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# Convergent origin and accelerated evolution of vesicle-associated RhoGAP proteins in two unrelated parasitoid wasps

### 6

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### 21

## 22 Abstract

23 Animal venoms and other protein-based secretions that perform a variety of functions, from predation 24 to defense, are highly complex cocktails of bioactive compounds. Gene duplication, accompanied by 25 modification of the expression and/or function of one of the duplicates under the action of positive 26 selection, followed by further duplication to produce multigene families of toxins is a well-documented 27 process in venomous animals. This evolutionary model has been less described in parasitoid wasps, 28 which use maternal fluids, including venom, to protect their eggs from encapsulation by the host 29 immune system. Here, we evidence the convergent recruitment and accelerated evolution of two 30 multigene families of RhoGAPs presumably involved in virulence in two unrelated parasitoid wasp 31 species, Leptopilina boulardi (Figitidae) and Venturia canescens (Icheumonidae). In both species, these 32 RhoGAPs are associated with vesicles that act as transport systems to deliver virulence factors, but are 33 produced in different tissues: the venom gland in Leptopilina sp. and the ovarian calyx in V. canescens. 34 We show that the gene encoding the cellular RacGAP1 is at the origin of the virulent RhoGAP families 35 found in Leptopilina sp. and V. canescens. We also show that both RhoGAP families have undergone 36 evolution under positive selection and that almost all of these RhoGAPs lost their GAP activity and 37 GTPase binding ability due to mutations in key amino acids. These results suggest an accelerated 38 evolution and functional diversification of these vesicle-associated RhoGAPs in the two phylogenetically 39 distant parasitoid species. The potential new function(s) and the exact mechanism of action of these 40 proteins in host cells remain to be elucidated.

41

42 Keywords: parasitoids, venom and other secretions, RhoGAP multigene family, evolution, gene

43 duplication, positive selection

### 44

### Introduction

45 Gene duplication is recognized as an important evolutionary process because it drives functional 46 novelty (Chen et al., 2013; Long et al., 2013). In particular, the formation of multigene families by repeated 47 gene duplication is a widely studied evolutionary process in several groups of venomous animals (Fry et al., 48 2009; Wong & Belov 2012; Casewell et al., 2013), but also in some parasites (Akhter et al., 2012; Arisue et 49 al., 2020). Such repeated duplication events of venomous or virulence protein-coding genes are often 50 accompanied by significant copy divergence through positive selection, allowing the acquisition of new 51 functions. In this work, we studied the process of accelerated evolution by duplication and divergence of 52 two multigene families encoding RhoGAPs presumably involved in virulence in two phylogenetically distant 53 parasitoid wasp species, Leptopilina species of the family Figitidae and Venturia canescens of the family 54 Icheumonidae.

55 The development of parasitoid wasps occurs at the expense of another arthropod, whose tissues are 56 consumed by the parasitoid larvae, usually resulting in the host death (Godfray 1994). For endoparasitoids 57 that lay eggs inside the host body, the host immune defense against parasitoids is usually based on the 58 formation of a multicellular melanized capsule around the parasitoid egg, resulting in its death (Carton et 59 al., 2008). To escape encapsulation, parasitoids have evolved several strategies, the main one being the 60 injection of maternal fluids together with the egg into the host at the time of oviposition (Pennacchio & 61 Strand 2006; Poirié et al., 2009). These maternal fluids contain (i) proteins synthesized in the venom glands 62 (Asgari & Rivers 2010; Poirié et al., 2014), some of which, as in Leptopilina, can be associated with 63 extracellular vesicles called venosomes and allow their transport to the targeted immune cells (Gatti et al., 64 2012; Wan et al., 2019), (ii) proteins synthesized in the ovarian calyx cells and associated with particles of 65 viral origin devoid of nucleic acid and named virus-like particles (VLPs) as in V. canescens (Reineke et al., 66 2006; Gatti et al., 2012; Pichon et al., 2015), and (iii) polydnaviruses (PDVs) integrated into the parasitoid 67 genome and found in some groups of parasitoids belonging to the families Braconidae and Ichneumonidae 68 (Drezen et al., 2014). Like VLPs, PDVs are synthesized in the ovarian calyx cells and are present in the 69 ovarian fluid along with the egg. These particles are unique in that they are formed by the integrated viral 70 machinery, but carry circular double-stranded DNA molecules that contain virulence genes that will be 71 expressed in the host cells (Drezen et al., 2014). The process of accelerated evolution by duplication and 72 divergence has been described for some of these virulence genes carried by PDVs, suggesting functional 73 diversification of the proteins produced (Serbielle et al., 2008; Serbielle et al., 2012; Jancek et al., 2013).

74 Leptopilina boulardi is a parasitoid wasp of Drosophila for which two lines have been well characterized 75 (Dubuffet et al., 2009). The ISm line of *L. boulardi* is highly virulent against *Drosophila melanogaster* but is 76 unable to develop in Drosophila yakuba whereas the ISy line can develop in both Drosophila species but its 77 success depends on the resistance/susceptibility of the host (Dubuffet et al., 2009). One of the most 78 abundant proteins in the venom of the L. boulardi ISm line belongs to the Rho GTPase activating protein 79 (RhoGAP) family and has been named LbGAP (Labrosse et al., 2005; Colinet et al., 2013). A LbGAP homolog 80 named LbGAPy is also present in the venom of the L. boulardi ISy line, but in lower amounts (Colinet et al., 81 2010; Colinet et al., 2013). While RhoGAPs are usually intracellular proteins composed of several different 82 domains (Tcherkezian & Lamarche-Vane 2007), LbGAP and LGAPy contain only one RhoGAP domain 83 preceded by a signal peptide that allows its secretion. LbGAP has been shown to be associated with and 84 transported by venosomes to target host hemocytes (Wan et al., 2019). LbGAP and LbGAPy specifically 85 interact with and inactivate two Drosophila Rho GTPases, Rac1 and Rac2 (Colinet et al., 2007; Colinet et al., 86 2010), which are required for hemocyte proliferation in response to parasitism, hemocyte adhesion around 87 the parasitoid egg, or the formation of intercellular junctions necessary for the capsule formation (Williams 88 et al., 2005; Williams et al., 2006). Combined transcriptomic and proteomic analyses have identified eight 89 additional RhoGAP domain-containing venom proteins in addition to LbGAP and LbGAPy for both L. 90 boulardi lines, suggesting that a multigene family has derived from repeated duplication (Colinet et al., 91 2013). One of these venom RhoGAPs, LbGAP2, has also been shown to be associated with venosomes in L. 92 boulardi ISm and to be released with LbGAP in host hemocytes (Wan et al., 2019). Interestingly, three 93 RhoGAP domain-containing venom proteins have also been identified in the closely related species 94 Leptopilina heterotoma (Colinet et al., 2013), suggesting that the recruitment of RhoGAPs in the venom 95 arsenal may have occurred before the separation of the two species.

96 Leptopilina venom RhoGAPs are not the only example of the possible use of a protein from this family 97 in parasitoid virulence (Reineke et al., 2006; Du et al., 2020). In the Ichneumonidae V. canescens, a RhoGAP 98 domain-containing protein known as VLP2 (named VcVLP2 in this paper) was found to be associated with 99 VLPs formed in the nucleus of ovarian calyx cells (Reineke et al., 2006; Pichon et al., 2015). VLPs, which 100 package proteic virulence factors wrapped into viral envelopes, are then released into the ovarian lumen 101 to associate with eggs and protect them from the host immune response by as yet unknown mechanisms 102 (Feddersen et al., 1986). These observations suggest convergent recruitment of proteins belonging to the 103 same family and injected into the host with the egg in two phylogenetically distant parasitoid species.

104 In this work, we showed that VcVLP2 in V. canescens, like LbGAP in Leptopilina, is not unique but is part 105 of a multigene family of virulent RhoGAPs associated with extracellular vesicles. RhoGAPs from different 106 organisms are grouped into distinct subfamilies based on similarities in RhoGAP domain sequence and 107 overall multi-domain organization (Tcherkezian & Lamarche-Vane 2007). Our analyses indicate that an 108 independent duplication of the RacGAP1 gene, a member of the large RhoGAP family, was the origin of the 109 virulent RhoGAP multigene families found in Leptopilina and V. canescens. We then performed 110 comparative analyses to understand the duplication events at the origin of these two multigene families. 111 Finally, we demonstrated evolution under positive selection for both RhoGAP multigene families in L. 112 boulardi and V. canescens, suggesting accelerated evolution and functional diversification.

113

### Methods

### 114 Biological material

115 The origin of the *L. boulardi* isofemale lines ISm (Lbm; Gif stock number 431) and ISy (Lby; Gif stock 116 number 486) has been described previously (Dupas et al., 1998). Briefly, the ISy and ISm founding females 117 were collected in Brazzaville (Congo) and Nasrallah (Tunisia), respectively. The L. heterotoma strain (Lh; GIF 118 stock number 548) was collected in southern France (Gotheron). The Japanese strain of 119 Leptopilina victoriae (Lv), described in Novković et al., 2011, was provided by Pr. M. T. Kimura (Hokkaido 120 University, Japan). All parasitoid lines were reared on a susceptible D. melanogaster Nasrallah strain (Gif 121 stock number 1333) at 25 °C. After emergence, the wasps were maintained at 20 °C on agar medium 122 supplemented with honey. All experiments were performed on 5- to 10-day-old parasitoid females.

### 123 Phylogeny of members of the Cynipoidea superfamily

The phylogeny of selected members of the Cynipoidea superfamily was constructed using internal transcribed spacer 2 (ITS2) sequences available at NCBI (Supplementary Table S1). Multiple sequence alignment was performed using MAFFT with the --auto option (Katoh & Standley 2013). Poorly aligned regions were removed using trimal with the -automated1 option (Capella-Gutierrez et al., 2009). Phylogenetic analysis was performed using maximum likelihood (ML) with IQ-TREE (Minh et al., 2020).

### 129 Search for candidate venom RhoGAPs in Leptopilina transcriptomes

130 L. victoriae venom apparatus, corresponding to venom glands and associated reservoirs, were 131 dissected in Ringer's saline (KCl 182 mM; NaCl 46 mM; CaCl<sub>2</sub> 3 mM; Tris-HCl 10 mM). Total RNA was 132 extracted from 100 venom apparatus using TRIzol reagent (Invitrogen) followed by RNeasy Plus Micro Kit 133 (QIAGEN) according to the manufacturers' instructions. The quality of total RNA was checked using an 134 Agilent BioAnalyzer. Illumina RNASeq sequencing (HiSeq 2000, 2 × 75 pb) and trimming were performed 135 by Beckman Coulter Genomics. The raw data are available at NCBI under the BioProject ID PRJNA974978. 136 Sequence assembly was performed using the "RNASeq de novo assembly and abundance estimation with 137 Trinity and cdhit" workflow available on Galaxy at the BIPAA platform (https://bipaa.genouest.org).

Leptopilina clavipes (whole body), Ganaspis sp. G1 (female abdomen), Andricus quercuscalicis (whole body) and Synergus umbraculus (whole body) transcriptomes were obtained from NCBI under accession numbers GAXY00000000, GAIW00000000, GBNY00000000 and GBWA000000000, respectively. Coding regions in the transcripts were identified and translated using TransDecoder (Haas et al., 2013), which is available on the BIPAA Galaxy platform. Searches for RhoGAP domain-containing sequences in translated coding sequences were performed using hmmsearch from the HMMER package (Eddy 2009) with the RhoGAP (PF00620) HMM profile.

### 145 Completion of *L. boulardi* and *L. heterotoma* venom RhoGAP sequences

146 Venom apparatus, corresponding to venom glands and associated reservoirs, were dissected in Ringer's 147 saline. Total RNA was extracted from venom apparatus using the RNeasy Plus Micro Kit (QIAGEN) according 148 to the manufacturers' instructions. To obtain the full-length coding sequence of LbGAP1.3, LbGAP5, 149 LbGAPy6, LhGAP1 and LhGAP2, rapid amplification of cDNA ends (RACE) was performed using the SMART 150 RACE cDNA Amplification Kit (Clontech). For LhGAP3, 3' RACE could not be applied due to the presence of 151 poly(A) stretches in the sequence. LhGAP3-matching sequences were searched for in the transcriptome 152 assembly obtained by Goecks et al., 2013) from the abdomen of *L. heterotoma* females (GenBank accession 153 number GAJC00000000) using the command line NCBI-BLAST package (version 2.2.24). The resulting 154 complete LhGAP3 sequence was then verified by RT-PCR using the iScript cDNA Synthesis Kit (BioRad) and 155 the GoTaq DNA Polymerase (Promega), followed by direct sequencing of the amplified fragment. Geneious 156 software (Biomatters) was used for sequence editing and assembly.

### 157 Obtention of Leptopilina RacGAP1 coding sequences

158 A combination of RT-PCR and RACE using conserved and specific primers was used to clone the full-159 length ORF sequence of L. boulardi and L. heterotoma RacGAP1. The amino acid sequence of Nasonia 160 vitripennis RacGAP1 (Supplementary Table S2) was used to search for similar sequences among 161 Hymenoptera species using BLASTP at NCBI (http://www.ncbi.nlm.nih.gov/blast/). A multiple sequence 162 alignment of the RacGAP1 amino acid sequences found in Hymenoptera was then performed using 163 MUSCLE (Edgar 2004). Two pairs of RacGAP1-specific degenerate primers were designed from the 164 identified conserved regions (Supplementary Figure S1). Total RNA was extracted from Lbm and Lh 165 individuals using the TRIzol reagent (Invitrogen), and RT-PCR experiments were performed using the 166 RacGAP1-specific degenerate primers. After direct sequencing of the amplified fragments, Lbm- and Lh-167 specific primers were designed to complete the sequences obtained by RT-PCR and RACE (Supplementary 168 Figure S1). The coding sequences of *L. victoriae and L. clavipes* RacGAP1 were obtained by BLAST searches 169 in the corresponding transcriptomes using *L. boulardi* and *L. heterotoma* RacGAP1 sequences as queries.

### 170 Obtention of the genomic sequences of Leptopilina venom RhoGAPs and RacGAP1

The genomic sequences of *Leptopilina* venom RhoGAPs and RacGAP1 were obtained by BLAST and Exonerate searches (Slater & Birney 2005) using the corresponding coding sequences as queries. *L. boulardi* ISm, *L. boulardi* ISy, *L. heterotoma* and *L. clavipes* genome assemblies were obtained from NCBI under accession numbers GCA\_011634795.1, GCA\_019393585.1, GCF\_015476425.1 and GCA\_001855655.1 respectively.

### 176 Search for candidate V. canescens RhoGAP calyx sequences

Coding regions were identified from the *V. canescens* calyx transcriptome published by Pichon et al., and translated into protein sequences using TransDecoder (Haas et al., 2013). The search for RhoGAP domain-containing sequences was performed using hmmsearch from the HMMER package (Eddy with the RhoGAP (PF00620) HMM profile. The identified RhoGAP calyx sequences were used as a database for Mascot to explore MS/MS data obtained from purified *V. canescens* VLPs (Pichon et al., 2015) with a false discovery rate of 1%. Only proteins identified by two or more peptides were considered.

- 183 The *V. canescens* RacGAP1 coding sequence was obtained by BLAST searches in the *V. canescens* calyx 184 transcriptome using the *N. vitripennis* RacGAP1 sequence as query.
- The *V. canescens* calyx RhoGAP and RacGAP1 genome sequences were obtained by BLAST and
   Exonerate searches (Slater and Birney 2005) using the corresponding coding sequences as queries. The *V. canescens* genome assembly was obtained from NCBI under accession number GCA\_019457755.1.

### 188 Identification of domains and motifs and prediction of subcellular localization

189 The presence and position of signal peptide cleavage sites in the identified RhoGAP domain-containing 190 sequences were predicted using the SignalP server 191 (https://services.healthtech.dtu.dk/service.php?SignalP). The searches for domains and motifs were 192 performed using InterProScan 5 (Jones et al., 2014). Coiled-coil regions were predicted using COILS (Lucas 193 et al., 1991).

Prediction of protein subcellular localization and sorting signals for *Venturia* RacGAP1 and calyx
 RhoGAPs was performed using the DeepLoc2 server (https://services.healthtech.dtu.dk/services/DeepLoc 2.0/).

197 Nuclear localization for VcRacGAP1 and calyx RhoGAPs was also predicted using the following three 198 tools: NucPred (Brameier et al., 2007) was used to indicate whether a protein spends part of its time in the 199 nucleus (https://nucpred.bioinfo.se/nucpred/), LocTree3 (Goldberg et al., 2014) to provide subcellular 200 localization and Gene Ontology terms (https://rostlab.org/services/loctree3/), and PSORT II (Nakai & 201 Horton 1999) to detect sorting signals and subcellular localization (https://psort.hgc.jp/form2.html). 202 PSORT II results corresponding to ER membrane retention signals and cleavage sites for mitochondrial 203 presequences were included as they may be associated with nuclear localization. Mitochondrial 204 presequences have no sequence homology but possess physical characteristics that allow them to interact 205 with the outer membranes of mitochondria and thus allow the targeting of proteins to the mitochondrial 206 matrix. Regarding mitochondrial targeting, it has been shown from plants to humans that some proteins 207 with mitochondrial presequences are dually targeted to mitochondria and the nucleus (Millar et al., 2006; 208 Mueller et al., 2004). Indeed, anchoring signals to the ER membrane are known to allow baculoviral 209 proteins to migrate to the inner nuclear membrane (Braunagel et al., 1996). Proteins forming VLPs in V. 210 canescens could thus follow the same pathways as their homologous proteins in baculoviruses, nudiviruses, 211 hytrosaviruses, and bracoviruses due to the conservation of this mechanism among these viruses belonging 212 to the order Lefavirales (Braunagel et al., 1996; Hong et al., 1997; Braunagel et al., 2004; Abd-Alla et al., 213 2008; Bézier et al., 2009; Braunagel et al., 2009).

### 214 Phylogeny of Leptopilina venom and Venturia calyx RhoGAPs

215 Searches for *N. vitripennis* RhoGAP sequences were performed using BLASTP at NCBI 216 (http://www.ncbi.nlm.nih.gov/blast/) and hmmsearch from the HMMER package (Eddy 2009) with the 217 RhoGAP (PF00620) HMM profile on the *N. vitripennis* v1.2 proteome database (Rago et al., 2016).

218 Multiple amino acid sequence alignments were performed using MAFFT with the --auto option (Katoh 219 and Standley 2013). For codon-based analysis of selection (see below), codon-based alignments of 220 complete coding sequences were performed using RevTrans 221 (https://services.healthtech.dtu.dk/service.php?RevTrans) with the amino acid alignments as templates. 222 Poorly aligned regions were removed using trimAl with the -automated1 option (Capella-Gutierrez et al., 223 2009). Phylogenetic analyses were performed using maximum likelihood (ML) with IQ-TREE (Minh et al., 224 2020). ModelFinder was used to select the best model selection for phylogeny (Kalyaanamoorthy et al., 225 2017).

### 226 Codon-based analysis of selection

Codon-based alignments of complete coding sequences were performed with RevTrans using the
 amino acid alignments as templates (see above). Phylogenetic analyses were performed using maximum
 likelihood (ML) with PhyML (Guindon et al., 2010). Smart Model Selection was used to choose the best
 model selection for the phylogeny (Lefort et al., 2017).

To detect evolutionary selective pressures acting on RhoGAP sequences, the ratios of non-synonymous substitutions (dN) to synonymous substitutions (dS) were compared using different ML frameworks: the CODEML program in the PAML (Phylogenetic Analysis by Maximum Likelihood) package (Yang 1997; Yang 2007), the HyPhy software implemented at http://www.datamonkey.org/ (Delport et al., 2010), and the Selecton server (Stern et al., 2007) available at http://selecton.tau.ac.il/index.html.

236 Five different methods were used to detect codons under positive selection. In the first, codon 237 substitution models implemented in CodeML were applied to the codon-based alignment using the F3x4 238 frequency model. Two pairs of site models were used to determine whether some codons were under 239 positive selection: M1a (neutral) versus M2a (selection) and M7 ( $\beta$ ) versus M8 ( $\beta$  &  $\omega$ ). The models were 240 compared using a likelihood ratio test (LRT) with 2 degrees of freedom to assess the significance of 241 detection of selection (Yang et al., 2000). Bayes Empirical Bayes (BEB) inference was then used to identify 242 amino acid sites with a posterior probability >95% of being under positive selection (Yang et al., 2005). We 243 then applied four other methods of detection of selection available in the HyPhy package: Single Likelihood 244 Ancestor Counting (SLAC), Fixed Effect Likelihood (FEL), Mixed Effects Model of Evolution (MEME) and Fast 245 Unbiased Bayesian AppRoximation (FUBAR) methods (Kosakovsky Pond & Frost 2005; Murrell et al., 2012;

Murrell et al., 2013). To eliminate false-positive detection, only codons identified by CodeML M2a and M8 and at least one of the other methods were considered under positive selection. Radical or conservative replacements were then determined based on whether they involved a change in the physicochemical properties of a given amino acid, such as charge or polarity (Zhang 2000).

Four different methods were used to detect codons under negative selection: the Selecton server using the pair of site models M8a ( $\beta\&$   $\omega$ =1) versus M8 ( $\beta\&$   $\omega$ ), and the FEL, SLAC and FUBAR methods mentioned above. Only codons identified by Selecton and at least one other method were considered under purifying selection.

To identify specific lineages with a proportion of sites evolving under positive or purifying selection, we performed branch-site REL analyses using the HyPhy package (Kosakovsky Pond et al., 2011). Unlike the branch and branch-site lineage-specific models available in CodeML, branch-site REL does not require *a priori* identification of foreground and background branches.

### 258 Structural analysis

259 Molecular modeling of VcVLP2 RhoGAP was performed using the Phyre server with default parameters 260 (http://www.sbg.bio.ic.ac.uk/phyre2/) (Kelley et al., 2015). The model with the highest confidence (100%) 261 and coverage (46%) was obtained using the crystal structure with PDB code 2OVJ as a template. Model 262 quality was evaluated using the QMEAN server (http://swissmodel.expasy.org/qmean/) (Benkert et al., 263 2011). Briefly, the QMEAN score is a global reliability score with values ranging between 0 (lower accuracy) 264 and 1 (higher accuracy). The associated Z-score relates this QMEAN score to the scores of a non-redundant 265 set of high-resolution X-ray structures of similar size, with ideal values close to 0. Visualization of LbGAP 266 (Colinet et al., 2007) and VcVLP2 structures and mapping of sites under selection were performed using 267 PyMol (http://sourceforge.net/projects/pymol/). Secondary structure assignment was performed with the 268 DSSP program (Kabsch & Sander 1983). Accessible surface area (ASA) or solvent accessibility of amino acids 269 was predicted using the ASAView algorithm (Ahmad et al., 2004).

### 270 In vitro mutagenesis

The S76K, V124K, L137D, T143L, S150L, I185K, E200G, and V203R mutations were introduced into the
 LbGAP cDNA using the QuickChange XL Site-Directed Mutagenesis Kit (Stratagene). The results of *in vitro* mutagenesis were verified by sequencing.

### 274 Yeast two-hybrid analysis

275 Interactions between LbGAP and LbGAP mutants and mutated forms of Drosophila RhoA, Rac1, Rac2 276 and Cdc42 GTPases were individually examined by mating as previously described (Colinet et al., 2007). 277 The plasmids expressing the GTPase proteins were tested against the pGADT7-T control vector, which 278 encodes a fusion between the GAL4 activation domain and the SV40 large T-antigen. Reciprocally, plasmids 279 producing LbGAP and LbGAP mutants were tested against the pLex-Lamin control vector. Interactions 280 between LbGAP and the Rac1 and Rac2 GTPases were used as positive controls (Colinet et al., 2007). 281 Interactions were initially tested by spotting five-fold serial dilutions of cells on minimal medium lacking 282 histidine and supplemented with 3-amino-triazole at 0.5 mM to reduce the number of false positives. ß-283 galactosidase activity was then detected on plates (Fromont-Racine et al., 1997).

### 284 Purification of Leptopilina venosomes and mass spectrometry analysis

285 Twenty-five Lbm, Lby and Lh female venom apparatus were dissected, and the reservoirs were 286 separated from the gland and opened to release the venom content in 25  $\mu$ l of Ringer's solution 287 supplemented with a protease inhibitor cocktail (Sigma). The venom suspension was centrifuged at 500xg 288 for 5 min to remove the residual tissues, then centrifuged at 15,000xg for 10 min to separate the vesicular 289 fraction from the soluble venom proteins (supernatant fraction) (Wan et al., 2019). The vesicular fraction 290 was then washed twice by resuspension in 25  $\mu$ l of Ringer's solution followed by centrifugation at 15,000xg 291 for 10 min. The two samples were mixed with 4x Laemmli buffer containing  $\beta$ -mercaptoethanol (v/v) and 292 boiled for 5 min. Proteins were then separated on a 6–16% linear gradient SDS-PAGE and the gel was silver 293 stained as previously described (Colinet et al., 2013). Identification of proteins by nano-LC-tandem mass 294 spectrometry (MS/MS) was performed on bands excised from the gels as previously described (Colinet et 295 al., 2013). MS/MS data analysis was performed with Mascot software (http://www.matrixscience.com),

licensed in house using the full-length coding sequences of the *Leptopilina* venom RhoGAP sequences. Data
validation criteria were (i) one peptide with individual ion score greater than 50 or (ii) at least two peptides
of individual ion score greater than 20. The mascot score is calculated as -10Log (p). The calculated FDR
(based on an automated decoy database search) was less than 1%. The mass spectrometry proteomics
data have been deposited to the ProteomeXchange Consortium via the PRIDE (Perez-Riverol et al., 2022)
partner repository with the dataset identifier PXD041695.

### 302

### Results

### 303 Leptopilina venom RhoGAPs probably emerged in the ancestor of the genus

304 In the Figitidae, venom RhoGAPs have only been described for L. boulardi and L. heterotoma (Labrosse 305 et al., 2005; Colinet et al., 2013). A search for candidate homologs was performed in the transcriptomes of 306 L. victoriae and L. clavipes, two other Leptopilina species, Gasnaspis sp. G1, another member of the family 307 Figitidae, and Andricus quercuscalicis and Synergus umbraculus, two gall wasp species representatives of 308 the superfamily Cynipoidea outside of Figitidae (separated by approximately 50 Mya (Peters et al., 2017)). 309 The putative venom RhoGAPs were identified based on the presence of a signal peptide at the N-terminus 310 of the protein, followed by a RhoGAP domain (Table 1). Two transcript sequences encoding RhoGAP 311 domain-containing proteins predicted to be secreted were found in L. victoriae (LvGAP1 and LvGAP2) and 312 one in L. clavipes (LcGAP1), whereas none were found for Gasnaspis sp. G1, A. quercuscalicis and S. 313 umbraculus (Table 1 and Figure 1A). Our results therefore suggest that venom RhoGAPs are specific to the 314 genus Leptopilina although further taxon sampling would be required to fully support this hypothesis.

### 315 Leptopilina venom RhoGAPs likely evolved from a single imperfect duplication of RacGAP1

316 A phylogenetic analysis was performed to learn more about the evolutionary history of Leptopilina 317 venom RhoGAPs. The phylogeny was constructed based on the amino acid sequence of the RhoGAP 318 domain of these venom proteins and that of the 19 classical RhoGAPs we identified from the predicted 319 proteome of the jewel wasp Nasonia vitripennis (Supplementary Table S2). Prior to analysis, the coding 320 sequence was completed for two Lbm (LbGAP1.3 and LbGAP5), one Lby (LbGAPy6) and all three Lh venom 321 RhoGAPs. Two of the Leptopiling venom RhoGAPs, namely LhGAP3 and LcGAP1, were predicted to contain 322 two RhoGAP domains instead of one. Therefore, these two domains were separated for the analysis. The 323 resulting phylogeny identified NvRacGAP1 as the closest RhoGAP from N. vitripennis to all Leptopilina 324 venom RhoGAPs with confident support values (Figure 1B). Accordingly, the RhoGAP domain found in all Leptopilina venom RhoGAPs was predicted to belong to the MgcRacGAP subfamily typically found in 325 326 RacGAP1-related proteins (Table 1).

327 The RacGAP1 coding sequence of L. boulardi, L. heterotoma, L. victoriae and L. clavipes was obtained 328 either by cloning and sequencing or by searching genomic data, and its domain organization was compared 329 with that of venom RhoGAPs (Figure 1C, Table 1 and Supplementary Figure S2). Leptopilina RacGAP1 has a 330 typical domain organization consisting of a coiled-coil region and a C1 motif (protein kinase C-like zinc 331 finger motif, a ~ 50 amino-acid cysteine-rich domain involved in phorbol ester/diacylglycerol and Zn<sup>2+</sup> 332 binding) followed by a RhoGAP domain (Tcherkezian & Lamarche-Vane 2007). A large part of the RacGAP1 333 sequence spanning the coiled-coil region and the C1 motif is replaced by the signal peptide in all Leptopilina 334 venom RhoGAPs (Figure 1C, Table 1 and Supplementary Figure S2). This suggests that a unique incomplete 335 duplication of the RacGAP1 gene occurred in the ancestor of the Leptopilina genus, resulting in a truncated 336 duplicate encoding a RhoGAP with an altered N-terminus.

337 To further investigate this duplication event and the possible origin of the signal peptide, the genomic 338 sequences of L. boulardi, L. heterotoma and L. clavipes venom RhoGAPs and RACGAP1 were obtained, 339 either by cloning and sequencing or by searching in genomic data, and compared (Supplementary Table 340 S2). The genomic sequence of Leptopilina RacGAP1 consists of 10 exons with a "CC region", "C1 motif" and 341 "RhoGAP domain" encoded by exons 2 and 3, exons 5 and 6 and exons 7 and 8, respectively. The RhoGAP 342 domain is also encoded by two exons for venom RhoGAPs. These two exons are preceded by one to three 343 exons, with the signal peptide encoded by the first exon for most of them. Sequence comparisons revealed 344 similarities between venom RhoGAPs and Leptopilina RacGAP1 only for the sequence spanning the two 345 exons encoding the RhoGAP domain and part of the following one. No significant similarities were found 346 for the preceding exons at either the nucleotide or amino acid level (Supplementary Figure S3). This

347 supports the hypothesis of a partial duplication of the RacGAP1 gene in the ancestor of the *Leptopilina* 348 genus, resulting in the loss of the sequence spanning exons 1 to 6, followed by further duplication of the 349 ancestrally duplicated gene during the diversification of the genus *Leptopilina*.

350 The two consecutive RhoGAP domains found in LhGAP3 were grouped together with confident support 351 values in the phylogenies (Figure 1B and Supplementary Figure S2). This suggests that a partial tandem 352 duplication spanning exons 2 and 3 has occurred for this L. heterotoma venom RhoGAP. In contrast, the 353 two RhoGAP domains found in LcGAP1 did not group together (Figure 1B and Supplementary Figure S2). 354 However, the genomic sequence of LcGAP1 was not complete and we could not find the region 355 corresponding to the second RhoGAP domain in the L. clavipes genome (Supplementary Table S3). 356 Therefore, it is unclear whether the LcGAP1 coding sequence found in the transcriptome assembly is true 357 or artifactual.

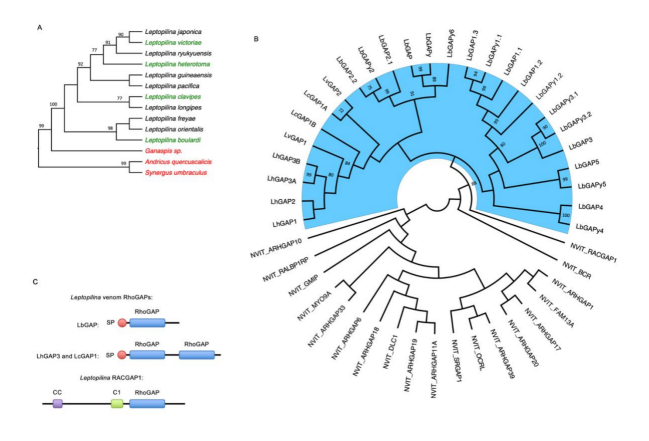
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360

361 362 **Table 1.** Motif and domain organization of *Leptopilina* venom RhoGAPs. The signal peptide was predicted using SignalP at CBS. The RhoGAP domain (PF00620) and the RacGAP1 domain (PTHR46199) were identified using InterProScan. Two successive RhoGAP domains were found in LhGAP3 and LcGAP1. Lbm: *L. boulardi* ISm ; Lby: *L. boulardi* ISy ; Lh: *L. heterotoma* ; Lc: *L. clavipes* ; Lv: *L. victoriae.* 

			Signal pe	eptide	RhoGAP domain					RacGAP1 domain			
							F00620)				HR46199	)	
		Total	From	То	From	То	Length	E-value	From	То	Length	E-value	
		length											
Lbm	LbGAP	282	1	20	52	193	142	2.9e-32	38	267	230	7.0e-37	
	LbGAP1.1	292	1	23	58	205	148	1.0e-19	42	255	214	7.0e-30	
	LbGAP1.2	280	1	23	58	205	148	1.1e-19	42	255	214	1.3e-29	
	LbGAP1.3	287	1	23	58	205	148	7.2e-17	42	253	212	1.3e-27	
	LbGAP2.1	251	1	23	44	183	140	9.9e-26	26	234	209	5.5e-37	
	LbGAP2.2	263	1	23	56	196	141	6.6e-26	49	246	198	1.3e-34	
	LbGAP3	295	1	23	56	175	120	3.2e-12	42	266	225	1.2e-27	
	LbGAP4	246	1	20	42	178	137	9.1e-11	38	218	181	5.6e-18	
	LbGAP5	275	1	20	60	207	148	6.3e-23	41	251	211	2.4e-33	
Lby	LbGAPy	286	1	20	52	193	142	1.6e-32	38	267	230	3.4e-37	
	LbGAPy1.1	287	1	23	58	205	148	5.6e-17	42	255	214	9.9e-28	
	LbGAPy1.2	325	1	23	56	203	148	2.9e-17	42	293	252	1.7e-26	
	LbGAPy2	256	1	23	44	185	142	8.3e-27	26	234	209	1.5e-37	
	LbGAPy3.1	281	1	23	56	175	120	1.8e-11	42	237	196	1.5e-25	
	LbGAPy3.2	266	1	23	41	160	120	5.8e-12	28	222	195	6.4e-25	
	LbGAPy4	274	1	20	42	177	136	7.4e-09	30	218	189	2.1e-17	
	LbGAPy5	275	1	23	60	207	148	4.4e-24	41	251	211	1.2e-35	
	LbGAPy6	239	1	20	53	194	142	8.2e-25	49	231	183	9.4e-34	
Lh	LhGAP1	293	1	20	43	180	138	5.5e-36	41	289	249	1.0e-38	
	LhGAP2	336	1	20	43	181	139	1.5e-29	45	275	231	8.2e-31	
	LhGAP3	467	1	20	35	174	140	1.4e-22	34	221	188	1.3e-38	
					257	396	140	2.2e-17					
Lc	LcGAP1	497	1	20	43	184	142	3.2e-30					
					249	384	136	3.5e-28	247	478	232	8.46e-71	
Lv	LvGAP1	370	1	20	38	179	142	2.4e-30					
	LvGAP2	266	1	20	38	181	144	8.5e-23	33	224	192	3.7e-28	

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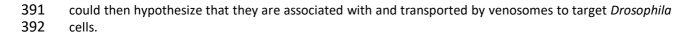


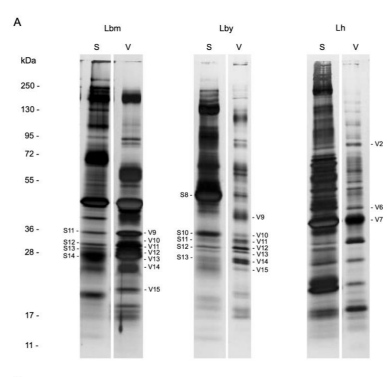


365 Figure 1. Origin of Leptopilina venom RhoGAPs (A) Maximum-likelihood phylogenetic tree of 366 selected members of the Cynipoidea superfamily. The phylogenetic tree was obtained with IQ-TREE 367 using internal transcribed spacer 2 (ITS2) sequence and displayed as a cladogram. Species in red are 368 those for which venom RhoGAPs were not found. No data were available for species in black. (B) 369 Maximum-likelihood phylogenetic tree of Leptopilina venom RhoGAPs along with N. vitripennis 370 classical RhoGAPs. The phylogenetic tree was obtained with IQ-TREE using the RhoGAP domain amino 371 acid sequence and displayed as a cladogram. For A and B, numbers at corresponding nodes are 372 bootstrap support values in percent (500 bootstrap replicates). Only bootstrap values greater than 373 70% are shown. Leptopiling venom RhoGAPs are highlighted in blue. (C) Comparison of protein 374 domain organization between LbGAP, representative of most Leptopilina venom RhoGAPs, LhGAP3 375 and LcGAP1, the two venom RhoGAPs that contain two RhoGAP domains, and Leptopilina RacGAP1. 376 C1: protein kinase C-like zinc finger motif, CC: coiled-coil region, MgcRacGAP: RhoGAP domain found 377 in RacGAP1-related proteins, SP: signal peptide.

### 378 L. boulardi and L. heterotoma venom RhoGAPs are associated with venosomes

379 LbGAP and LbGAP2 have been shown to be associated with vesicles named venosomes produced in L. 380 boulardi venom that transport them to Drosophila lamellocytes (Wan et al., 2019). Our next goal was to 381 investigate whether Leptopiling venom RhoGAPs other than LbGAP and LbGAP2 could be associated with 382 venosomes. Proteomic analysis was performed on the supernatant and vesicular fractions separated from 383 the venom by centrifugation. Comparison of the electrophoretic profiles obtained on a 6-16% SDS-PAGE 384 for L. boulardi ISm (Lbm), L. boulardi ISy (Lby) and L. heterotoma (Lh) revealed an important variation 385 between both fractions for all three wasps (Figure 2). All the major bands on the electrophoretic patterns 386 of Lbm, Lby and Lh supernatant and vesicular fractions, as well as several minor bands (35 bands in total 387 for Lbm, 34 for Lby and 37 for Lh), were excised and tryptic peptides were analyzed by mass spectrometry. 388 The coding sequences of Leptopilina venom RhoGAPs were used to perform Mascot searches on the mass 389 spectrometry data obtained from both venom fractions. All L. boulardi and L. heterotoma venom RhoGAPs 390 were detected in the vesicular fraction, where most of them were found to be enriched (Figure 2). We





				Supernat	ant	Vesicular		
		Length	Predicted Mw	Protein band	Mascot Matches	Protein band	Mascot Matches	
	LbGAP	282	32.13	<b>\$11</b> /\$12	11	V9/V10/V11/V12/ V13/V14/V15	57	
	LbGAP1.1	292	33.02	\$11/ <b>\$12</b> /\$14	14	V9/ <b>V10</b> /V11/V12/ V14/V15	85	
	LbGAP1.2	280	31.99	\$13	3	V10/V11/V12/V13	36	
	LbGAP1.3	287	32.80	\$12/ <b>\$13</b> /\$14	12	V9/V10/ <b>V11</b> /V12/ V13/V14/V15	85	
Lbm	LbGAP2.1	251	28.81	\$11/\$12/\$13/ <b>\$14</b>	14	V10/V11/V12/ V13/V14/V15	60	
Lbm	LbGAP2.2	263	30.18	\$12/\$13/ <b>\$14</b> 9		V9/V10/V11/ <b>V12</b> / V13/V14/V15	85	
	LbGAP3	295	34.01	<b>\$12</b> /\$13	3	V10/V11/V12/V13	15	
- 1	LbGAP4	246	28.57			V13/ <b>V14</b>	3	
	LbGAP5	275	31.72			V10/V11	Mascot Matche 57 85 36 85 60 85 15	
	LbGAPy	286	32.56	<b>S8</b>	1	V10/V11	4	
	LbGAPy1.1	287	32.81	\$10/\$11/ <b>\$12</b> /\$13	19	V11/V12/V13	19	
	LbGAPy1.2	325	37.47			V13	1	
	LbGAPy2	256	29.37	<b>S8</b> /S13	8	V12/V13/V14	8	
lby	LbGAPy3.1	281	32.58	S11	1	V10/V11/V12/V13	5	
LUY	LbGAPy3.2	266	30.74	S11	1	V10/V11/V12/V13	5	
	LbGAPy4	274	31.92	S10/ <b>S11</b> /S12/S13	9	V9/V10/ <b>V11</b> /V12/ V13/V14/V15	36	
	LbGAPy5	275	32.04			V12	1	
	LbGAPy6	239	27.83			V15	3	
	LhGAP1	293	33.16			V7	13	
Lh	LhGAP2	336	38.23			V6	4	
	LhGAP3	467	53.92			V2	7	

**Figure 2. Proteomic analysis of** *Leptopilina* **venom supernatant and vesicular fractions.** (A) The supernatant (S) and vesicular (V) fractions obtained from 25 Lbm, Lby and Lh venom reservoirs were separated on a 6-16% SDS-PAGE under reducing conditions and visualized by silver staining. Protein bands in which *Leptopilina* venom RhoGAPs were identified by mass spectrometry are numbered on the gel. Molecular weight standard positions are indicated on the left (kDa). (B) For each *Leptopilina* venom RhoGAP, length (number of amino acids), predicted Mw (kDa), bands in which specific peptides were found by mass spectrometry and number of total Mascot matches according to the S and V venom fractions are given. Numbers in bold correspond to the bands in which each RhoGAP was identified as the most abundant according to the number of Mascot matches.

403 V. canescens calyx RhoGAPs probably evolved from two or more imperfect RacGAP1 duplication events

404 Analysis of the V. canescens calyx transcriptome allowed the identification of a total of 13 RhoGAP 405 domain-containing protein coding sequences, including VcVLP2 (Table 2). Matches with peptides obtained 406 from a proteomic analysis of V. canescens VLPs (Pichon et al., 2015) were detected for all 13 calyx RhoGAPs, 407 indicating that they are associated with VLPs (Table 2).

408 Similar to the Leptopilina venom RhoGAPs, the V. canescens calyx RhoGAPs contain a MgcRacGAP1 409 domain (Table 2) and are closely related to NvRacGAP1 (Figure 3A). In contrast to the Leptopilina venom 410 RhoGAPs, none of the V. canescens calyx proteins were predicted to be secreted. Most were predicted to 411 be localized in the nucleus and/or to contain a nuclear localization signal (Supplementary Tables S4 and 412 S5). The absence of a predicted nuclear localization signal for VcGAP7 may be explained by the incomplete 413 coding sequence at the 5' end (Table 2). There is greater variation in domain organization for V. canescens 414 calyx RhoGAPs compared to Leptopilina venom RhoGAPs (Figure 3B and Supplementary Figure S4). Eight 415 V. canescens calyx RhoGAPs have retained the C1 motif. In addition, VcGAP12 still contains a CC region in 416 the N-terminal part of the sequence although the total sequence length is shorter than RacGAP1 (Figure 417 3B and Supplementary Figure S4).



420

421 422 Table 2. Motif and domain organization of Venturia calyx RhoGAP proteins associated with VLPs. The coiled-coil (CC) region was predicted using COILS (Lucas et al., 1991). The protein kinase C-like zinc finger (C1) motif (PS50081), the RhoGAP domain (PF00620) and the RacGAP1 domain (PTHR46199) were identified using InterProScan. Mascot 5: number of matches with peptides from purified V. canescens VLPs. <sup>a</sup>Coding sequence not complete at 5' end. <sup>b</sup>Coding sequence not complete at 3' end.

					C2 (PS50				AP doma F00620)	in			AP1 doma HR46199)	
	Mascot	Total length	From	То	From	To	From	То	Length	E-value	From	То	Length	E-value
VcVLP2	26	485			56	106	134	277	144	7.4e-36	28	366	339	3.1e-98
VcGAP1	9	207					30	168	139	5.7e-25	12	199	188	5.1e-49
VcGAP2	19	280					23	161	139	6.9e-29	8	195	188	1.5e-50
VcGAP3	4	270			22	69	92	231	140	2.6e-25	14	265	252	2.9e-63
VcGAP4	21	382			37	86	116	240	125	2.2e-16	20	298	279	2.7e-68
VcGAP5	19	307			15	63	91	236	146	4.0e-24	7	281	275	2.6e-60
VcGAP6	16	284			17	67	88	215	128	3.3e-19	14	270	257	8.0e-55
VcGAP7	14	225ª					15	156	142	5.3e-29	4	212	209	9.2e-57
VcGAP8	5	369			64	113	141	281	141	4.6e-28	20	334	315	3.6e-84
VcGAP9	9	319 <sup>b</sup>			54	104	123	266	144	1.1e-28	38	306	269	5.5e-73
VcGAP10	5	233					30	166	137	1.6e-22	7	219	213	7.1e-45
VcGAP11	5	199					27	166	140	4.3e-19	8	198	191	5.7e-42
VcGAP12	8	402	21	48	79	131	159	296	138	3.3e-35	54	372	319	9.7e-91

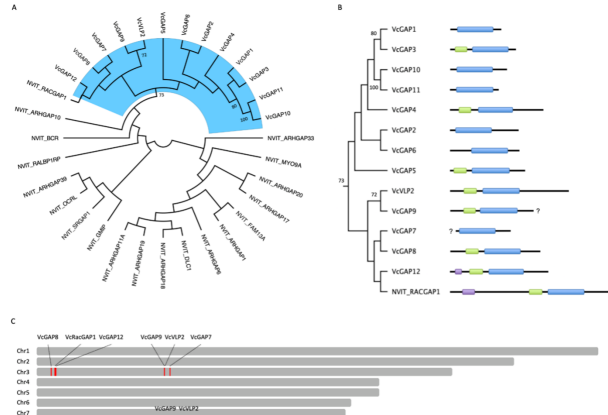
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424 The genomic coding sequence of VcRacGAP1 was obtained by genomic data mining. It comprised 9 425 exons with the CC region, C1 motif and RhoGAP domain encoded by exons 2 and 3, exons 5 and 6 and exon 426 7, respectively (Supplementary Table S6). In contrast to Leptopilina, the RhoGAP domain is encoded by a 427 single exon in V. canescens RacGAP1 as well as in all calyx RhoGAPs. Interestingly, the VcGAP12 gene is 428 located close to the VcRacGAP1 gene on chromosome 3, but is three exons shorter, suggesting a recent 429 incomplete duplication (Supplementary Table S6). In contrast, VcVLP2 and the other VcGAP genes are more 430 widely distributed in the V. canescens genome (Supplementary Table S6). It is therefore possible that 431 incomplete duplication events of the VcRacGAP1 gene, followed by further tandem and dispersed

### 432 duplication of the ancestrally duplicated genes, led to the current number of calyx RhoGAPs in V. canescens

433 organized in several clusters comprising 2 or more genes in tandem arrays (Figure 3C).

### 434



Chr7	VCGAP9 VCVLP2					
Chr8	$\setminus$ /		j i i i i i i i i i i i i i i i i i i i			
Chr9	$\sim$					
Chr10						
Chr11						
	SMBp 10MBp 15MBp VCGAP5 VCGAP2 VCGAP6 VCGAP4 VCGAP3 VCGAP1	20Mbp	25Mbp	30Mbp	35Mbp	40Mbp

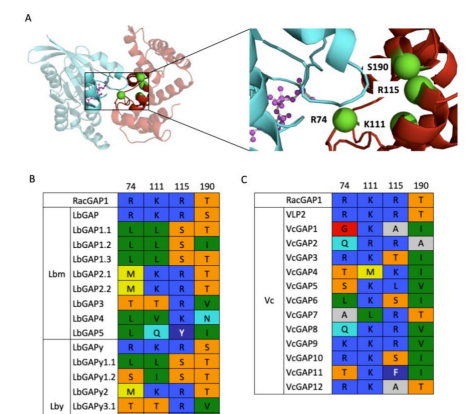
### 435

436 Figure 3. Origin of V. canescens calyx RhoGAPs (A) Maximum-likelihood phylogenetic tree of V. 437 canescens calyx RhoGAPs along with N. vitripennis classical RhoGAPs. The phylogenetic tree was 438 obtained with IQ-TREE using the RhoGAP domain amino acid sequence and displayed as a cladogram. 439 Numbers at corresponding nodes are bootstrap support values in percent (500 bootstrap replicates). 440 Only bootstrap values greater than 70% are shown. V. canescens calyx RhoGAPs are highlighted in 441 blue. (B) Comparison of protein domain organization between V. canescens calyx RhoGAPs and N. 442 vitripennis RacGAP1 using the RhoGAP domain as reference. The color code is as follows: purple for 443 the coiled-coil region (CC), green for the protein kinase C-like zinc finger motif (C1) and blue for the 444 RhoGAP domain. A (?) indicates that either the N- or C-terminal part of the sequence is unknown. (C) 445 V. canescens chromosome map with the position of gene loci corresponding to RacGAP1 and calyx 446 RhoGAPs visualized using chromoMap R package (Anand & Rodriguez Lopez 2022).

447

### 448 Evidence of positive selection in L. boulardi venom and V. canescens calyx RhoGAP sequences

449 In a previous work, we identified four amino acid residues involved in the interaction of LbGAP with 450 Rho GTPases (Colinet et al., 2007), including the key arginine residue (R74 in LbGAP) required for the GAP 451 catalytic activity (Figure 4A). The other 8 venom RhoGAP sequences found in L. boulardi Ism and ISy were 452 all mutated at this arginine residue (Figure 4B). Most also contained mutations in one or more of the other 453 three amino acids involved in Rho GTPase interaction (Figure 4B). Similarly, all V. canescens calyx RhoGAP 454 sequences, except VcVLP2, are mutated in one or more of the sites essential for GAP activity and/or 455 involved in the interaction with Rho GTPases (Figure 4C). These observations suggest that only LbGAP and 456 its homolog LbGAPy in L. boulardi and VcVLP2 in V. canescens are functional as RhoGAPs.



457

458	<b>Figure 4. Mutations in the essential sites for GAP activity and/or involved in interaction with Rho</b>
459	<b>GTPases</b> (A) Tertiary structure of LbGAP (red) in complex with Rac1 (blue) and the transition-state
460	analogue GDP.AIF3 (modeled by homology for sequence spanning amino acid residues 51 to 216 in
461	Colinet et al., 2007). The four sites essential for GAP activity and/or involved in interaction with
462	RhoGTPases are colored green. (B) Amino acids found at the four sites essential for GAP activity
463	and/or involved in the interaction with RhoGTPases for <i>Leptopilina</i> RacGAP1 and <i>L. boulardi</i> venom
464	RhoGAP sequences. The numbering corresponds to the positions in the LbGAP sequence. (C) Amino
465 466 467 468	acids at the four sites essential for GAP activity and/or involved in interaction with RhoGTPases for <i>V. canescens</i> RacGAP1 and calyx RhoGAP sequences. The numbering corresponds to the positions in the LbGAP sequence. In B and C, amino acids are colored according to their properties following the RasMol amino acid color scheme.

R

K N

Q

F κ R

LbGAPy3.2

LbGAPy4

LbGAPy5

LbGAPy6

469 A search for positive selection was performed to detect possible functional divergence of L. boulardi 470 venom and V. canescens calyx RhoGAP sequences. PAML codon-based models with (M2a and M8) and 471 without (M1a and M7) selection were compared using likelihood ratio tests (LRTs). Both M1a/M2a and

472 M7/M8 comparisons resulted in the rejection of the null hypothesis suggesting that a fraction of codons in473 RhoGAP sequences are under positive selection (Table 3).

474**Table 3.** Positive selection analysis among sites using CodeML for *L. boulardi* venom and *V. canescens*475calyx RhoGAP sequences. InL is the log likelihood of the model. p-value is the result of likelihood ratio476tests (LRTs). Global  $\omega$  is the estimate of the dN/dS ratio under the model (given as a weighted477average). Parameters ( $\omega > 1$ ) are parameters estimates for a dN/dS ratio greater than 1.

478

L. boulardi				
Model	lnL	p-value	Global ω	Parameters ( $\omega > 1$ )
M1a: neutral	-5735.55		0.71	
M2a: selection	-5688.59	< 0.001	1.65	ω = 4.70, p = 0.21
M7: ß	-5741.07		0.70	
M8: ß&w	-5688.19	< 0.001	1.56	$\omega$ = 4.4, p = 0.22
V. canescens				
Model	lnL	p-value	Global ω	Parameters ( $\omega > 1$ )
M1a: neutral	-12616.14		0.78	
M2a: selection	-12574.17	< 0.001	1.31	ω = 2.72, p = 0.28
M7: ß	-12612.64		0.73	
M8: ß&w	-12564.70	< 0.001	1.21	ω = 2.30, p = 0.34

479

Consistently, a total of 7 and 11 branches of the phylogenetic tree constructed with the *L. boulardi* venom and *V. canescens* calyx RhoGAP sequences, respectively, were detected by the REL branch-site test as corresponding to lineages on which a subset of codons has evolved under positive selection (Figure 5A and 6A). For *L. boulardi*, 6 of the 7 lineages under positive selection were internal branches, indicating that selection occurred primarily before the separation of the *L. boulardi* Ism and ISy strains (Figure 5A).

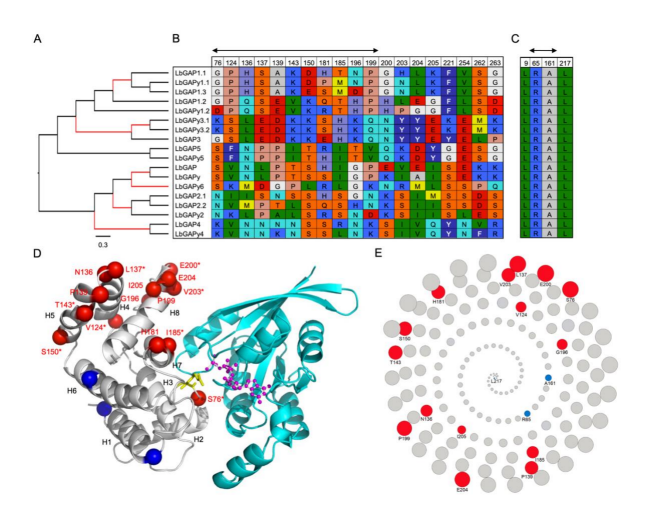
The combined use of five different methods identified a total of 19 codons as candidates under positive selection for the *L. boulardi* venom RhoGAP sequences, most of which were found in the region corresponding to the RhoGAP domain (Figure 5B). In contrast, only 4 amino acids were detected as evolving under negative selection in the *L. boulardi* venom RhoGAPs, respectively (Figure 5C). The leucine at position 9 in LbGAP is located in the signal peptide, demonstrating the importance of this region. The other three amino acids in *L. boulardi* and most of those in *V. canescens* under negative selection are buried in the protein and are probably important for the structural stability (Figures 5D and 5E).

Interestingly, at least one radical change in charge and/or polarity of the corresponding amino acids was found for all selected codons (Figure 5B). For example, the polar uncharged serine residue at position 76 in LbGAP was replaced by a negatively charged lysine residue in LbGAP4 (Figure 5B). Since radical changes are more likely to modify protein function than conservative changes, this suggests that the identified non-synonymous substitutions may be adaptive. Interestingly, the corresponding amino acids are mostly exposed on the surface of the protein and therefore likely to interact with partners (Figures 5C and 5D).

We therefore generated site-specific mutants of LbGAP for 8 of the 19 amino acids under positive selection and compared their binding capabilities to Rho GTPases. Two-hybrid analysis revealed a lack of interaction for five of the mutants, indicating that the corresponding amino acids are essential for interaction with Rac GTPases (Table 4). On the other hand, the remaining three mutants were still able to interact strongly with Rac1 and Rac2, suggesting that the corresponding amino acids are not involved in the interaction with Rac GTPases (Table 4).

For the *V. canescens* calyx RhoGAP sequences, six amino acids were identified as candidates under positive selection and 19 under negative selection (Figures 6B and 6C). Four of the amino acids under positive selection are predicted to be exposed on the surface of the protein, indicating functional diversification in relation to interaction with partners as in *L. boulardi* (Figures 6D and 6E). Most of the amino acids under negative selection are buried within the protein and are likely to be important for structural stability (Figures 6D and 6E).

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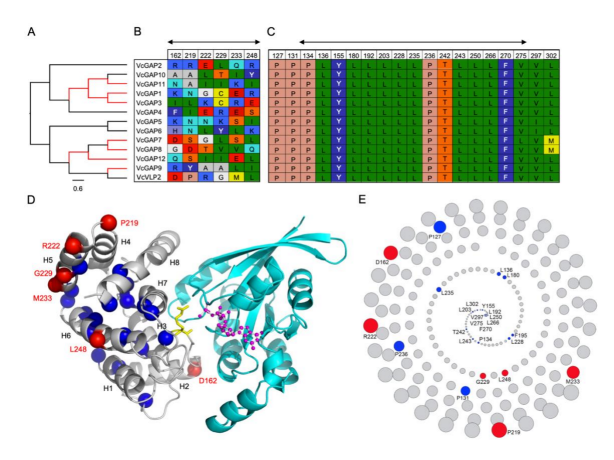
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Figure 5. Identification of positively selected branches and codons in L. boulardi venom RhoGAPs. (A) Cladogram of L. boulardi venom RhoGAPs. Branches identified under positive selection by the HyPhy BSR method are colored in red. (B) Identified positively selected codons numbered according to LbGAP amino acid sequence. Amino acids are colored according to their properties following the RasMol amino acid color scheme. The two-sided arrow indicates sites located in the RhoGAP domain. (C) Identified negatively selected codons numbered according to LbGAP amino acid sequence. (D) Positively- and negatively selected sites displayed on the tertiary structure of LbGAP (grey) in complex with Rac1 (blue) and the transition-state analogue GDP.AIF3 (modeled by homology for sequence spanning amino acid residues 51 to 216 in Colinet et al., 2007). Helices are shown as ribbons and numbered according to their location in the amino-acid sequence of LbGAP. Positively selected sites are displayed as red-colored spheres and numbered according to the LbGAP sequence. A star indicates amino acid residues tested in mutagenesis and two-hybrid interaction assays. Negatively selected sites are displayed as spheres and colored in blue. Protruding LbGAP Arg74 is shown as sticks and colored in yellow. GDP.AIF3 is shown as ball-and-sticks models and colored in magenta. (E) Spiral view of LbGAP amino acids in the order of their solvent accessibility. Most accessible amino acids come on the outermost ring of the spiral whereas the buried amino acids are occurring in the innermost ring. Radius of the circles corresponds to the relative solvent accessibility. Red and blue colors are used for positively- and negatively selected amino acids respectively. All other amino acids are in grey.

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Figure 6. Identification of positively selected branches and codons in V. canescens calyx RhoGAPs. (A) Cladogram of V. canescens calyx RhoGAPs. Branches identified under positive selection by the HyPhy BSR method are colored in red. (B) Identified positively selected codons numbered according to VcVLP2 amino acid sequence. Amino acids are colored according to their properties following the RasMol amino acid color scheme. The two-sided arrow indicates sites located in the RhoGAP domain. (C) Identified negatively selected codons numbered according to VcvLP2 amino acid sequence. (D) Positively- and negatively selected sites displayed on the tertiary structure of VcVLP2 (grey) in complex with Rac1 (blue) and the transition-state analogue GDP.AIF3 (modeled by homology for sequence spanning amino acid residues 117 to 316). Helices are shown as ribbons and numbered according to their location in the amino-acid sequence of VcVLP2. Positively selected sites are displayed as red and negatively selected sites blue spheres, respectively, and numbered according to the VcVLP2 sequence. Protruding VcVLP2 Arg156 is shown as sticks and colored in yellow. GDP.AIF3 is shown as ball-and-sticks models and colored in magenta. (E) Spiral view of VcVLP2 amino acids in the order of their solvent accessibility. Most accessible amino acids come on the outermost ring of the spiral whereas the buried amino acids are occurring in the innermost ring. Radius of the circles corresponds to the relative solvent accessibility. Red and blue colors are used for positively- and negatively selected amino acids respectively. All other amino acids are in grey.

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**Table 4.** Summary of the results of interaction assays for the LbGAP mutants. Results are based on growth on selective medium lacking histidine and qualitative ß-galactosidase overlay assays. -: no interaction; (+): very weak interaction; +++: strong interaction.

										r
	LbGAP	т								
	LUGAP	S76K	V124K	L137D	T143L	S150L	I185K	E200G	V203R	-
Rac1	+++	-	-	+++	+++	+++	-	-	-	I
Rac2	+++	-	-	+++	+++	+++	-	-	-	-
CDC42	(+)	(+)	(+)	(+)	(+)	(+)	(+)	(+)	(+)	(+)
RhoA	-	-	-	-	-	-	-	-	-	1
Lamin	-	-	-	-	-	-	-	-	-	-

### 554

### Discussion

### 555 Independent convergent recruitment of vesicle-associated RhoGAP proteins in distantly related 556 parasitoid species

557 The process of duplication of a gene encoding a protein usually involved in a key physiological process 558 whose function will be hijacked is a major mechanism of toxin recruitment (Fry et al., 2009; Wong & Belov 559 2012; Casewell et al., 2013). The results of phylogeny and domain analyses indicate that the Leptopilina 560 venom RhoGAPs and the calyx RhoGAPs found in V. canescens VLPs originated from duplication of the gene 561 encoding the cellular RacGAP1 in these species. This independent convergent recruitment may suggest a 562 similar function in virulence in these two evolutionarily distant parasitoid species. Consistently, our results 563 indicate that venom RhoGAPs in L. boulardi and L. heterotoma and calyx RhoGAPs in V. canescens are 564 associated with extracellular vesicles, named venosomes and VLPs, respectively. While these two types of 565 vesicles have different tissue origins and formation mechanisms, Leptopilina venosomes and V. canescens 566 VLPs would act as transport systems to deliver virulent proteins, including RhoGAPs, into host hemocytes 567 to alter their function (Pichon et al., 2015; Wan et al., 2019). However, the role of RhoGAPs on host 568 hemocytes is still unclear or even unknown, although we have previously shown that LbGAP inactivates 569 Rac-like GTPases (Colinet et al., 2007), which are required for successful encapsulation of Leptopilina eggs 570 (Williams et al., 2005; Williams et al., 2006).

571 RacGAP1 is an intracellular multidomain protein consisting of a coiled-coil region followed by a C1 motif 572 (protein kinase C-like zinc finger motif) and a RhoGAP domain (Tcherkezian & Lamane-Vane 2007). 573 Leptopilina venom RhoGAPs, on the other hand, possess only a RhoGAP domain preceded by a secretory 574 signal peptide. Sequence comparisons at the genomic level support the hypothesis of a single partial 575 duplication of the RacGAP1 gene in the ancestor of the Leptopilina genus, resulting in the loss of a 5' portion 576 of the sequence encoding the coiled-coil region and the C1 motif, followed by the acquisition of the signal 577 peptide sequence. This initial evolutionary event would then have been followed by further duplication of 578 the ancestral duplicate gene throughout the diversification of the genus Leptopilina, leading to the current 579 number of venom RhoGAPs. A first hypothesis for the origin of the signal peptide would be that the loss of 580 part of the 5' coding sequence in the duplicate resulted in a new N-terminal end for the encoded protein 581 with signal peptide properties. Modification of the N-terminal end of the protein into a fully functional 582 signal peptide may have required some point mutations after partial duplication, as in the case of the 583 antifreeze protein of an Antarctic zoarcid fish (Deng et al., 2010). Another possibility is that the signal 584 peptide originated from a process of partial duplication with recruitment (Katju & Lynch 2006), in which a 585 "proto-peptide signal" coding sequence was acquired from the new genomic environment into which the 586 partial copy was integrated. Finally, it is also possible that the new 5' region of the duplicated copy results 587 from a chimeric duplication (Katju & Lynch 2006), e.g. from the partial duplication of another gene or by 588 exon shuffling (Vibranovski et al., 2006). The acquisition of the signal peptide probably facilitated the 589 neofunctionalization of the venom RhoGAPs according to the evolutionary model of protein subcellular 590 relocalization (PSR) proposed by Byun McKay & Geeta (2007), according to which a modification of the N-591 or C-terminal region of a protein can change its subcellular localization, by the loss or acquisition of a 592 specific localization signal, and enable it to acquire a new function.

593 In V. canescens, the N-terminal region of the calyx RhoGAPs is more variable in length, with some 594 proteins retaining the C1 motif upstream of the RhoGAP domain and one retaining a coiled-coil region 595 upstream of the C1 motif. Most of the V. canescens calyx RhoGAPs were predicted to contain a nuclear 596 localization signal, consistent with (i) the accumulation of VcVLP2 in the nucleus of the calyx cells prior to 597 its association with the virus-derived VLPs (Pichon et al., 2015) and (ii) our results from a proteomic analysis 598 indicating that all 13 calyx RhoGAPs are associated with VLPs. Human RacGAP1 has been described to 599 localize to the cytoplasm and nucleus (Mishima et al., 2002), suggesting that V. canescens calyx RhoGAPs 600 have retained the RacGAP1 nuclear localization signal, unlike Leptopilina venom RhoGAPs. One of the V. 601 canescens calyx RhoGAP genes is located close to the RacGAP1 gene in the genome, suggesting a recent 602 partial tandem duplication. The other genes, on the other hand, are scattered throughout the genome 603 suggesting that they originate from one or more older duplication events of the RacGAP1 gene. In contrast 604 to Leptopilina, it cannot be excluded that two or more partial duplication events of the RacGAP1-encoding 605 gene occurred in V. canescens, followed by further duplication of ancestral duplicates.

### 606 Accelerated evolution through duplication and divergence: pseudogenization and/or 607 neofunctionalization?

608 Our work revealed a significant divergence for the two multigene families of RhoGAPs in L. boulardi 609 and V. canescens compared to RacGAP1, illustrated by the presence of mutations on the arginine essential 610 for GAP activity and/or on one or more of the amino acids shown to be important, or even necessary, for 611 interaction with the Rac GTPases in all RhoGAPs except LbGAP (and its homolog LbGAPy) and VcVLP2. The 612 presence of mutations at key sites in the majority of L. boulardi venom and V. canescens calyx RhoGAPs 613 suggests a loss of function by pseudogenization. However, peptide matches were found in proteomic 614 analyses for each of these RhoGAPs, indicating that the corresponding genes are successfully transcribed 615 and translated. Furthermore, our results indicate that the ancestrally acquired secretory signal peptide, in 616 which we identified an amino acid under negative selection, is conserved in all L. boulardi RhoGAPs and 617 that most V. canescens RhoGAPs conserved the nuclear localization signal of RacGAP1. In addition, all 618 RhoGAPs retained the ability to associate with vesicles, namely venosomes in L. boulardi and VLPs in V. 619 canescens. Finally, we showed that several amino acids embedded in the protein structure evolved under 620 negative selection, suggesting that they are important for protein stability. Taken together, these 621 observations are not consistent with the hypothesis that any of the mutated RhoGAPs is undergoing a 622 pseudogenization process, although it cannot be excluded that this process is quite recent and could have 623 been initiated by the loss of the GAP active site.

624 An alternative hypothesis to pseudogenization would be functional diversification of the mutated 625 RhoGAPs independent of the Rho pathway. Indeed, we have evidenced an evolution under positive 626 selection for the majority of RhoGAPs in the two multigenic families. Most of the sites under positive 627 selection are located on the surface of the protein and are therefore likely to interact with partners. In 628 addition, directed mutagenesis and two-hybrid experiments in *L. boulardi* have shown that some of these 629 amino acids are not involved in the interaction with Rac GTPases. Although the majority of studies on the 630 RhoGAP domain concern the interaction with Rho GTPases, interactions with other proteins have also been 631 described (Ban et al., 2004; Xu et al., 2013), supporting the hypothesis of a neofunctionalization of the 632 mutated RhoGAPs. Further studies will be needed to determine which functions the mutated RhoGAPs 633 have acquired in relation to parasitism. Recently, a possible role has been proposed for one of the RhoGAPs 634 found in the venom of the ISy line of *L. boulardi* in the induction of reactive oxygen species (ROS) in the 635 central nervous system of *D. melanogaster* in the context of superparasitism avoidance (Chen et al., 2021). 636 However, ROS production is notably regulated by the GTPases Rac1 and Rac2 (Hobbs et al., 2014), whereas 637 this venom protein (named EsGAP1 in Chen et al., 2021 and corresponding to LbGAPy6 in our study) has 638 likely lost its RhoGAP activity due to a mutation on the arginine at position 74. Therefore, the mechanism 639 by which EsGAP1 would be involved in ROS induction is unclear.

640 Other potential viral factors involved in parasitic success have been described previously that are 641 encoded by large gene families and similarly correspond to truncated forms of cellular proteins that have 642 retained only a single conserved domain, such as protein tyrosine phosphatases (PTPs) or viral ankyrins (V-643 ANKs) of bracoviruses. In the case of bracovirus PTPs, some are active as phosphatases, while others are 644 mutated in their catalytic site and have been shown to be inactive as PTPs (Provost et al., 2004). In the 645 latter case, as well as for V-ANK, it has been suggested that by binding to their target, they may act as 646 constitutive inhibitors of the function of the corresponding cellular protein (Provost et al., 2004; 647 Thoetkiattikul et al., 2005). Moreover, the model of adaptive evolution by competitive evolution of 648 duplicated gene copies (Francino 2005) predicts that after a first step in which different copies explore the 649 mutation space, once a protein with an optimal function is obtained, the other copies will begin to decay 650 and undergo pseudogenization. This can lead to intermediate situations where both functional and non-651 functional proteins are produced, as observed for different virulence protein families of parasitoid wasps. 652 Some of the described features of L. boulardi and V. canescens RhoGAPs may suggest that a process of 653 competitive evolution is underway in V. canescens and L. boulardi, although our results are also consistent 654 with the hypothesis of a possible neofunctionalization of these vesicle-associated proteins as discussed 655 above.

In conclusion, we evidenced the independent convergent origin and accelerated evolution of a
 multigene family of vesicle-associated RhoGAP proteins in two unrelated parasitic wasps. Strikingly, these
 vesicles, which are similarly involved in parasitism, are produced in distinct organs: the venom apparatus
 in *Leptopilina* and the ovarian calyx in *V. canescens*. In the case of *Leptopilina* RhoGAPs, the acquisition of

660 a secretory signal peptide after incomplete duplication of the RacGAP1 gene allowed secretion into the 661 venom where the vesicles are formed. V. canescens RhoGAPs, on the other hand, probably retained the 662 RacGAP1 nuclear localization signal, because it is in the nucleus of calyx cells that VLPs are formed. Another 663 striking point is that our results suggest a possible functional diversification of vesicle-associated RhoGAPs 664 in both species, with the exception of LbGAP in L. boulardi and VcVLP2 in V. canescens, which are probably 665 the only ones with RhoGAP activity. An open question would be whether all RhoGAPs are important for 666 parasite success, but by different mechanisms, independent of the Rho pathway or not, depending on 667 whether RhoGAPs are mutated or not.

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### 669

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### **Data and supplementary information availability**

- 677 Raw data from the Illumina RNASeq sequencing of the *L. victoriae* venom apparatus are available at 678 NCBI under the BioProject ID PRJNA974978.
- The mass spectrometry proteomics data from *L. boulardi* and *L. heterotoma* venosomes have been deposited to the ProteomeXchange Consortium via the PRIDE (Perez-Riverol et al., 2022) partner repository
- 681 with the dataset identifier PXD041695.
- 682 Supplementary information is available online at Data INRAE: https://doi.org/10.57745/K82IWQ

### 683 Conflict of interest disclosure

- 684 The authors declare that they comply with the PCI rule of having no financial conflicts of interest in 685 relation to the content of the article.
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931