enabled by a rapid transcriptional plasticity of duplicated genes (Mathers et al. 2017). By contrast, very little changes in gene expression have been measured in A. pisum lineages when shifted from one suitable legume plant to another (Eyres et al. 2016). Therefore, while large gene families, including salivary effectors, may be key in the rapid diversification of specialized A. pisum on various plant species, transcriptional plasticity may enable ecological generalism in M. persicae. These hypotheses require further investigation. In particular, comparative genomics using more species of the M. persicae and A. pisum lineages in combination with functional analyses of candidate effectors, would help to precisely date the origins of genetic changes and to establish more robust correlations between these changes and ecological shifts.

Supplementary Material
Supplementary data are available at Genome Biology and Evolution online.

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