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Genomes of trombidid mites reveal novel predicted allergens and laterally transferred genes associated with secondary metabolism

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Abstract

Background: Trombidid mites have a unique life cycle in which only the larval stage is ectoparasitic. In the superfamily Trombiculoidae (“chiggers”), the larvae feed preferentially on vertebrates, including humans. Species in the genus *Leptotrombidium* are vectors of a potentially fatal bacterial infection, scrub typhus, that affects 1 million people annually. Moreover, chiggers can cause pruritic dermatitis (trombiculiasis) in humans and domesticated animals. In the Trombidioidea (velvet mites), the larvae feed on other arthropods and are potential biological control agents for agricultural pests. Here, we present the first trombidid mite genomes, obtained both for a chigger, *Leptotrombidium deliense*, and for a velvet mite, *Dinothrombium tinctorium*. **Results:** Sequencing was performed using Illumina technology. A 180 Mb draft assembly for *D. tinctorium* was generated from two paired-end and one mate-pair library using a single adult specimen. For *L. deliense*, a lower-coverage draft assembly (117 Mb) was obtained using pooled, engorged larvae with a single paired-end library. Remarkably, both genomes exhibited evidence of ancient lateral gene transfer from soil-derived bacteria or fungi. The transferred genes confer functions that are rare in animals, including terpene and carotenoid synthesis. Thirty-seven allergenic protein families were predicted in the *L. deliense* genome, of which nine were unique. Preliminary proteomic analyses identified several of these putative allergens in larvae. **Conclusions:** Trombidid mite genomes appear to be more

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dynamic than those of other acariform mites. A priority for future research is to determine the biological function of terpene synthesis in this taxon and its potential for exploitation in disease control.

Keywords: chigger; trombiculid; scrub typhus; terpenes; isoprenoids; horizontal transfer; *Leptotrombidium*; *Dinothrombium*; *Tetranychus*; Trombidiformes

Background

The Acari (mites and ticks) are the most speciose group within the subphylum Chelicerata, with approximately 55,000 described species in both terrestrial and aquatic habitats and an estimated total diversity of up to 1 million species [1]. This assemblage is paraphyletic and is composed of two major divisions, the Parasitiformes and the Acariformes, which both contain species of medical, veterinary, and agricultural importance. For example, the Parasitiformes harbor predatory mites used in the control of agricultural pests (e.g., *Metaseiulus occidentalis*); ectoparasites of honey bees (*Varroa destructor* and *Tropilaelaps mercedesae*) that transmit pathogenic viruses; and, most famously, the ticks (Ixodida). The Acariformes include major ectoparasites and sources of allergens for humans and other animals, such as the scabies mite (*Sarcoptes scabiei*) and the dust mites (*Dermatophagoïdes* spp. and *Euroglyphus maynei*).

The role of lateral gene transfer (LGT) in the evolution of animals has remained controversial since the first metazoan genomes were sequenced. Despite the explosion of new genomic resources across a wider variety of metazoan phyla in recent years, some claims of large-scale LGT in animal genomes have been shown to be the result of flawed data-analysis methods that failed to exclude sequences from bacterial contaminants [2–5]. Attempts to conduct meta-analyses across diverse metazoan genomes using consistent criteria have also been criticized for reliance on unsound assumptions [6, 7]. However, it is irrefutable that several metazoan taxa are reliant on functions obtained via LGT for essential physiological processes, including digestion of complex and/or toxic materials (especially in the case of herbivores and plant parasites) and the avoidance of host defenses [8–11]. One group of acariform mites, the spider mites (superfamily Tetranychoidea: order Trombidiformes; Fig. 1), are major pests of various crops and rely on well-characterized lateral gene transfers of β -cyanoalanine synthase (from bacteria) and carotenoid biosynthesis enzymes (from fungi) to detoxify hydrogen cyanide in their diet [12] and to control diapause [13], respectively. Moreover, a recent in-depth analysis of LGT in the two-spotted spider mite (*Tetranychus urticae*) revealed in silico evidence that this species can synthesize pantothenic acid using genes of bacterial origin incorporated into its genome [14]. Currently, it is unknown whether LGT is a key feature of the evolution of the spider mites only or is more widespread among the Trombidiformes.

In addition to the Tetranychoidea, the Trombidiformes contain two other superfamilies of economic or clinical importance, the Trombidioidea (the velvet mites) and Trombiculoidae (the “chiggers” and related groups) [15]. These two taxa, known collectively as trombidid mites (the Trombidiae; Fig. 1), have a unique natural history among arthropods in that only the larval stage is ectoparasitic, whereas the nymphs and adults are predators of other arthropods (Fig. 2). However, the Trombidioidea and Trombiculoidae differ in their host preferences. Larvae of the Trombiculoidae are exclusively parasites of other arthropods, and some species feed on insects of medical, veterinary, and agricultural importance, including mosquitoes [16], the New World screwworm fly [17], and aphids [18] (Fig. 2). On dipteran hosts,

heavy infestations can reduce flight ability, while certain aphid species can be killed in a few days by as few as two feeding larvae [19, 20].

The more heavily studied larval stages of the Trombiculoidae (commonly referred to as chiggers or berry bugs) primarily feed on terrestrial vertebrates, including humans [21], although some little-known taxa in this superfamily are ectoparasites of invertebrate hosts in common with the Trombidioidea [22–24] (Fig. 2). Importantly, the only major mite-transmitted disease of humans, scrub typhus or tsutsugamushi disease, is vectored by chiggers in the genus *Leptotrombidium* [25], while other chigger genera have only been implicated epidemiologically as locally important vectors [26]. Scrub typhus is a severe febrile illness caused by infection with an obligate intracellular bacterium (*Orientia* spp.) in the order Rickettsiales and features an epidemiological cycle that includes wild small mammals, which are the primary hosts for many chigger species [27]. This disease has a fatality rate of 6% if not treated promptly with antibiotics [28] and has increased in incidence globally in recent years, with the annual minimum incidence reaching >17/100,000 in South Korea and Thailand in 2012–2013 [29]. In the so-called tsutsugamushi triangle within the Asia-Pacific region, scrub typhus has a median seroprevalence of 22.2% [29]; however, endemic scrub typhus has emerged in several other parts of the world within the past decade, including South America [30], the Middle East [31], and possibly sub-Saharan Africa [32]. Chiggers have also been implicated in the transmission of hantaviruses [33], *Bartonella* spp. [34], and *Rickettsia* spp. [35]. Furthermore, chiggers have direct impacts worldwide by causing trombiculiasis, which is a highly pruritic dermatitis that can afflict humans, companion animals, and domestic ruminants, potentially leading to severe hypersensitivity [36–39].

A remarkable second unique feature of trombidid mites is that the larvae induce the formation of a feeding tube or “stylostome” at the attachment site that is extraneous to the larval mouthparts [40]. These larvae are not blood feeders but ingest tissue exudates (in the case of vertebrate hosts) or arthropod hemolymph [41]. The life history of trombidid nymphs and adults has been poorly studied. However, in the Trombiculoidae, the eggs of Collembola (springtails) and other arthropods are an important part of the diet [42] (Fig. 2). Arthropod eggs may also serve as food items for adults and nymphs in the Trombidioidea [43], although some species have potential roles in biological control, as they feed on pest arthropods such as spider mites, scale insects, aphids, and termites [20, 44–46].

To date, research on trombidid mites has suffered from a dearth of molecular data that could facilitate studies on speciation; population structure; host-vector and vector-pathogen interactions; and life-history evolution in this group. To address this deficit, we present a comparative analysis of the genomes of the chigger *Leptotrombidium deliense* (the primary scrub typhus vector in Southeast Asia [47]) and the giant red velvet mite, *Dinothrombium tinctorium* (the world’s largest acarine species [48]). We show that these trombidid mites form a distinct branch of the Trombidiformes that exhibit two classes of LGT for secondary metabolism: the previously identified carotenoid

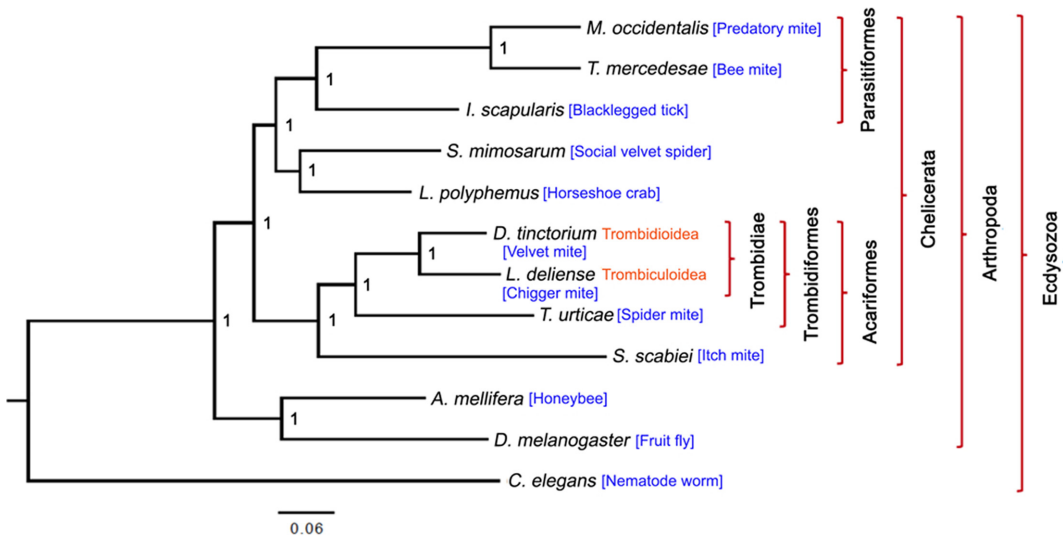


Figure 1: Phylogenetic tree based on the amino acid sequences of 476 one-to-one orthologous genes in 12 species of Ecdysozoa using Bayesian methods. The taxonomy of the trombidid mites follows the scheme of Lindquist et al. [15].

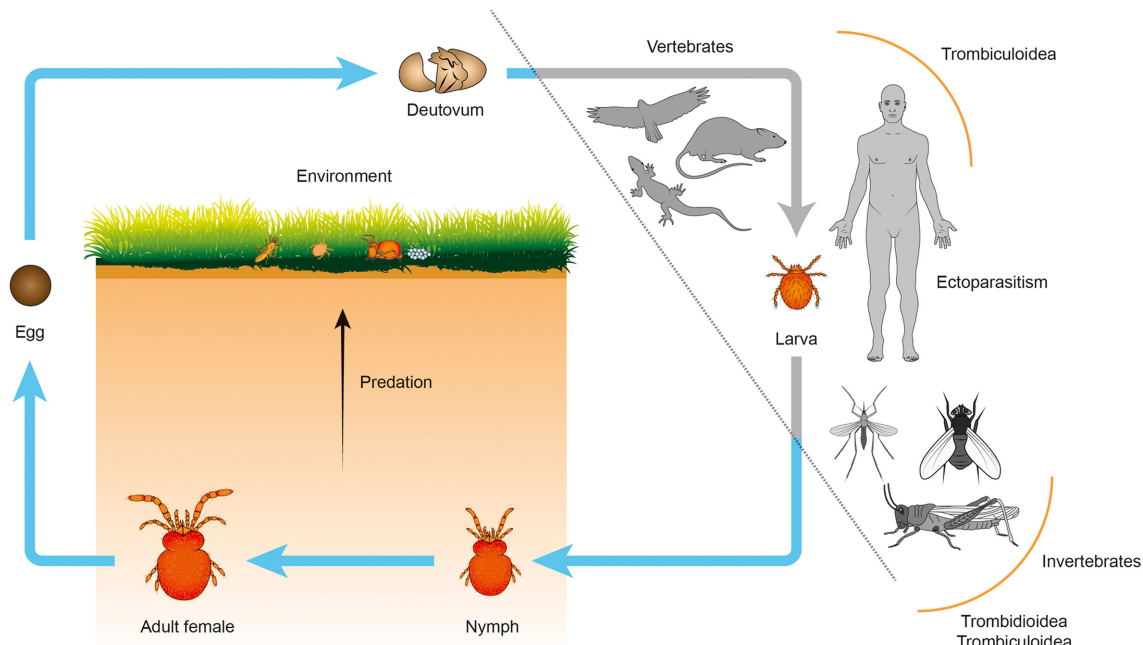


Figure 2: Simplified life cycle of trombidid mites. In the Trombidioidea (velvet mites), the larvae are parasitic on other arthropods; whereas in the Trombiculoidae ("chiggers"), the larvae feed on a variety of vertebrates or (more rarely) other invertebrates. The deutonymph and adult are free-living, edaphic stages that predate soft-bodied arthropods (e.g., termites, springtails, and other mites) or consume their eggs. Trombidid eggs are laid in the environment and produce questing larvae that congregate and seek a host. For clarity, only the deutonymph and active instars are shown; the protonymph (between the larva and deutonymph stages) and the tritonymph (between the deutonymph and adult stages) are calyptostatic.

biosynthesis enzymes of fungal origin and the much larger terpene synthase gene families, which probably derive from soil-associated bacteria. We also identify unique clusters of predicted allergens in *L. deliense* that may contribute to the symptoms of trombiculiasis in humans and domestic animals.

Data Description

Since the specimens were highly disparate in physical size (adult *D. tinctorium* can reach ~16 mm in length, whereas larval *L.*

deliense do not normally exceed 250 µm, even when engorged), a tailored approach to sequencing was necessary in each case. For the velvet mite, DNA from a single adult was used to generate two Illumina TruSeq libraries (insert sizes, 350 bp and 550 bp) and one Nextera mate-pair library (insert size, 3 Kb). The TruSeq libraries were bar coded, indexed, and paired-end (PE) sequenced (2×100 bp) on one lane, and the Nextera library was PE sequenced (2×250 bp) on an additional lane, both on the Illumina MiSeq platform. For *L. deliense*, DNA from a pool of engorged larvae (obtained from Berdmore's ground squirrels in

Thailand) was used to produce one New England Biolabs (NEB) Next Ultra DNA library (insert size, 550 bp) and PE sequenced (2×150 bp) on the Illumina MiSeq.

The total number of trimmed reads generated was ~ 362 million for *D. tinctorium* and ~ 38 million for *L. deliense*. For the former, PE reads were assembled using Abyss (v. 1.5.2) [49, 50]. For the *L. deliense* data, a preliminary assembly to contig level was performed using Velvet (v. 1.2.07) [51]. Reads derived from mammalian host genomic DNA were removed from the preliminary genome assembly using blobtools (v0.9.19), which generates a GC-coverage plot (proportion of GC bases and node coverage) [52] (Fig. 3). The *L. deliense* genome was then reassembled using SPAdes assembler (v. 3.7.1) [53] with default settings. For gene calling, the MAKER pipeline [54] was used to integrate *ab initio* gene predictions from Augustus (v. 3.2.2) [55], SNAP (v. 2013-11-29) [56], and GeneMark (v. 2.3e) [57] with evidence-based gene models. The Methods section provides more details on how downstream genome analyses were performed.

Similar to the genome sequencing strategy, preliminary proteomic analyses of the trombicid mites was customized to the sample types available. A single adult *D. tinctorium* was subjected to protein extraction in sodium dodecyl sulfate (SDS) buffer and tryptic digestion using the filter-aided sample preparation method [58]. The digested sample was split into eight fractions using the High pH Reversed-Phase Peptide Fractionation Kit (Pierce) prior to nano liquid chromatography-electrospray ionization-tandem mass spectrometry (LC MS ESI MS/MS) analysis on a Q-Exactive mass spectrometer (Thermo Fisher Scientific). A total of 137,638 spectra were generated across the eight fractions. For *L. deliense*, a soluble protein extract was obtained from a small pool of ethanol-fixed engorged larvae ($n = 10$) collected from several species of wild rodents in Thailand [27]. Following tryptic digestion, downstream analyses proceeded as for *D. tinctorium*, producing 18,059 spectra. The Methods section provides details on how MS spectra searches and Pfam enrichment analyses were performed.

Analyses

Genome statistics and phylogenomics

Assembled genome sizes were 180.41 Mb for *D. tinctorium* and 117.33 Mb for *L. deliense* (Supplementary Table S1), whereas *k*-mer analysis placed the genome size estimates much closer together but was of a similar scale to the assemblies (143.52–147.09 and 158.31–160.95 Mb, respectively; Supplementary Fig. S1). The estimate for *L. deliense* was slightly smaller than those determined for *Leptotrombidium pallidum* and *Leptotrombidium scutulare* using DNA from laboratory-reared adult specimens, which were 191 ± 7 Mb and 262 ± 13 Mb (by quantitative polymerase chain reaction [qPCR]), or 175 Mb and 286 Mb (by *k*-mer analysis), respectively [59]. The repeat content of the new genomes presented a significant challenge, with unclassified repeats alone accounting for 19%–23% of the total size, which is approximately double the proportion of the *S. scabiei* [60] and *T. urticae* [61] genomes (Supplemental Table S2). As expected from sequencing a pool of *L. deliense* larvae compared with a single adult of *D. tinctorium*, and the corresponding DNA library strategies employed, the chigger genome was considerably less contiguous than that of the velvet mite. Nevertheless, the estimated completeness of the predicted gene set for *L. deliense* (67% complete, 7.1% duplicated, 12% fragmented, 19% missing) based on the Benchmarking Universal Single-Copy Orthologues (BUSCO,

[RRID:SCR_015008](#)) criteria [62] compared favorably with that of other arachnid genomes (Supplementary Table S1).

Notably, the *D. tinctorium* genome contained the greatest number of protein-coding genes from the Acariformes sequenced to date and a very high rate of duplicated conserved genes (34%) according to BUSCO (BUSCO, [RRID:SCR_015008](#)) analysis (Supplementary Table S1). To address possible assembly artifacts caused by high levels of heterozygosity leading to collapsed haplotypes, we ran the Redundans pipeline [63] on the genomic scaffolds for *D. tinctorium*, which reduced the total scaffold number by 45% and the gene content by 36% (Supplementary Table S3). Accordingly, the BUSCO duplicated gene proportion was decreased from 34% to 12% without a detrimental impact on the number of complete conserved genes (Supplementary Table S3). However, this result seemed to contradict the *k*-mer analysis, which had not identified high levels of heterozygosity in this genome (Supplementary Fig. S1b). Moreover, reciprocal Basic Local Alignment Search Tool (BLAST) analysis of the genomic scaffolds revealed that while the duplicated genes were identical at the nucleotide level, they were always located on scaffolds with distinct sequence context outside the gene boundaries, even if this sometimes amounted to only individual single-nucleotide polymorphisms or insertions/deletions (Supplementary Table S4). As we had achieved high genome coverage ($175\times$) and a genuine large-scale chromosomal duplication event was deemed plausible (see Discussion), the Abyss assembly was used for downstream analysis without removal of “redundant” scaffolds.

Maximum likelihood and Bayesian phylogenomic analyses based on 476 one-to-one orthologous genes exhibited complete concordance, placing *L. deliense* and *D. tinctorium* together as sister taxa and placing *T. urticae* as their closest relative among sequenced species (Figs. 1 and 4). Our divergence time estimates accord closely with those previously published for arachnids [64], with the Parasitiformes and Acariformes separating approximately 430 million years ago (Mya) (Fig. 4). We estimate that the trombicid mites *sensu stricto* (velvet mites and chiggers) diverged from the phytophagous Tetranychoidae 265 Mya, and finally the Trombidioidea and Trombiculoidae last shared a common ancestor approximately 133 Mya (Fig. 4).

Gene family expansions

When gene families were compared among the Acariformes and reference invertebrate genomes, the *D. tinctorium* genome was shown to contain a substantially greater number of unique paralogous groups than the other sequenced acariform mites, while only 56 gene clusters were shared among all Acariformes (Supplementary Fig. S2). Moreover, the *D. tinctorium* genome displayed a greater number of multicopy (“N:N:N”) and patchy orthologues when analyzed alongside the published arachnid genomes (Fig. 4). The gene family expansion in *D. tinctorium* also dwarfed that seen among other members of the Arachnida, including *L. deliense* (Supplementary Fig. S3).

Relative to other acariform mites, *D. tinctorium* exhibited a significant expansion of 56 gene families, including a large family (ORTHOMCL104) of uncharacterized proteins containing 47 members in this species but no representatives in the other acariform genomes (Supplementary Table S5b). Examination of conserved domains in this gene family revealed a major facilitator superfamily domain with some weak but significant similarity by BLAST-protein (P) ($\sim 25\%$ amino-acid identity, $> 95\%$ coverage) to feline leukemia virus subgroup C receptor-related protein (FLVCR)-1 from various Metazoa. Two other orthologous clusters

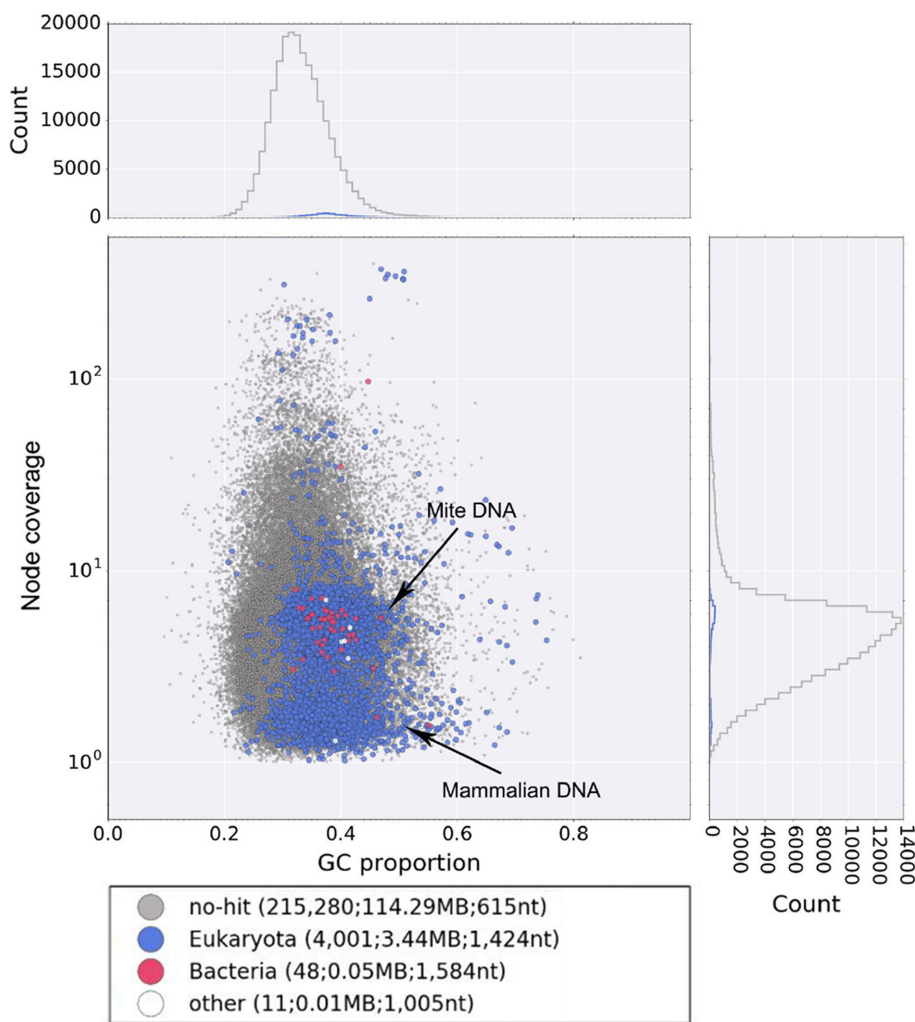


Figure 3: Blob-plot of contigs assembled from sequence data derived from engorged *Leptotrombidium deliense* larvae. Gray = no hit; blue = Eukaryota; red = bacteria; white = other.

displayed an identity of up to 30% with FLVCR2 and showed a statistically significant expansion in this species, containing 27 (ORTHOMCL316) and 21 (ORTHOMCL484) members, compared with only one and three members in *L. deliense*, respectively (Supplementary Table S5b).

Overrepresented protein families detected by LC-MS/MS

From the extremely small sample of engorged *L. deliense*, 522 mite proteins were identified by shotgun LC-MS/MS with at least one unique peptide against a background of 292 proteins of putative host origin (Supplementary Table S6b). Of the *L. deliense* proteins, 290 were considered as high-confidence identifications (≥ 2 unique peptides; Supplementary Table S6a) and were subjected to Pfam domain enrichment analysis. The most overrepresented protein domains were derived from ATP synthase (PF00006 and PF02874) and both muscle and nonmuscle isoforms of myosin or paramyosin (PF01576), although key enzymes of the citric acid cycle (ATP citrate synthase and succinate-CoA ligase, PF00549) were also well represented (Fig. 5). Several other proteins with probable origins from muscle tissue contained calponin homology domains (PF00307) and/or spec-

trin repeat domains (PF00435), including muscle protein 20 [65], myophilin, actinin, and spectrin subunits (Fig. 5). Myophilin is an invertebrate-specific protein that has previously been characterized as an immunogenic muscle component from parasitic platyhelminths [66], suggesting that it may contribute to the inflammatory response during trombiculiasis.

In a preliminary proteomic analysis of *D. tinctorium*, processing of a single adult led to the robust identification (≥ 2 unique peptides) of 1,636 proteins, as the large size of the specimen was conducive to peptide fractionation prior to LC-MS/MS (Supplementary Table S7). Laminin B (PF00052) was the most enriched domain (Fig. 5), which was present in several isoforms of perlecan (the basement membrane-specific heparan sulfate proteoglycan core protein). Perlecan is highly expressed during embryonic development in *Drosophila* [67], suggesting that this mite specimen may have contained fertilized eggs. Indeed, this specimen was undoubtedly a female, as it contained vitellogenins (yolk proteins) (Supplementary Table S7). Lyase-1 domains (PF00206) were also highly enriched (Fig. 5) and were found in mitochondrial fumarate hydratase (an enzyme of the citric acid cycle) and in adenylosuccinate lyase of the purine nucleotide cycle. The peptidase M20 (PF01546) and M20 dimer domains (PF07687) were both present in cytosol nonspecific dipep-

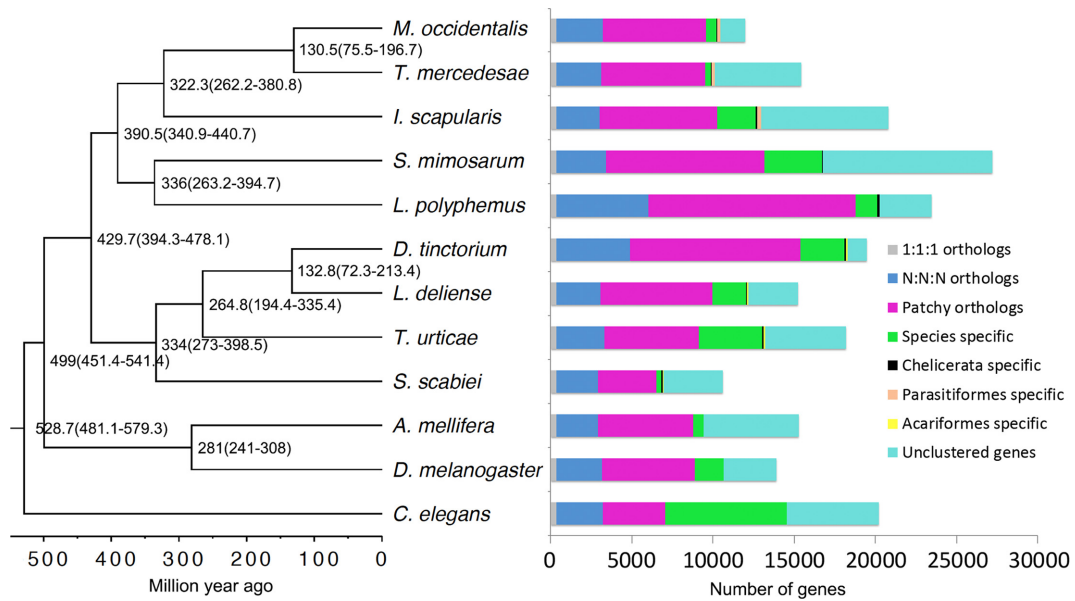


Figure 4: Estimated divergence times using a relaxed molecular clock with fossil calibration time and classification of protein-coding genes between 12 species of Ecdysozoa. *Caenorhabditis elegans* was used as the outgroup, and the bootstrap value was set as 10,000,000. The 1:1:1 orthologs comprise the common orthologs with the same copy numbers in different species, and the N:N:N orthologs comprise the common orthologs with different copy numbers in these species. Patchy orthologs are shared between more than one, but not all, species (excluding those belonging to the previous categories). Unclustered genes are those that cannot be clustered into gene families.

tidases, which have a key role in protein digestion in the midgut [68] and in N-fatty-acyl-amino acid synthase-hydrolases, which regulate thermogenesis via uncoupled respiration [69]. Another group of proteins with a putative role in thermoregulation were the α -crystallin-like small heat shock proteins (PF00525); in ticks, these are highly immunogenic proteins expressed in tick salivary glands and exhibit thermoprotective activity [70]. Finally, the inhibitor I29 domain (PF08246) represented digestive cysteine proteinases [71] and cathepsin L-like proteases (Fig. 5), which are highly expressed in feeding stages of *T. urticae* [72] and are a major protein component of spider mite feces [73].

Lateral gene transfers and mobile elements

To determine the origin of the bright coloration of the trombidid mites, we searched both genomes for the fused carotenoid synthases-cyclases that were reported to have been laterally transferred into the *T. urticae* genome from zycomycete fungi, perhaps via aphids [61]. In common with *T. urticae*, two of these carotenoid synthases-cyclases were observed in the *L. deliense* genome, while *D. tinctorium* harbored 12 copies (Supplementary Fig. S4). However, the single largest gene family expansion observed in the *L. deliense* genome was within an orthologous cluster annotated as “pentalenene synthase,” which contained 39 members (Supplementary Table S5a). This cluster (ORTHOMCL65) also contained 21 genes in the *D. tinctorium* genome but lacked orthologues in the genomes of other arachnids. A second orthologous cluster (ORTHOMCL1284) of terpene synthases contained 17 members and was unique to *L. deliense* (Supplementary Table S5a).

The capacity to generate terpenoids (also known as isoprenoids) *de novo* in metazoans is extremely unusual. While some millipedes (for instance, the Japanese species *Niponia nodulosa*) are known to produce terpenes such as geosmin and 2-methylisoborneol in defensive secretions, these secondary metabolites are assumed to be derived from microbial sym-

bionts [74]. The absence of terpene synthases in the fully sequenced diplopod genome from the rusty millipede, *Trigoniulus corallinus*, certainly supports this interpretation [75]. Dust mites also produce a monoterpane, neryl formate, which has been demonstrated to act as an aggregation pheromone [76]. However, BLAST analysis of the *Dermatophagoides farinae* genome assembly [77] using the trombidid terpene synthases failed to identify significant homologues, suggesting that dust mites rely on microbial symbionts for terpene production or that terpene synthases in mite genomes are evolving too rapidly to be identified by homology searches. To the best of our knowledge, the only animals known to harbor terpene synthase genes in their nuclear genomes are a very restricted number of beetle species (predominantly flea beetles of the subfamily Galerucinae, which produce (6R,7S)-himachala-9,11-diene as a male aggregation pheromone [78]) and the collembolan *Folsomia candida*, in which the metabolites produced and their function are unknown [79]. While terpene synthases are widespread in plants, fungi, and bacteria, the terpene synthases from flea beetles do not resemble those from non-metazoan taxa and appear to have evolved from arthropod trans-isoprenyl diphosphate synthases [78]. Moreover, the terpene synthases of *F. candida* are more similar to those from the flea beetles than they are to the non-metazoan enzymes [79].

We generated phylogenies for the trombidid terpene synthases, which clearly showed close affinities to bacterial and fungal homologues and not to those from other arthropods (Figs. 6 and 7). For ORTHOMCL65, the *L. deliense* and *D. tinctorium* enzymes formed distinct groups, and the closest homologues from other taxa included a monoterpane synthase from *Micromonospora* spp. (phylum Actinobacteria), a genus that is known to synthesize 2-methylenebornane [80], as well as related genes from agaricomycete fungi (Fig. 6). In the case of the *L. deliense*-specific ORTHOMCL1284, the nearest homologues were distributed among Actinobacteria and other bacterial phyla, including the Chloroflexi, Proteobacteria, and Bacteroidetes (Fig.

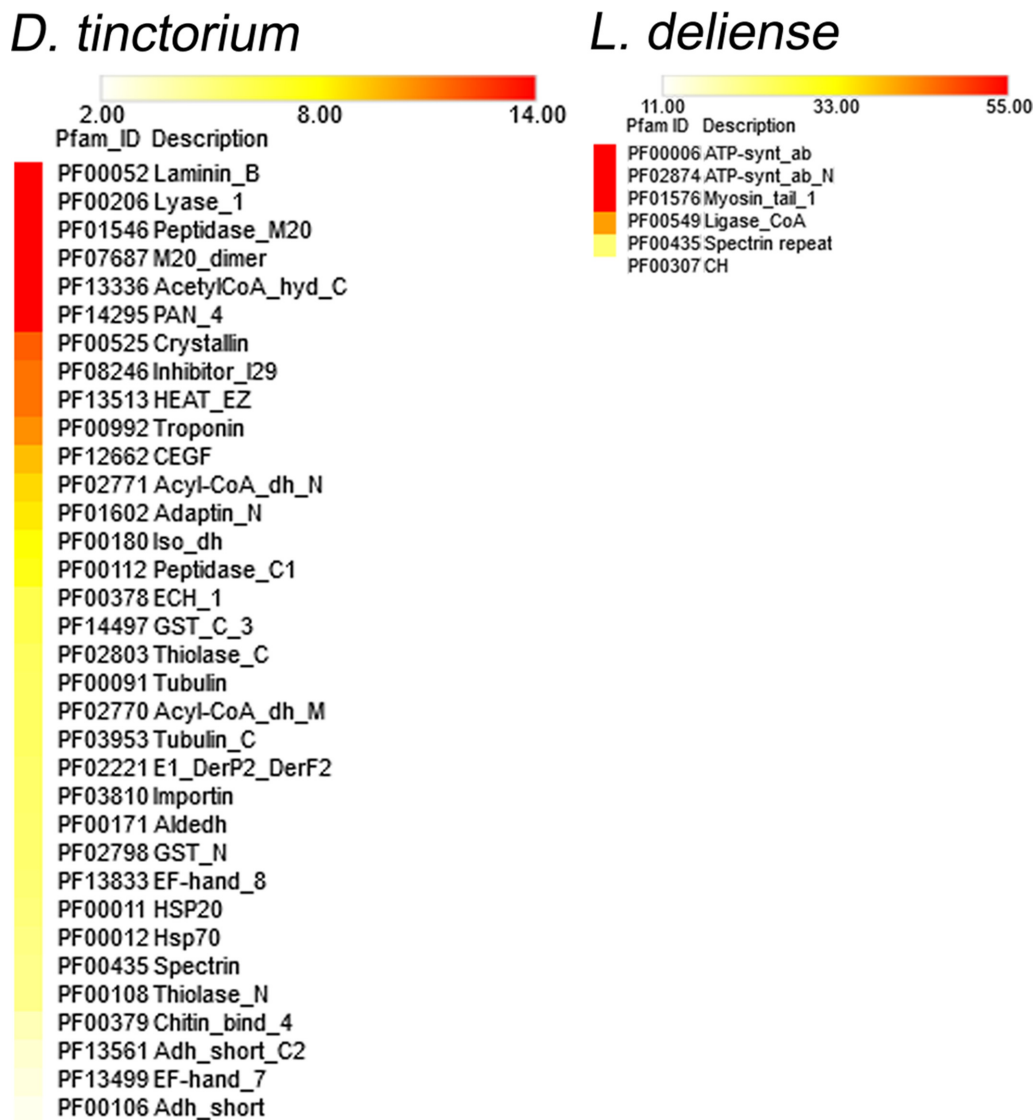


Figure 5: Overrepresented Pfam domains in proteomic datasets generated from a single adult *Dinothrombium tinctorium* and a pool of engorged *Leptotrombidium deliense* larvae. The color scale represents fold enrichment.

7). These terpene synthases were clearly separated from the flea beetle proteins and consisted predominantly of germacrene or geosmin synthases (Fig. 7). However, for both clusters of trombiculid terpene synthases, the amino acid identity with their nearest bacterial or fungal homologues was low ($\leq 30\%$). Nevertheless, the majority fulfilled the criteria of Crisp et al. [6] as high-confidence (“class A”) lateral gene transfers due to the absence of sufficiently closely related homologues in other Metazoa.

To exclude the possibility that the trombiculid terpene synthases were contaminating sequences of bacterial or fungal origin from the environment or derived from microbial symbionts, we examined the genomic context of each terpene synthase to determine if they were sometimes found adjacent to an incontrovertible metazoan gene. Due to the low contiguity of the *L. deliense* assembly, it was not possible to find other genes on the same contig as a terpene synthase. However, one of the members of ORTHOMCL65 in *D. tinctorium* was located 5.2 kb downstream of a gene with a top BLAST hit to a translational elongation factor-2 mRNA from *Dinothrombium pandorae* [81] (Supple-

mentary Fig. S5). Furthermore, blob-plot analysis revealed that the GC content and read coverage of the contigs containing terpene synthases lay close to the overall mean for both trombiculid genomes (Fig. 8) and the coverage for each gene and its parent scaffold were very similar (Supplementary Table S8). Importantly, sequences of unambiguous bacterial origin were very rare in both genomes (Supplementary Table S9), with no evidence for a high-titer symbiont or environmental contaminant that may have impacted significantly on the genome assemblies. Of these candidate laterally transferred genes in the trombiculid mites, expression at the protein level could not be detected in the small *L. deliense* sample. However, in *D. tinctorium*, a high-confidence identification of a single terpene synthase from ORTHOMCL65 was achieved based on two unique peptides, while a third peptide was shared with an additional terpene synthase (Supplementary Fig. S6).

In addition to these lateral gene transfers, the trombiculid mite genomes exhibited further evidence for dynamism in the form of endogenous retroviruses (ERVs). Both genomes showed sig-

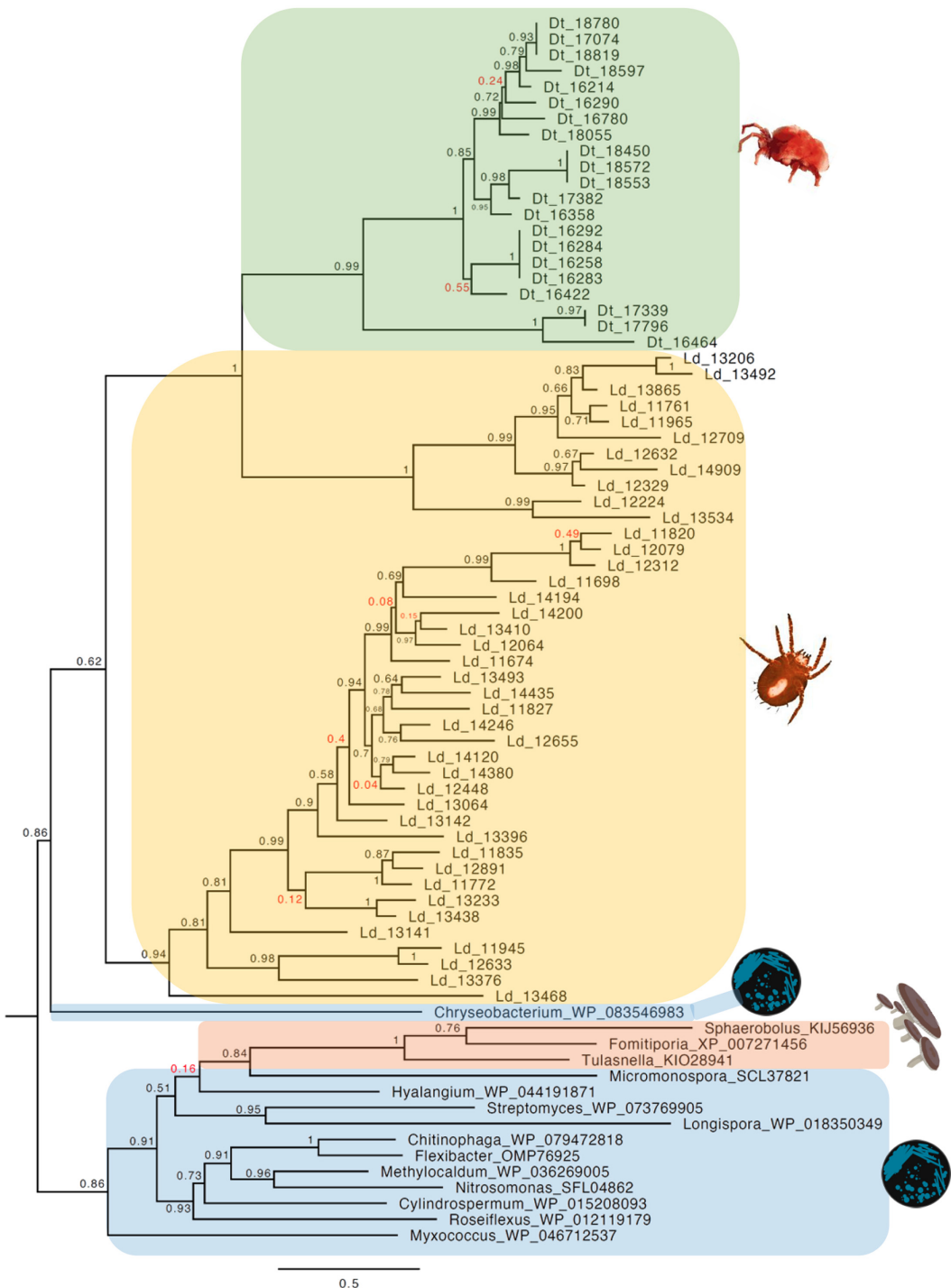


Figure 6: Phylogeny of terpene synthases from cluster ORTHOMCL65 of trombiculid mites alongside related genes from bacteria and fungi. The tree was constructed using a maximum likelihood method; poorly supported nodes are red.

nificant expansions of reverse ribonuclease integrases and Pol polyprotein-like genes, while in *L. deliense* only, a 21-member family of Gag polyprotein-like genes was apparent (Supplementary Table S5a). Interestingly, the closest homologues of the Gag-like polyproteins in *L. deliense* were found in rodents, bats, lagomorphs, small carnivores, and colugos (a taxon restricted to Southeast Asia [82]) (Fig. 9); all of which are known or likely hosts for chigger mites. Unfortunately, the low contiguity of the

L. deliense genome and the metazoan context of ERVs (with similar GC content to the host) militated against bioinformatic attempts to exclude the possibility of an origin from host contamination, i.e., from squirrel-derived cells on mite mouthparts or in the gut. In contrast, there was no evidence of a vertebrate origin of the Pol polyprotein or integrase genes in *L. deliense*, as they were clearly related to sequences from arachnids and other arthropods (Supplementary Figs. S7b and S8b).

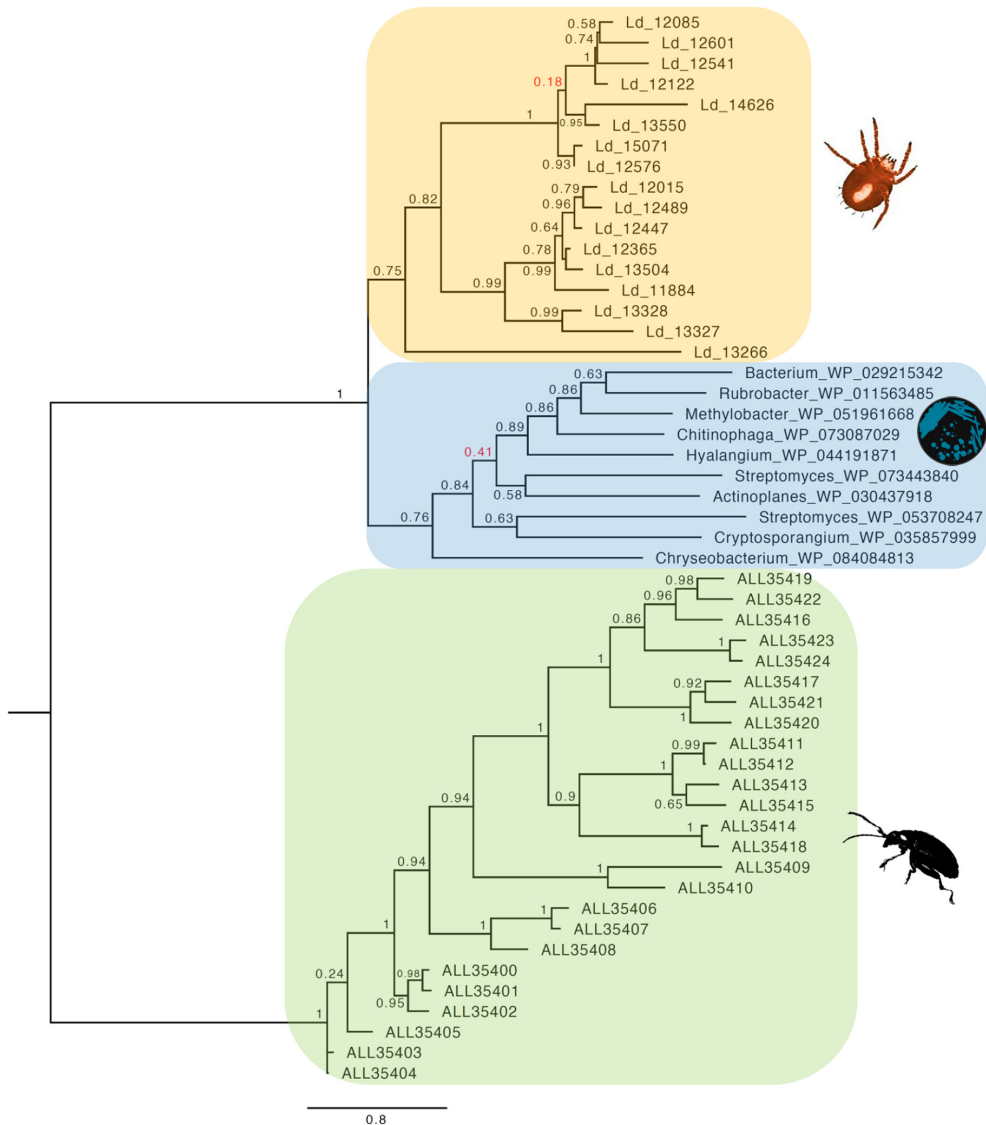


Figure 7: Phylogeny of terpene synthases from cluster ORTHOMCL1284 of *Leptotrombidium deliense* alongside related genes from bacteria. Metazoan terpene synthases from galerucid beetles are shown as an outgroup. The tree was constructed using a maximum likelihood method; poorly supported nodes are red.

In the *D. tinctorium* genome, most members of the ERV protein families each clustered in a monophyletic clade, suggesting expansion within the mite genome from a single origin (Supplementary Figs. S7a and S8a). The closest homologues of the Pol-like polyproteins were found mainly in other arthropods (especially cladocerans and ticks) and more distantly in fungi (Supplementary Fig. S8a); whereas the reverse ribonuclease integrases were most similar to those from ants, moths, nematodes, and other mites (Supplementary Fig. S7a). A single transposase family was also expanded in the *D. tinctorium* (ORTHOMCL4926; Supplementary Table S5b), with top BLAST hits in a microsporidian (*Anncaliia algerae* [83]), collembolan (*Orchesella cincta*), and other arthropods. Thus, endogenous retroviruses in *D. tinctorium* might originate from eukaryotic pathogens, larval hosts, prey species, or soil microorganisms, although the fact that some velvet mites feed on other Acari [19] renders phylogenetic analyses of potential lateral gene transfers especially problematic. Unique among chelicerate genomes sequenced to date, the *D. tinctorium* genome contained hepatitis D ribozyme-like genes

(Rfam RF01787; Supplementary Table S10), which in *Anopheles* mosquitoes, have been suggested to be involved in processing of non-long terminal repeat(LTR) retrotransposons [84].

Immune system

Many Acari act as vectors of plant or animal pathogens, and their life histories expose them to a multitude of microorganisms in their diets and in the environment. Thus, how they interact with pathogens and commensals via their immune system is likely to be a critical aspect determining their success as a group. The canonical humoral immune response gene networks in *Drosophila* are the Toll signalling pathway (responding to β -1,3-glucans from fungi and lysine-type peptidoglycan from gram-positive bacteria) and the immune deficiency (IMD) pathway (responding to diaminopimelic acid-type peptidoglycan from gram-negative bacteria). These pathways are activated when upstream transmembrane receptors (peptidoglycan recognition proteins [PGRPs] and β -glucan recognition proteins) bind to the

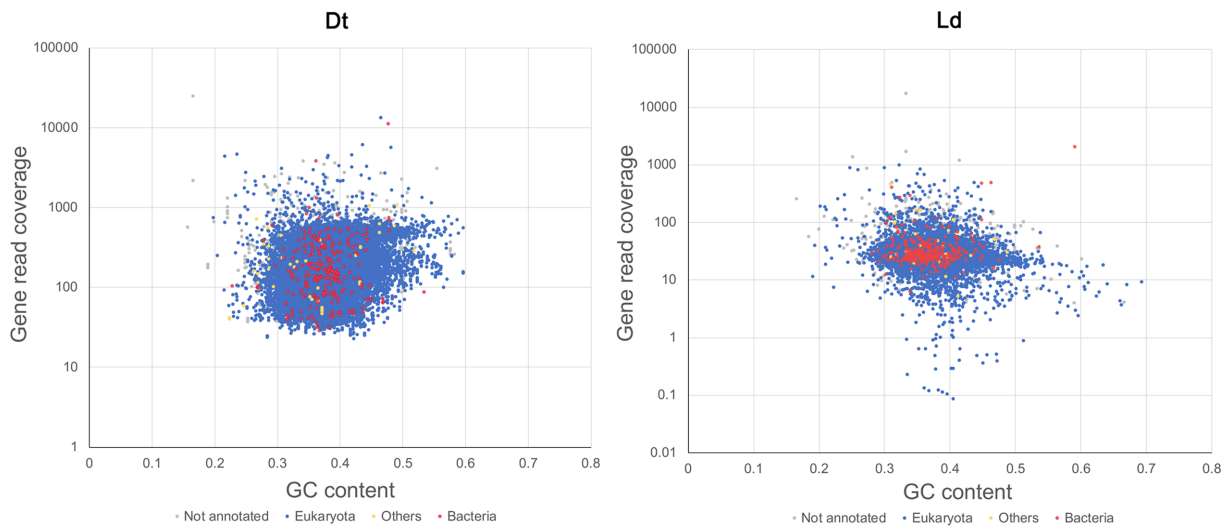


Figure 8: Blob-plot of read coverage and GC content for terpene synthase genes in *Dinothrombium tinctorium* (Dt) and *Leptotrombidium deliense* (Ld). Terpene synthases (in red, “Bacteria”) are shown in relation to all other mite genes (in blue).

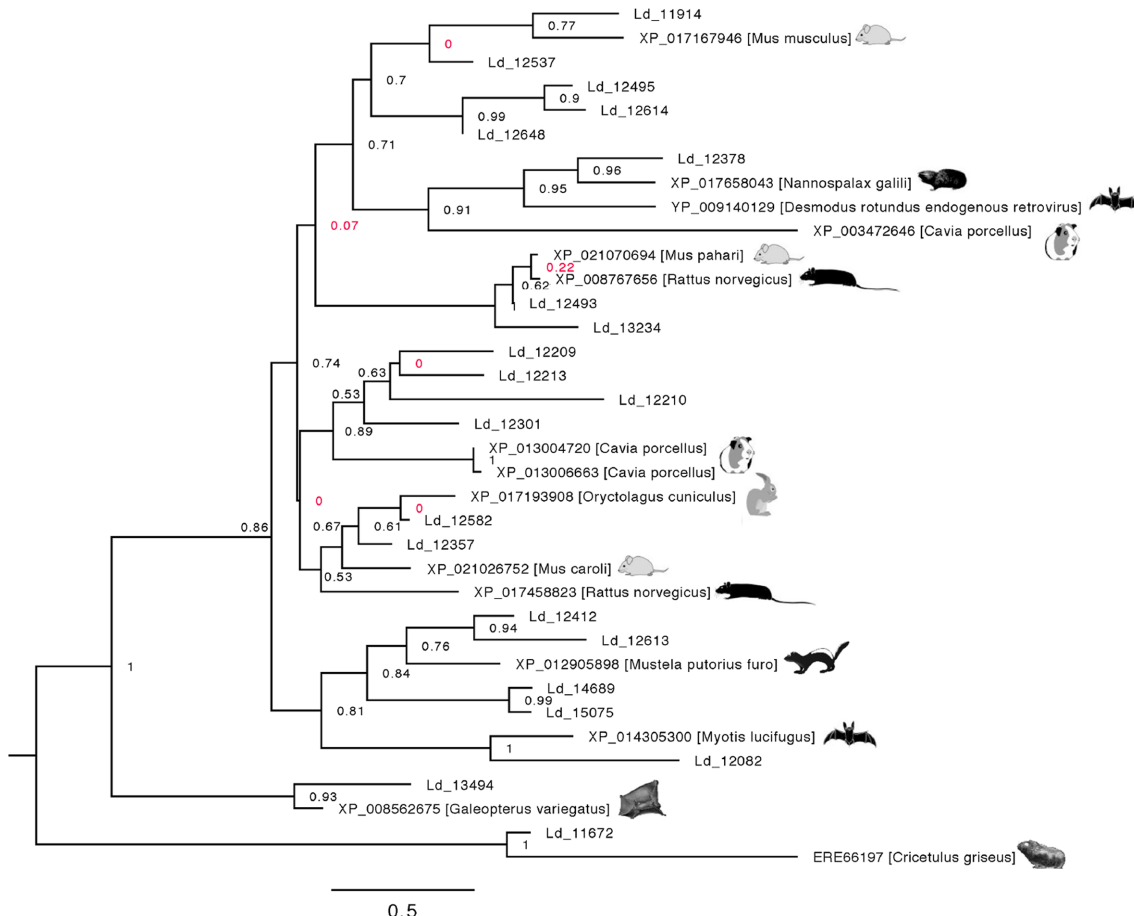


Figure 9: Phylogeny of Gag-like polyproteins from *Leptotrombidium deliense* in relation to homologous sequences from small mammals. The tree was constructed using a maximum likelihood method; poorly supported nodes are red.

pathogen-derived molecules [85]. Recently, the expanding number of arthropod genomes from outside the class Insecta has highlighted key disparities in the immune pathway genes between the Pancrustacea (Hexapoda and Crustacea) vs the Che-

licerata and Myriapoda. The most striking difference pertains to the IMD signaling pathway, which was thought to be absent in chelicerates [86]. However, genomic analyses and experimental data from *Ixodes scapularis* have revealed an alternative IMD

Table 1: Characteristics of C-type lectin domain proteins in the *Leptotrombidium deliense* genome

Cluster	No. genes in cluster	No. of genes with characteristic			Motifs		
		Secretion (%) ^a	Transmembrane domains (%) ^b	N-glycosylation (%) ^c	EPN (%)	WND (%)	QPD (%)
ORTHOMCL223	32	15 (46.9)	5 (15.6)	6 (18.8)	6 (18.8)	1 (3.1)	1 (3.1)
ORTHOMCL584	22	14 (63.6)	0	1 (4.5)	16 (72.7)	13 (59.1)	1 (4.5)
ORTHOMCL1095	16	10 (62.5)	2 (12.5)	1 (6.3)	11 (68.8)	10 (62.5)	0
ORTHOMCL1094	18	8 (44.4)	3 (16.7)	4 (22.2)	2 (11.1)	2 (11.1)	0

^aDetermined by SecretomeP 2.0 (combined percentage for both classic signal peptides and nonclassic internal secretion signatures) [94].

^bDetermined by TMHMM 2.0 [95].

^cDetermined by NetNGlyc 1.0 [96].

pathway in which interactions between IMD and Fas-associated protein with a death domain (both absent in the tick) are complemented by a E3 ubiquitin ligase (X-linked inhibitor of apoptosis protein [XIAP]) and its ligand, the E2 conjugating enzyme Bendless [87]. This pathway recognizes bacterial-derived lipids and restricts the growth of *Anaplasma phagocytophilum* and *Borrelia burgdorferi* in ticks. Although these data indicate that ticks (and perhaps other Parasitiformes) have a parallel IMD pathway distinct from that characterized in insects, we were unable to identify an XIAP homologue in acariform mites, including the trombicid genomes. This suggests that the evolution of the IMD pathway diverged along different lineages of the Chelicerata, including the paraphyletic Acari assemblage.

There are two nonexclusive scenarios that can be postulated to explain how the immune system of acariform mites operates in the apparent absence of an IMD pathway. The first is that these taxa might use the Toll pathway to respond to gram-negative bacteria as well as gram-positive bacteria and fungi. Indeed, crosstalk and synergistic immune responses to individual pathogens in *Drosophila* indicate that the two pathways are functionally interconnected, even in insects [85], and the IMD pathway may have become redundant during the evolution of acariform mites. Second, expansions in other gene families associated with the immune response may provide alternative pathogen recognition and signaling pathways to tackle gram-negative bacterial infections. This second scenario is supported in the trombicid mite genomes by large repertoires of Dscam genes (Supplementary Fig. S9), which have previously been described to have undergone expansions in the Chelicerata and Myriapoda compared to the Pancrustacea [86]. In insects, Dscam is involved in phagocytosis of bacteria by hemocytes, and the *D. melanogaster* Dscam-hv gene exhibits a remarkable capacity to generate >150,000 alternatively spliced isoforms, perhaps conferring some level of specificity to the insect immune response (although this remains highly controversial [88]). Even relative to other acarine genomes, those of the trombicid mites display a substantially greater complement of Dscam genes (~40 in *D. tinctorium*; Supplementary Fig. S9), rivaling the 60 gene family members observed in the *Strigamia maritima* (coastal centipede) genome [86, 89].

Several other expanded gene families in one or both of the trombicid mite genomes may have roles in the immune response. In common with *I. scapularis*, these genomes lack the transmembrane PGRPs that are activated in the presence of peptidoglycan in insects but contain several other PGRP genes with putative extracellular or intracellular roles. However, these soluble PGRP genes are present in larger numbers in the trombicid mite genomes than in those of *I. scapularis* and *T. urticae* (Sup-

plementary Fig. S10). Since soluble PGRP fragments can have a co-receptor function as shown in insects [90], they might work in concert with as-yet-unidentified components of the acarine immune system to recognize pathogens. This is particularly important in the case of *L. deliense*, as evidence for a peptidoglycan-like structure has recently been reported for *Orientia tsutsugamushi* [91]. Moreover, a much larger expansion in a second class of proteins with putative roles in the immune system, the C-type lectin domain (CTLD) proteins, was apparent in *L. deliense* (Supplementary Table S5a; Table 1). The CTLD protein family is a large and diverse group, most members of which do not bind carbohydrates and are thus not lectins [92]. If a CTLD protein does have lectin activity, the carbohydrate-recognition domain usually contains the amino acid motif “WND,” together with “EPN” if the specificity is for mannose, and QPD if the specificity is for galactose. However, several exceptions to this pattern do exist [92]. The expanded *L. deliense* CTLD proteins belong to four orthologous groups containing a total of 88 genes, of which one cluster (ORTHOMCL1094) contains very few genes with signatures of carbohydrate-binding activity (Table 1). The other three groups mainly contain proteins with EPN motifs, suggesting specificity for mannose, although a small proportion of QPD-motif CTLD proteins were apparent in two of the clusters, which might bind galactose (Table 1). The majority of the *L. deliense* CTLD proteins that were predicted to bind carbohydrates exhibited classic or internal secretion signatures, while only a small proportion (10–20%) contained transmembrane domains (Table 1). In common with many members of the CTLD protein family, including those in other arthropods, N-glycosylation sites were predicted in a significant minority of the *L. deliense* CTLD proteins [93] (Table 1).

Photoreceptor and chemosensory systems

Unlike insects, chelicerates lack compound eyes. Mites and ticks may be eyeless or can possess one or more pairs of simple dorsal ocelli. The Parasitiformes sequenced to date are all eyeless species (*I. scapularis*, *M. occidentalis*, and *T. mercedesae*), whereas the trombicid mites and *T. urticae* have two pairs of ocelli on the prodorsum in the adult stage [48]. However, the genomes of both eyeless and eyed Acari exhibit a variable complement of opsins that, in combination with the chromophore retinal, form light-sensitive proteins termed rhodopsins. The genomes of eyeless ticks and mites, as well as that of *T. urticae*, contain one or more genes of the “all-trans-retinal” peropsin class, which in spiders have been shown to encode nonvisual photosensitive pigments with combined G-protein coupled receptor and retinal photoisomerase activity [97]. Since even the eyeless species show evidence for reproductive and diapause behaviors that respond

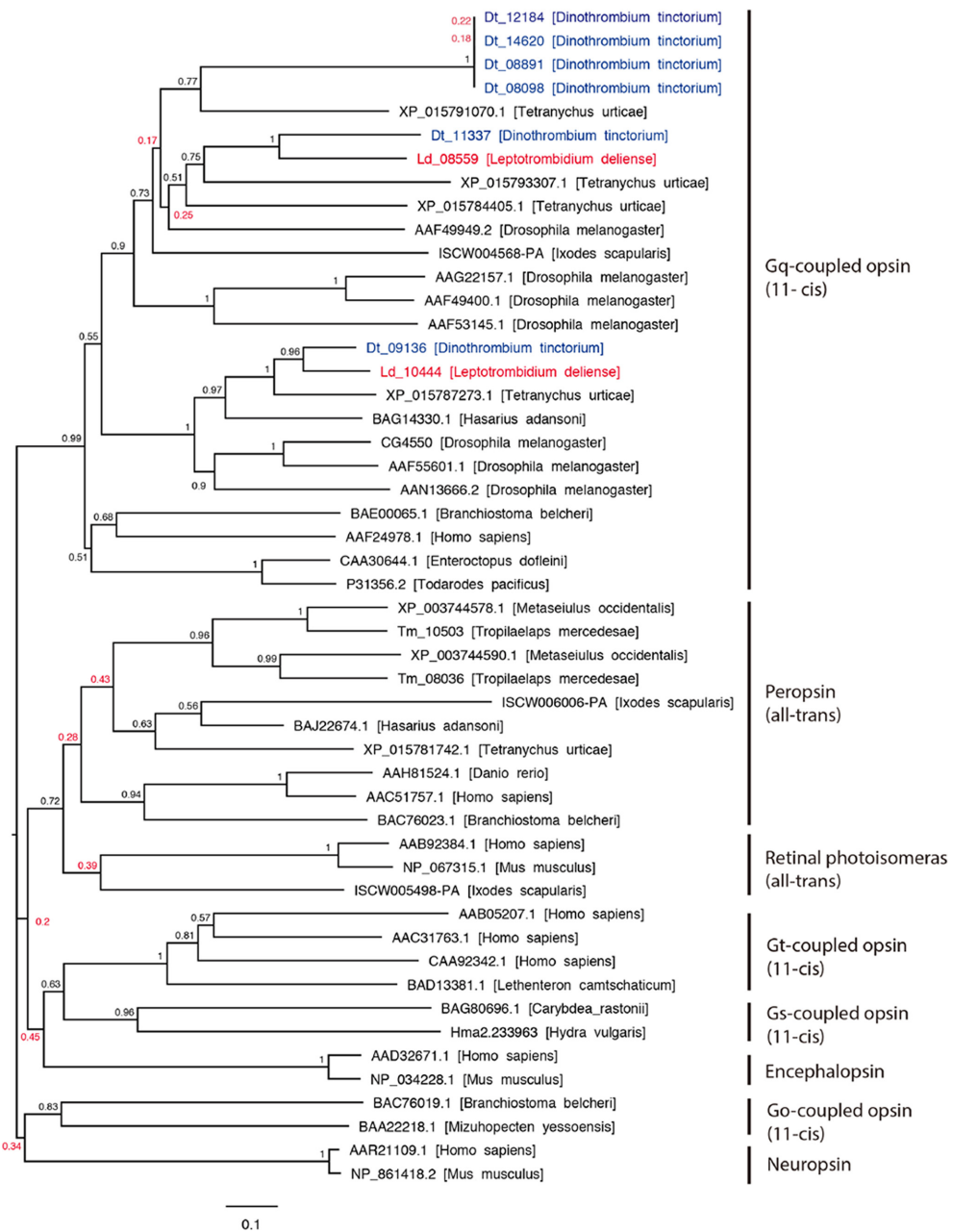


Figure 10: Phylogeny and classification of metazoan opsins. The tree was constructed using a neighbor-joining method. Poorly supported nodes are red.

to day length, it has been suggested that peropsins are important for the maintenance of circadian rhythms [98, 99]. Notably, we found no evidence of peropsin genes in the trombidid mite genomes, but did find orthologues of *T. urticae* rhodopsin-1 and -7 in both *L. deliene* and *D. tinctorium* (Fig. 10). In the latter, an additional four rhodopsin-7-like paralogues were apparent, three of which were identical at the amino-acid level (Fig. 10).

In contrast with insects but in common with crustaceans and myriapods, the Acari appear to have a scant repertoire of chemosensory protein classes, lacking both odorant-binding proteins (OBPs) and odorant receptors. Moreover, the small chemosensory proteins that have expanded considerably in some insect orders (especially Lepidoptera [100]) are completely absent in the mite genomes, although a gene encoding one such protein was identified in the *I. scapularis* genome (Table 2). Thus,

mites rely primarily on gustatory and ionotropic receptors for chemosensation. The repertoire of gustatory receptors (GRs) in *L. deliense* (42 members) and *D. tinctorium* (105 members) was in a similar range to that of most mites and ticks (albeit from the Parasitiformes) and for the Mandibulata (Table 2); hence, there was no evidence for the massive expansion in this gene family recently reported for the *T. urticae* genome, with almost 700 members [101].

Ionotropic glutamate receptors (iGluRs) are glutamate-gated ion channels that are divided into two subtypes based on sensitivity to N-methyl-D-aspartic acid (NMDA). The canonical iGluRs do not have direct roles in chemosensation. Rather, at least in *D. melanogaster*, the NMDA-sensitive channels are expressed in the brain and are involved in associative learning and memory [103]. The non-NMDA channels have fundamental roles in synaptic

Table 2: Comparison of chemosensory receptor repertoires between trombidid mites and 11 other arthropods

Species	Chemosensory receptor ^a				
	GR	OR	IR	OBP	CSP
<i>D. tinctorium</i>	105	0	7	0	0
<i>L. deliense</i>	42	0	8	0	0
<i>T. urticae</i>	689	0	4	0	0
<i>T. mercedesae</i>	5	0	8	0	0
<i>M. occidentalis</i>	64	0	65	0	0
<i>I. scapularis</i>	60	0	22	0	1
<i>S. maritima</i>	77	0	60	0	2
<i>D. pulex</i>	53	0	85	0	3
<i>D. melanogaster</i>	73	62	66	51	4
<i>A. mellifera</i>	10	163	10	21	6
<i>B. mori</i>	56	48	18	44	18
<i>A. pisum</i>	53	48	11	15	13
<i>P. humanus</i>	8	10	12	5	7
<i>humanus</i>					

^aCSP, chemosensory protein; GR, gustatory receptor; IR, ionotropic receptor; OBP, odorant-binding protein; OR, olfactory receptor. Data for other genomes were obtained from [89, 98, 99, 101, 102].

transmission in the neuromuscular junction within muscle tissue or in the nervous system [104], and certain receptor subunits have been shown to be involved in the regulation of sleep (GluR1 [105]) or vision (Clumsy, CG5621, CG3822, CG11155, CG9935 [106, 107]). Strikingly, the *D. tinctorium* genome harbored seven NMDA-type iGluRs and 61 non-NMDA iGluRs, representing substantially greater repertoires than those observed for the *L. deliense*, *T. urticae*, and *D. melanogaster* genomes (especially for the non-NMDA iGluRs) (Fig. 11). The chemosensory ionotropic receptors (IRs), which exhibit sequence similarity to iGluRs but do not bind glutamate [108], also showed interesting differences in gene family size compared with *T. urticae* and *D. melanogaster*. Notably, while *D. melanogaster* has one gene encoding an IR25a protein, *T. urticae* has three such genes, and the trombidid mites have five copies each (Fig. 11). The *D. melanogaster* IR25a is a widely expressed co-receptor that couples with stimulus-specific IRs to facilitate sensitivity to a diverse range of acids and amines. Recently, IR25a in combination with IR21a and IR93a were demonstrated to function as a thermosensory complex expressed by the dorsal organ cool cells of *D. melanogaster* larvae, which mediates avoidance behavior to cool temperatures (<20°C) [109, 110]. Sequences that cluster with *D. melanogaster* IR21a and IR93a in the “antennal and first leg” class of IRs were identified in the trombidid mite genomes, with one copy in *D. tinctorium* (as for *T. urticae*) and three copies in *L. deliense* (Fig. 11). Although chelicerates lack antennae, the orthologues of IR93a and/or IR25a have been shown to be highly expressed exclusively in the first pair of legs in *T. mercedesae* [99] and *Varroa destructor* [111], suggesting functional parallels between insects and mites.

Predicted allergens

Although the propensity of chiggers to cause pruritic dermatitis is well recognized in humans and other animals [36–39], the identity of the allergens involved has not been established [112]. The *L. deliense* and *D. tinctorium* genomes were predicted to encode 37 and 33 groups of protein allergens, respectively, which is substantially more than other sequenced mites in the Acariformes with the exception of the dust mites, *D. farinae* [77], and *E. maynei* [113]. Since velvet mites rarely come into contact with

humans, only the *L. deliense* allergens were subjected to further analysis. The *L. deliense* predicted allergen clusters included nine groups that were unique to this species and six that were shared with *D. tinctorium* only (Fig. 12), while an additional 28 putative allergen genes in the *L. deliense* genome did not cluster in orthologous groups (Supplementary Table S11b). The *L. deliense*-unique groups included five distinct clusters of trypsin-like serine proteases and one cluster each of subtilases, papain-like cysteine proteases, enolases, and cyclophilins, all of which could be classified into recognized allergen families listed in the AllFam database [114] (Fig. 13). The nonclustered allergens belonged to a variety of structural and enzymatic protein groups, but cathepsins, serine proteases, and peptidylprolyl isomerases were the most common annotations (Supplementary Table S11b).

The major allergens in *D. farinae* are the 25-kDa Der f 1, a papain-like cysteine protease, and Der f 2, a 14-kDa uncharacterized protein with an ML (lipid-binding) domain [77]. However, many other minor allergens have been detected by immunoproteomic studies [115, 116] or predicted by homology searches in the *D. farinae* genome [77]. In *L. deliense*, five distinct clusters of papain-like cysteine proteases were identified (AllFam AF030), of which three were shared with *D. farinae*, one was shared only with *D. tinctorium*, and one was unique (Fig. 13). No orthologue of Der f 2 (AF111) was apparent.

Recently, an alpha-enolase has been reported as a novel minor allergen in *D. farinae* [116]. However, the two enolases (AF031) with predicted allergenic properties in the *L. deliense* genome formed an orthologous cluster that was absent from other mites sequenced to date, with homologues in parasitic nematodes and distant chelicere relatives (e.g., horseshoe crabs; Supplementary Table S11a). A similar pattern was observed for the cyclophilins (AF038), which have previously been considered a class of allergens restricted to fungal and plant sources [117], although these peptidyl-prolyl cis-trans isomerasers are universally present across all domains of life. In an immunoproteomic study, a cyclophilin was newly identified as a dust mite allergen, Der f 29 [115], but this was not closely related to the *L. deliense* cyclophilins, which exhibited a greater affinity (~75% identity) to homologues in fungi and fish (Supplementary Table S11a). The *L. deliense* subtilases (serine proteases with a peptidase S8/S53 domain, AF021) were also absent from other mite genomes but showed 40%–50% identity to subtilases from fungi and bacteria (Supplementary Table S11a). These proteases have been identified as major allergens produced by ascomycete fungi such as *Curvularia lunata* [118] and *Trichophyton* spp. [119]. Finally, the five clusters of trypsin-like serine proteases (AF024) exhibited the closest homologues (40%–50% identity) in a diverse range of organisms, including *T. urticae* (but too distant to cluster in ORTHOMCL31), Diptera and scorpions (ORTHOMCL32), fish and lizards (ORTHOMCL88), bugs and ants (ORTHOMCL89), and acorn worms and Diptera (ORTHOMCL90) (Supplementary Table S11a). Thus, these predicted allergens were distinct from the *D. farinae* molecules classified in AF024 (Der f 3, 6 and 9 [120]), although within ORTHOMCL29, ORTHOMCL30, and ORTHOMCL43, *L. deliense* does possess additional trypsin-like proteases that are orthologous to these *D. farinae* allergens (Fig. 13).

A label-free quantitative analysis of protein content in the *L. deliense* indicated that muscle-derived allergens related to the *D. farinae* paramyosin Der f 11 (AF100 [121]) and to *T. urticae* tropomyosin isoforms (AF054) were most abundant (Fig. 13). Although single unique peptides were detected for several *L. deliense*-specific allergen clusters and unclustered allergenic proteins, only one *L. deliense*-specific allergen was present in quantifiable amounts (an enolase in AF031), and this was consider-

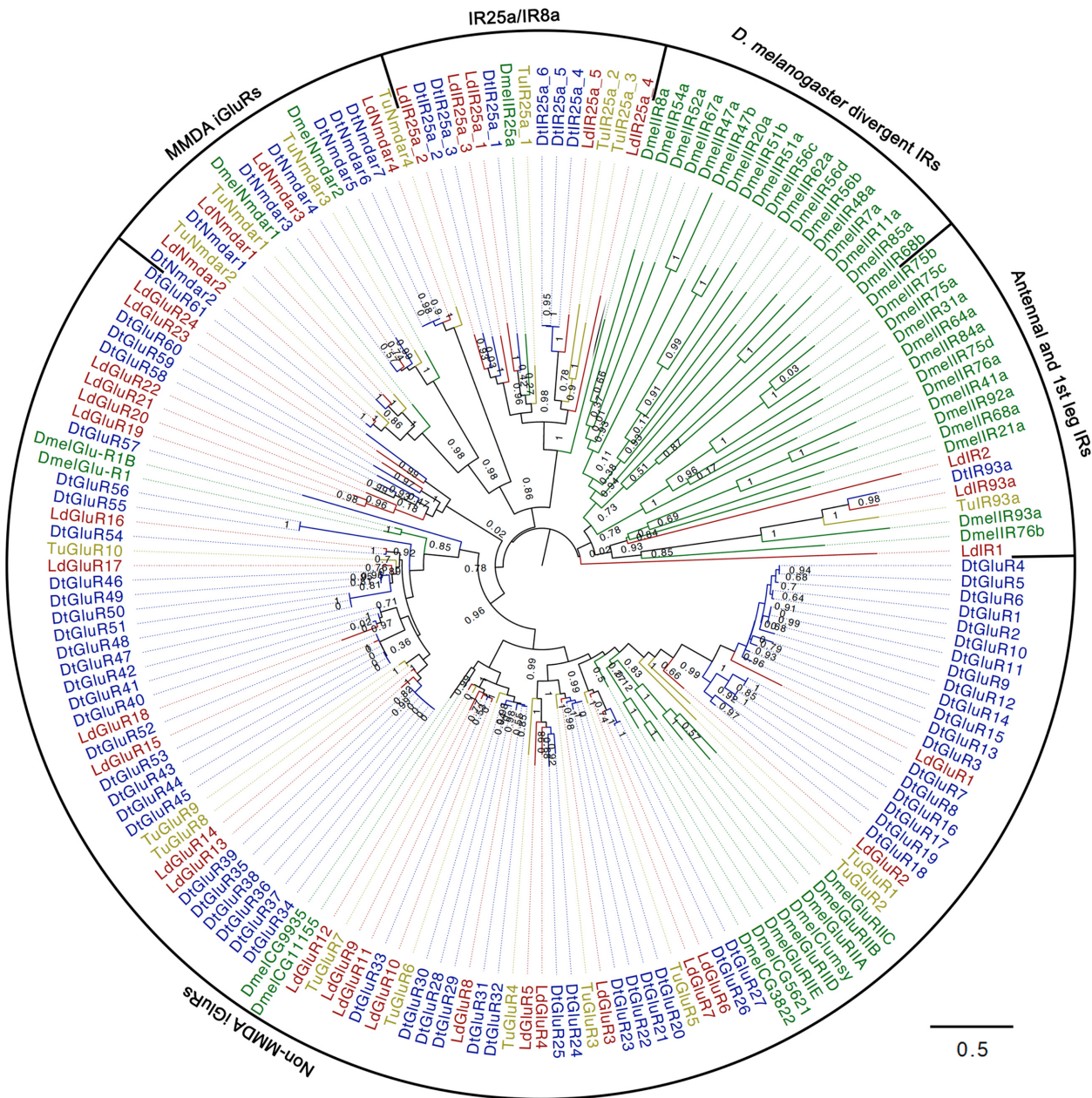


Figure 11: Phylogeny of *Dinlothrombium tinctorium*, *Leptotrombidium delense*, *Tetranychus urticae*, and *Drosophila melanogaster* ionotropic receptors and ionotropic glutamate receptors. The tree was constructed using a maximum-likelihood method.

ably less abundant than the shared allergens (Supplementary Table S6a, Fig. 13). However, as allergenicity is not dictated entirely by allergen quantity and can vary markedly between individuals, validating the identity of the most important allergens in chiggers will require screening of sera from trombiculiasis patients.

Putative salivary proteins

Due to the diminutive size of chiggers and the absence of any artificial feeding mechanism for laboratory colonies that might allow collection of saliva, the chigger sialome has not been characterized to date. However, numerous high-throughput studies of tick saliva have been conducted on several genera and multiple life cycle stages [122–127], and recently an elegant proteomic

analysis of *T. urticae* saliva was published, in which mite salivary secretions were collected in an artificial diet substrate [128]. Using proteomic datasets from this *T. urticae* study and a recent *I. scapularis* sialome analysis conducted over several time-points [122], we identified one-to-one orthologues of the salivary proteins from both sources in the tick, spider mite, and trombiculid mite genomes. We reasoned that as *T. urticae* is phylogenetically close to the trombiculid mites and *I. scapularis* is very distant, protein families shared by the tick and the trombiculid mites but not present in the *T. urticae* genome are likely to represent proteins required for ectoparasitism on animal hosts (as opposed to phytophagy). Indeed, 24 orthologous clusters were shared among the animal ectoparasites but not with *T. urticae*, whereas only five clusters were shared between all mites at the exclusion of the tick (Fig. 14). These 24 animal-ectoparasite clusters are can-

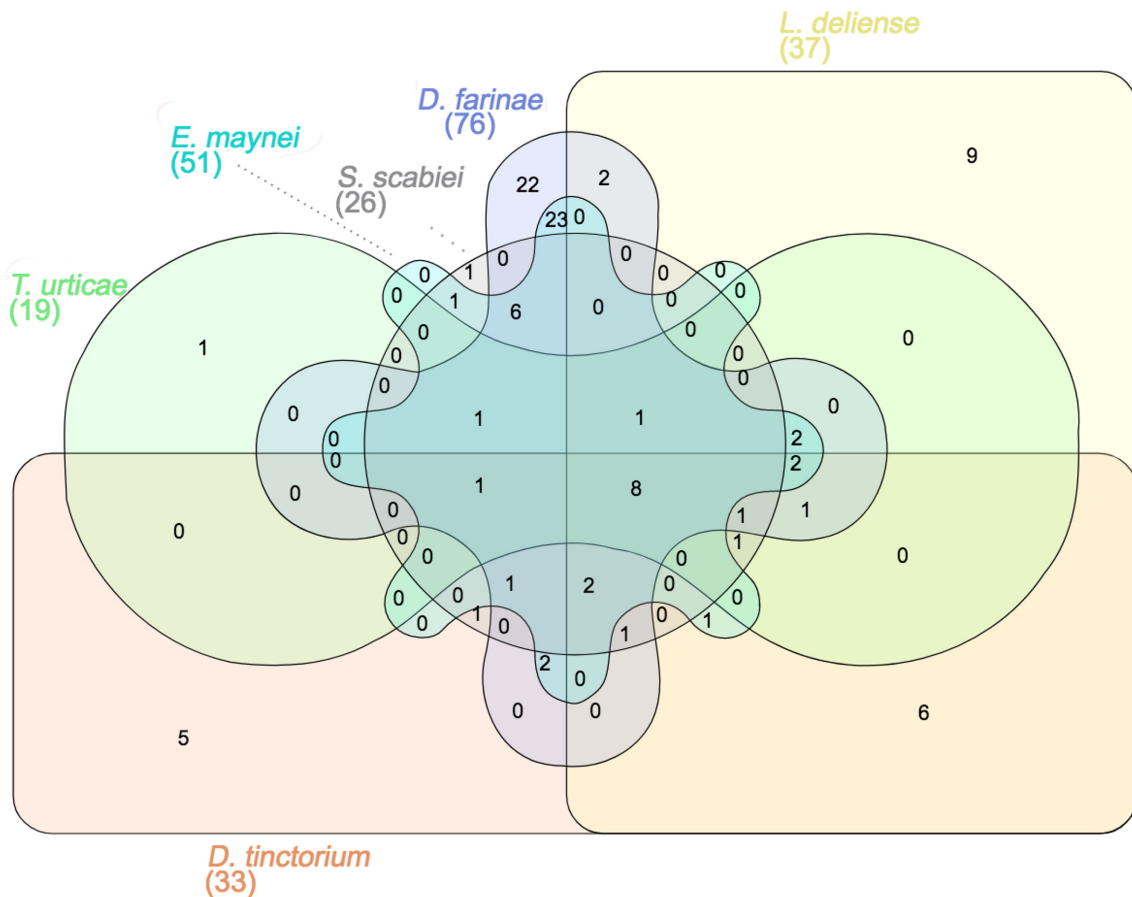


Figure 12: Venn diagram of orthologous clusters of predicted allergens from six species of acariform mites.

candidates as key salivary components of trombiculid mites. An additional two clusters were shared exclusively by *I. scapularis* and *L. deliense*, suggesting that they might be important for feeding on vertebrate hosts (Fig. 14).

To feed successfully, ticks must suppress local immune responses and prevent the clotting of blood. Although trombiculid mites feed on tissue exudates or hemolymph rather than blood and do not feed for as long as some hard tick species, they face similar challenges as ectoparasites that provoke an inflammatory response in their hosts. Interestingly, in accordance with low levels of heme in the trombiculid mite diet, we did not find orthologues of tick salivary proteins involved in heme detoxification (ferritins and hemelipoproteins [124]) in the trombiculid mite genomes. Several lipocalins with histamine-binding activity have been identified in tick saliva from multiple different species [122, 124, 125], but orthologues of these small proteins were also not present. However, genes encoding two enzymes involved in catabolism of the histamine precursor histidine, urocanate hydratase and formiminotransferase-cyclodeaminase, were detected in both trombiculid mite genomes (Table 3). The degradation of histidine feeds into the one carbon pool by folate, and this process is mediated in part by formyltetrahydrofolate dehydrogenase, an enzyme that is also present in multiple copies in the *D. tinctorium* genome (Table 3). While the presence of folate biosynthesis enzymes in tick saliva has been reported previously (and not only for *I. scapularis* [124]), the functional significance of their secretion is unclear. One possibility is that ectoparasitic Acari not only utilize bacterial symbionts as folate

“factories” [129, 130] but they can scavenge it at the source from precursors in their B vitamin-deficient diets.

Several other protein clusters with potential roles in immune evasion or the regulation of salivation and the ingestion of host fluids were identified in the trombiculid mite genomes. The presence of an expanded acetylcholinesterase gene family in tick genomes has been noted previously, and acetylcholinesterases have been detected in the saliva of *Rhipicephalus microplus* [132] and *Amblyomma americanum* [124], as well as *I. scapularis* [122]. It has been proposed that salivary acetylcholinesterases could interfere with cholinergic signaling between host immune cells and might facilitate pathogen establishment [132]. However, the trombiculid mites have only a single gene copy each that clusters with the *I. scapularis* acetylcholinesterases (Table 3). Similarly, ATPase inhibitors (Table 3) in saliva could impact on local immune responses [124] since extracellular purine metabolites act as “alarmins” [133]. The massive expansion of sulfotransferases in the *I. scapularis* genome and the secretion of some members of this family in saliva is particularly enigmatic. However, recently it has been proposed that sulfotransferases could control salivation and feeding cycles in ticks by sulfating the neurotransmitters dopamine and octopamine [134]. Alternatively, or in addition, they might be involved in increasing the activity of small cysteine-free thrombin inhibitors in tick saliva by sulfation of tyrosine residues [135]. Notably, only two of these sulfotransferases were present in each trombiculid mite genome compared with >70 members of this family in *I. scapularis* (Table 3).

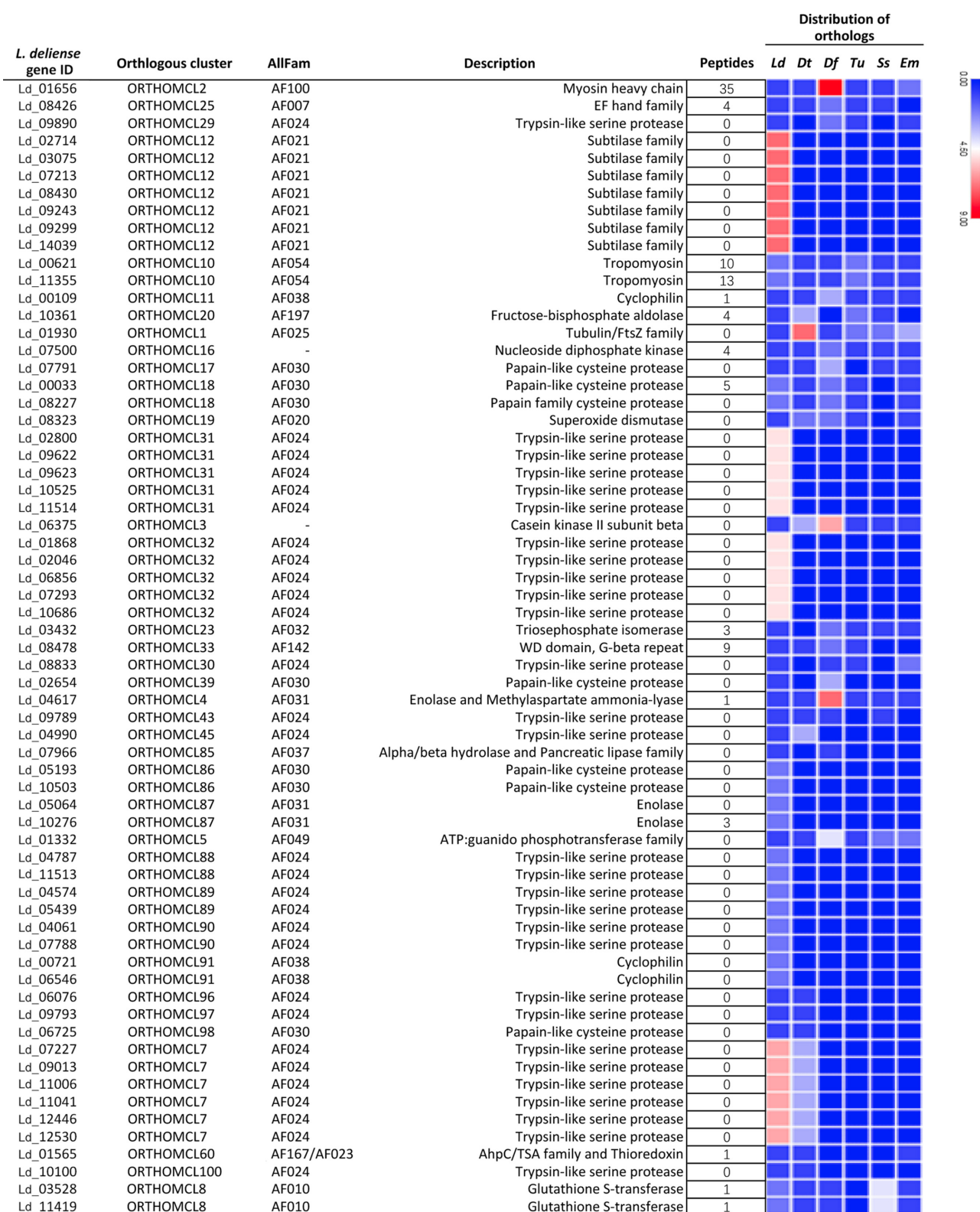


Figure 13: Orthologous clusters of predicted allergens in *Leptotrombidium deliense* as classified by the AllFam database. "Peptides" refers to the number of unique peptides from each allergen detected by mass spectrometric analysis of a pool of *L. deliense* larvae. The heat map indicates the number of orthologues for each predicted allergen in the genomes of six acariform mites. Ld, *Leptotrombidium deliense*; Dt, *Dinlothrombium tinctorium*; Df, *Dermatophagoides farinae*; Tu, *Tetranychus urticae*; Ss, *Sarcoptes scabiei*; Em, *Euroglyphus maynei*.

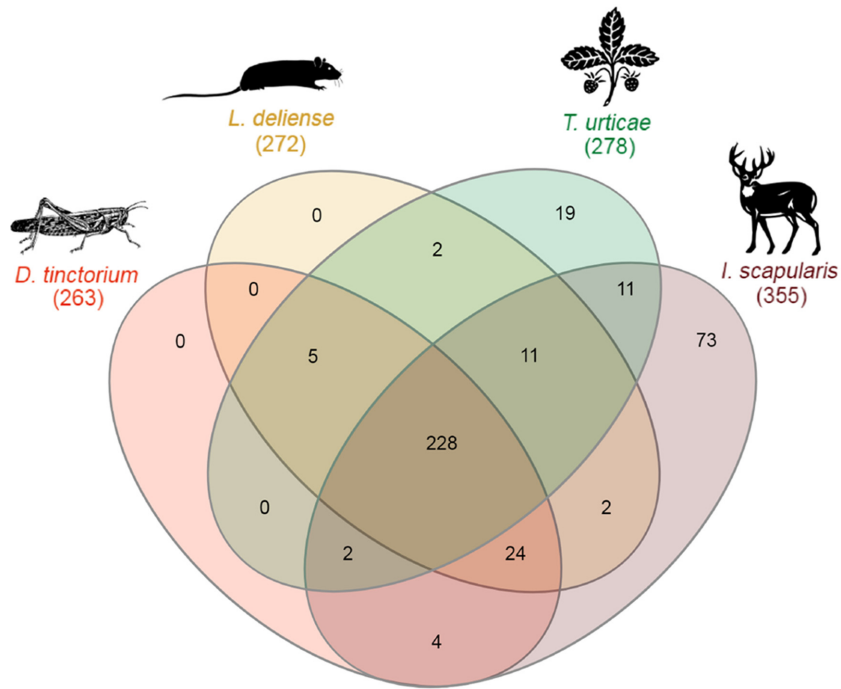


Figure 14: Venn diagram of orthologous clusters of putative salivary proteins in four species of Acari. One-to-one orthologues of salivary proteins from *Ixodes scapularis* [122] and *Tetranychus urticae* [128] were identified in the genomes of *Leptotrombidium deliense* and *Dinothrombium tinctorium*. Thumbnail images indicate representative host species.

Of the two salivary protein families restricted to *L. deliense* and *I. scapularis* (Table 3), the secreted trypsin inhibitor-like cysteine-rich domain proteins are among a wide diversity of serine protease inhibitors produced by ticks [136]. This specific class of trypsin inhibitor-like protein includes ixodidin, an antimicrobial peptide, which is expressed in the hemocytes of *R. microplus* [137] and BmSI-7 and BmSI-6, two peptides from the same species of tick that inhibit cuticle-penetrating proteases secreted by entomopathogenic fungi [138]. The BmSI-7 peptide is expressed in multiple tissues, including the salivary glands [138], but its role in saliva is unknown. However, one possibility is that it helps prevent the tick bite site from becoming infected. In contrast with *I. scapularis*, which harbors 27 trypsin inhibitor-like proteins in its genome, only one orthologue was identified in *L. deliense* (Table 3). The second cluster restricted to *I. scapularis* and *L. deliense*, a signal sequence receptor subunit (Table 3), was unexpected as it has a canonical function in trafficking secretory proteins through the endoplasmic reticulum (ER) [139]. This appears to have a moonlighting role in tick saliva since it generates strong immune responses in rabbits parasitized by *A. americanum* [124].

Feeding ticks secure their mouthparts in the skin of the host for days or weeks using a cement-like substance that forms a cone in the bite wound. Superficially, the stylostome generated at the feeding site of trombicid mites resembles the tick cement cone, although the structure is tubular (in the Trombiculidae) or highly branched (in the Trombidioidea) [41]. In both of the trombicid mite genomes, we found an orthologue of a glycine-rich protein present in the sialome of *Rhipicephalus pulchellus* [126] (Table 3). The tick glycine-rich proteins are related to spider silk proteins and form the main structural component of tick cement as determined by proteomic studies [140]. To determine if the trombicid mite genomes may contain other cement-associated proteins not detected in the *I. scapularis* salivary pro-

teomics study [122], we searched for orthologues of all tick cement proteins in the National Center for Biotechnology Information (NCBI) database. We found orthologues of an *I. scapularis* glycine-rich cement protein in both *L. deliense* (one copy) and *D. tinctorium* (three copies) that was distinct from the *R. pulchellus* orthologue; moreover, the velvet mite also possessed an orthologue of a second *I. scapularis* cement protein (Supplementary Table S12). In addition, both trombicid mites harbored a gene related to a cement protein transcript identified in the sialotranscriptome of *Amblyomma triste* [127] (Supplementary Table S12). Finally, orthologues of *A. americanum* acidic chitinases involved in conferring stability to the tick cement cone were present in both *D. tinctorium* (four copies) and *L. deliense* (one copy) [141] (Supplementary Table S12).

Discussion

Genome features and trombicid mite evolution

In this study, we exploited the close phylogenetic relationship between the Trombicidoidea and the Trombiculidoidea in order to obtain a genome from a single adult velvet mite that could be used to corroborate data derived from a suboptimal trombiculid mite sample (i.e., a pool of engorged larvae). This strategy proved successful because, in almost all cases, the unusual features of the trombicid mite genomes were shared between the two sequenced taxa. In contrast with other acariform mites, the trombicid mite genomes were substantially larger, contained a greater proportion of repeats, and exhibited expansions of mobile elements. These features, coupled with heterozygosity and host contamination in the case of *L. deliense*, proved challenging for accurate genome size estimation but did not prevent the annotation of protein-coding genes, which was sufficient (even for *L. deliense*) for an initial protein expression study.

Table 3: Orthologues of *Ixodes scapularis* salivary proteins in trombiculid mite genomes

Orthologous cluster	Number of genes in cluster ^a			Representative gene ID	Representative annotation ^b
	<i>I. scapularis</i>	<i>L. deliense</i>	<i>D. tinctorium</i>		
ORTHOMCL5799	1	2 (0)	1 (0)	EEC06447	CNDP dipeptidase
ORTHOMCL8488	1	1 (0)	1 (0)	EEC03480	Uncharacterized protein (peptidase M17, leucine aminopeptidase/peptidase B domain)
ORTHOMCL9	73	2 (0)	2 (0)	EEC08659	Sulfotransferase
ORTHOMCL2940	2	1 (0)	2 (2)	ISCW018873	Short-chain alcohol dehydrogenase
ORTHOMCL8622	1	1 (1)	1 (1)	JAA64643	Glycine-rich cell wall structural protein, partial (<i>Rhipicephalus pulchellus</i>)
ORTHOMCL3018	1	1 (0)	3 (0)	EEC14134	Calponin
ORTHOMCL5249	2	1 (0)	1 (0)	JAB84323	Stretch-regulated skeletal muscle protein (<i>Ixodes ricinus</i>)
ORTHOMCL7670	1	1 (1)	1 (0)	EEC14106	Multifunctional chaperone
ORTHOMCL8293	1	1 (0)	1 (0)	EEC09272	Uncharacterized protein (aspartate dehydrogenase domain)
ORTHOMCL262	5	5 (4)	5 (4)	ISCW002881	UDP-sugar hydrolase
ORTHOMCL2418	1	1 (0)	4 (4)	EEC13206	Cytochrome C
ORTHOMCL5815	2	1 (0)	1 (0)	ISCW011750	Succinyl-CoA ligase beta subunit
ORTHOMCL3911	1	1 (0)	3 (0)	JAB73948	Formyltetrahydrofolate dehydrogenase, partial (<i>Ixodes ricinus</i>)
ORTHOMCL3015	2	1 (0)	2 (2)	ISCW001951	ATPase inhibitor
ORTHOMCL8	66	12 (6)	7 (0)	EEC10817	Acyl-CoA synthetase
ORTHOMCL260	13	1 (1)	1 (0)	ISCW001079	Acetylcholinesterase
ORTHOMCL6164	2	1 (0)	1 (0)	ISCW022662	NS1-binding protein
ORTHOMCL4904	2	1 (1)	1 (0)	ISCW006538	60S acidic ribosomal protein LP1
ORTHOMCL907	6	1 (0)	1 (0)	EEC05404	Translation initiation inhibitor UK114/IBM1
ORTHOMCL4900	2	1 (0)	1 (1)	JAB75945	Ribosomal protein LP2 (<i>Ixodes ricinus</i>)
ORTHOMCL3820	3	1 (1)	1 (1)	ISCW002028	IMP-GMP specific 5'-nucleotidase
ORTHOMCL4006	1	1 (0)	3 (0)	EEC13628	Urocanate hydratase
ORTHOMCL3946	1	1 (0)	3 (0)	EEC20451	Uncharacterized protein (pseudouridine-5'-phosphate glycosidase domain)
ORTHOMCL5111	2	1 (0)	1 (0)	ISCW010907	Formiminotransferase-cyclodeaminase
ORTHOMCL75	27	1 (0)	0	EEC02489	Cysteine-rich secreted protein (trypsin inhibitor-like cysteine rich domain)
ORTHOMCL8850	2	1 (1)	0	ISCW012704	Signal sequence receptor beta

^aNumbers in parentheses refer to genes identified with either classic signal peptides or nonclassic internal secretion signatures by SecretomeP 2.0 [94].

^bAnnotations are from *I. scapularis* unless otherwise stated. Protein domain information was obtained from the Conserved Domain Database [131]. Tick salivary protein data were obtained from Kim et al. [122].

Obtaining high-quality genomes for the Chelicerata has proven very challenging across multiple orders. The smaller acariform mite species (e.g., *T. urticae*) have compact genomes, but the low DNA yields from individual specimens require the use of pooling strategies. Conversely, the larger arachnids such as the ticks, spiders, and scorpions tend to have very large (several Gb), highly repetitive genomes, sometimes displaying evidence of whole-genome duplication [142]. Recently, several acarine genomes utilizing long-read technology have been published. One of these had the advantage of using DNA from a continuous tick (*I. scapularis*) cell line and achieved a BUSCO score of 95% [143]. In contrast, the current draft assembly for the cattle tick (*Rhipicephalus microplus*) used relatively little long-read data in a hybrid assembly and attained a BUSCO gene score of only ~40% (including fragmented genes), although it should be emphasized that the estimated genome size for this species is enormous (7.1 Gb) and it contains 70% repetitive DNA [144]. Finally, a Pacific Biosciences assembly for the tiny genome (63.2 Mb) of another acariform mite (*Psoroptes ovis*) has been published with a BUSCO score of 91% but used a pool of >3,000 individuals to

obtain sufficient DNA [145]. This would never be a practicable option for chiggers. However, in future, a long-read assembly might be possible from a single *D. tinctorium* adult individual due to their massive size.

Importantly, as analyses using the BUSCO Arthropoda gene set consider highly conserved housekeeping genes that are single copy in insects (the vast majority of arthropod genomes are from this single class), it will underestimate lineage-specific expansions (rates of gene duplication) between taxa involving novel genes, while flagging duplications of conserved genes in chelicerates that may have a biological origin rather than being assembly artifacts. Elevated duplication rates often signify assembly errors caused by high levels of heterozygosity in the starting material, particularly when samples are pooled [63]. While DNA extraction from single individuals can mitigate these problems, species with large population sizes may display high heterozygosity at the individual level [146], and/or additional haplotypes may originate from developing embryos or stored sperm. Unfortunately, as *D. tinctorium* is extremely difficult to sex by morphological criteria, it is possible that the individual

that we sequenced was a fertilized female, perhaps explaining the high duplication rate flagged by the BUSCO analysis for this genome.

To explore duplication in the *D. tinctorium* genome further, we used the Redundans pipeline [63] and reciprocal BLAST of genomic scaffolds. This demonstrated conclusively that duplication in this genome is not caused by elevated heterozygosity but by identical gene copies located on distinct scaffolds, even if these scaffolds sometimes differed by only a few bases. While assembly errors cannot be excluded in the absence of structural context provided by long-read data, possible biological explanations for this surprising result include recent ectopic recombination of a chromosome segment or chromosome abnormalities (aneuploidy or B chromosomes [147]). This would be compatible with the higher repeat content in this genome compared with that for other acariform mites, which might facilitate large-scale duplications. Notably, whole genome duplication has been reported in other chelicerates, although this is thought to be ancient in the spider-scorpion lineage [142].

As previously reported on publication of the first spider genomes [64], the Acari are polyphyletic, with the superorder Parasitiformes (i.e., the ticks together with mesostigmatid and holothyrid mites) apparently more closely aligned to the spiders (order Araneae) than to the Acariformes (Fig. 1). Recent phylogenomic studies have demonstrated that certain arachnid orders, including the Acariformes and Parasitiformes, are evolving at accelerated rates, leading to long-branch attraction artifacts [148]. This can lead to paraphyletic models of arachnid evolution, including the nested placement of the horseshoe crabs (order Xiphosura) within the Arachnida as we observed here, and a node age for divergence from the Araneae (263.2–394.7 Mya; Fig. 4) that appears much too recent compared with the oldest xiphosuran fossil, which has been dated to the late Ordovician [149]. However, other key nodes, such as the divergence between the Acariformes and the (Araneae + Parasitiformes), as well as the Araneae-Parasitiformes split, were dated within the range of previous estimates based on the fossil record [150] or, more recently, nuclear genomes [64]. While the controversial phylogeny of the Chelicerata at higher taxonomic levels is beyond the scope of our study, at finer scales, the phylogenomic analyses supported the conventional morphology-based taxonomy within the Acariformes, confirming that the trombidid mites are closely related to the phytophagous Tetranychoidea [15]. Moreover, based on an earlier analysis using nuclear and mitochondrial markers [151], our dating of the split between the Tetranychoidea and the Trombididae is consistent with a secondary radiation within the Trombidiformes that is hypothesized to have occurred during the Carboniferous.

Interestingly, within the Trombididae, the divergence between the velvet mites and the chiggers apparently occurred much later (~133 Mya) than the emergence of the earliest terrestrial vertebrates (395 Mya [152]), coinciding perhaps with the appearance of crown-group mammals. If this scenario is correct, the ectoparasitism of non-mammalian vertebrates by chiggers that occurs today may be a product of secondary adaptation, as suggested by other authors [21]. Although the fossil record is devoid of trombidid mite specimens predating the Eocene [153], it has been speculated from palaeogeographical and comparative morphological evidence that trombiculid mites fed initially on other arthropods, with larval ectoparasitism on vertebrates evolving during the Paleocene, leading to an increase in chigger diversity [21]. Our data challenge this hypothesis, because it would imply that something other than host choice in the larval stage drove the split between the Trombidioidea and the Trombiculidoidea 60

million years before the latter began feeding on vertebrates. Only the discovery of more ancient trombidid fossils will help to resolve these uncertainties.

Potential roles for terpenes in trombidid mite biology

The most striking finding in the trombidid mite genomes was the presence of large families of laterally transferred terpene synthases. As the level of amino acid identity between the trombidid terpene synthases and their closest homologues in microbes is quite low, their end products cannot be inferred with any confidence. However, the question of whether compounds such as 2-methylisoborneol, geosmin, or germacrene might confer adaptive advantages to trombidid mites helps to frame hypotheses for experimental testing. Interestingly, all of these compounds are associated with odors and tastes that humans, and some arthropods, may sense as unpleasant or aversive. For instance, 2-methylisoborneol has a musty odor that humans associate with ripe cheeses [154]; whereas geosmin confers the smell of moist soil and is the cause of the muddy, “off” taste that the human olfactory system detects in spoiled water, wine, and the flesh of certain freshwater fish [155–157]. More importantly, geosmin released by *Penicillium* spp. or *Streptomyces* spp. on rotting fruit is strongly aversive to *Drosophila* because these organisms produce secondary metabolites that are directly toxic to the fly or to its primary food source, yeast [158]. Germacrene has also been implicated as an arthropod repellent among complex sesquiterpene mixtures found in the essential oils of various plants, which have been shown to be effective against several acarines, including the ticks *Rhipicephalus microplus* and *Ixodes ricinus* [159, 160] and the poultry red mite *Dermanyssus gallinae* [160]. Although it has not been assayed in isolation, germacrene is additionally a significant component of essential oils or crude leaf extracts that exhibit toxic effects against phytophagous mites (*Brevipalpus phoenicis* in the Trombidiformes [161]) and ants (*Solenopsis invicta* [162]).

These potential repelling and/or toxic effects of terpenoids would align closely with the apparent aposematic nature of trombidid mites, most species of which are brightly colored due to their carotenoid content. These mites have few natural enemies and have been reported to be rapidly regurgitated if offered to predators in the laboratory [163]. However, cannibalism between adults and even ectoparasitism of the free-living stages by trombidid larvae can occur, underlining the relevance of chemical communication within species and between closely related species [19]. To the best of our knowledge, only one report of a parasitoid affecting trombidid mites has been published (the acrocerid fly *Pterodontia flavipes* attacking *Podothrombium* spp. [164]), but the dearth of research on free-living trombidid stages means that no doubt other parasitoids exploiting these hosts do exist. It is important to note that not all mites are repelled by terpenoid compounds. Many plants use terpenoids as defense compounds to signal to the natural enemies of pest arthropods that the plant is under attack. For example, *Lotus japonicus* infested with *T. urticae* releases several terpenoids, including germacrene, that attract the predatory mite *Phytoseiulus persimilis* [165]. Communication by sex pheromones is also known to occur in *T. urticae*, although molecules other than terpenoids are suspected to mediate this [166]. Moreover, as noted above, neryl formate is an aggregation pheromone in dust mites [76]. In conclusion, while the conferment of a foul taste (and perhaps odor) to the aposematic trombidids appears to be the most likely evolutionary driver of *de novo* terpenoid synthesis capability, it is also possible that these compounds are used to commu-

nicate with potential mates during courtship or to repel members of the same (or closely related) species to reduce competition or deter cannibalistic behavior.

Diapause and seasonality

Many of the putative functions of laterally transferred and/or expanding gene families in the trombiculid mites appear to be relevant to the temporal regulation of the life cycle and the switching of metabolic demands between dormant and active stages. The life history of trombiculid mites features alternation between immobile calyptostases (the deutovum, protonymph, and tritonymph) and the active instars (larva, deutonymph, and adult) [167]. The calyptostases typically persist for 25–30 days, while the active stages in temperate species can undergo hibernation over the winter months, including larvae that have not fed until late in autumn. Diapause of eggs is common in temperate species and can exceed one year in the chigger, *Hirsutiella zakhvatkini*, without loss of viability [21]; while larvae of this species can also overwinter on their rodent hosts [168]. The life cycle of trombiculid mites in tropical and subtropical regions has been little studied, but several generations per year are possible [21]. In the case of *Dinothrombium* spp., although the adults are positively phototactic and diurnal if humidity is high (becoming crepuscular during drier conditions), circadian cycles of activity were maintained if mites were transferred to constant darkness in the laboratory [163]. Remarkably, adult *D. pandorae* of the Californian deserts may only emerge from their burrows during rainstorms to feed and mate for a few hours each year, migrating to the deepest extent of their subterranean refuges during the height of summer [44]. However, overcoming torpor by rapidly adjusting metabolic rate after the cold desert night to the early morning warmth when termite prey become active is critical to the life cycle of *D. pandorae* [44]. It may also be important for chiggers to avoid cool microclimates (and thus maintain peak metabolism) when questing for small mammals, since their hosts are highly motile and a suitable location for attachment must be targeted rapidly before grooming behavior leads to ingestion of the mite. Hence, the small expansions in the IR repertoire (IR25a, IR21a, and IR93a) that we observed in the trombiculid mite genomes might reflect more acute sensitivity to cool temperatures than for the phytophagous *T. urticae*.

With relevance to both regulation of metabolism and circadian cycles, the homologues of FLVCR proteins have been little studied in arthropods, but in vertebrates they export heme from the cytoplasm to the extracellular milieu (for FLVCR1) and from mitochondria into the cytoplasm (for FLVCR2) [169]. A FLVCR gene homologue in *Drosophila melanogaster*, CG1358, is involved in maintenance of circadian rhythms in the absence of light together with other genes with roles in iron metabolism [170]. Thus, it is intriguing that FLVCR homologues were significantly expanded in the *D. tinctorium* genome but not that of *L. deliense*, as adult chiggers in tropical environments exhibit more regular activity above ground than do *Dinothrombium* spp. [21]. Furthermore, the proteomic analysis of *D. tinctorium* revealed overrepresentation of putative digestive enzymes with peptidase M20 and inhibitor I29 domains, which is consistent with an imperative for adult velvet mites to obtain food reserves rapidly while foraging briefly above ground [44]. An elevated metabolic rate for *L. deliense* larvae was also suggested by the preponderance of mitochondrial enzymes responsible for aerobic energy production in protein extracts. Further studies on trombiculid mite metabolism are clearly warranted, as most metabolic studies in the Trombidiformes have focused on winter diapause in *T. ur-*

ticae [171], creating a knowledge gap around the physiology of tropical species.

In spider mites, carotenoids are essential for the control of diapause and sexual reproduction. However, reproductive behavior between the Tetranychidae and the trombiculid mites is radically different, since male *T. urticae* become developmentally arrested in close proximity to dormant female deutonymphs on leaf surfaces. This “guarding” behavior is stimulated by the intensity of the yellow coloration of the dormant females (derived from their carotenoid pigments) and allows the male to mate with the adult female immediately after ecdysis [172]. In contrast, in order to be inseminated, adult females of trombiculid mites must collect a spermatophore deposited on the ground by the male, the location of which is signposted by signaling threads. While coloration is not known to be a factor during courtship, the males of some species deposit spermatophores in specially constructed “gardens” and perform encircling dances with the female [19]. However, using genetic manipulation of a laterally transferred phytoene desaturase, it has recently been discovered that carotenoids have a second, distinct function in the regulation of diapause in *T. urticae* [13]. A lack of phytoene desaturase activity not only results in albinism but prevents overwintering strains from entering diapause, probably due to disrupted light (and thus photoperiod) perception caused by vitamin A deficiency.

To the best of our knowledge, anatomical and experimental studies on trombiculid mite vision have not been performed, but spider mites once again provide a closely related template. The eyes of *T. urticae* have been partially characterized and show bi-convex lenses in the anterior pair and simplex convex lenses in the posterior pair [173]. It has been proposed that the anterior eyes respond to ultraviolet (UV) and green light, whereas the posterior pair are sensitive to UV light only. However, laser ablation experiments have demonstrated that either pair can receive sufficient information to control the photoperiodic termination of diapause, while removal of both pairs prevents diapausal exit [174]. In *T. urticae*, the expression pattern of rhodopsins in the eyes has not been determined, but in the jumping spider *Hasarius adansoni*, green-sensitive rhodopsin-1 expressed in the lateral ocelli is important for the monochromatic detection of movement [175]. Until recently, the function of rhodopsin-7 was enigmatic, but experiments in *D. melanogaster* have shown that it operates in the circadian pacemaker neurons of the central brain and is responsible for their highly sensitive response to violet light [176]. Taken together, these experimental findings from other arthropods suggest that trombiculid mites might depend on rhodopsin-7 homologues rather than peropsins for control of circadian rhythms (although we failed to identify orthologues of the *Drosophila* clock gene in Acari). The relative roles of FLVCR gene homologues, phytoene desaturases, and the rhodopsin pigments in the control of diapause and other life history traits in trombiculid mites is evidently a key priority for future experimental studies.

Immune response and vector biology

As trombiculid mites are edaphic organisms with a parasitic larval stage, they are exposed to soil microorganisms, the exterior flora of their hosts, and pathogens contained in ingested body fluids. In the case of certain trombiculid mites (especially *Leptotrombiculidium* spp.), their role as biological vectors of *O. tsutsugamushi* highlights a specific infectious challenge that has resulted from feeding on small mammals. In other arthropods, CTLD proteins with lectin activity act as transmembrane or se-

creted pattern recognition receptors that bind to carbohydrates on the surface of pathogens. They can function as opsonins in the hemolymph that agglutinate unicellular pathogens and facilitate their phagocytosis by hemocytes [177] or may be expressed on the surface of tissues that form a barrier to infectious assaults, such as in the gut or on the gills of crustaceans [93, 178]. These findings are compatible with roles as secreted opsonins for most of the CTLD proteins identified in the current study, with a smaller number perhaps operating as immune surveillance receptors on the surface of cells or extracellular matrices.

Significant exposure to viral nucleic acids was suggested by the striking diversity of ERV-related sequences in the *L. deliense* genome. Among these, more than 20 Gag-like polyprotein genes of potential mammalian origin were identified. The fact that all other classes of retroviral proteins in the *L. deliense* assembly were closely related to arthropod sequences, and the phylogenetic proximity of the Gag-like polyproteins to ERV elements in non-sciurid mammals, rendered host DNA contamination in the mite gut or on mouthparts an unlikely source for these sequences; although this possibility cannot be excluded entirely. Despite this caveat, lateral transfers in the distant past originating from mammalian body fluids during the brief ectoparasitic stage is a working hypothesis that can be tested when a more contiguous chigger genome becomes available. In contrast, the horizontal transfer of the long-interspersed element BovB is postulated to have involved an opposite transmission route, i.e., between vertebrates by ticks [179].

To the best of our knowledge, no experimental studies on the immune response of trombiculid mites have been performed to date. However, a recent study in *T. urticae* involving experimental challenge with bacteria demonstrated that the spider mites were highly susceptible to systemic infection [180]. This contrasted with a more robust response to bacterial challenge in another acariform mite, *Sarcassania berlesei*, which unlike *T. urticae* has a saprophytic lifestyle. The authors of this study concluded that the ecology of spider mites, in which all life-cycle stages feed on plant phloem (a relatively aseptic food source), has led to a high degree of susceptibility to pathogen exposure. This failure to overcome infectious insults was associated with an apparent absence of many antimicrobial protein effectors in the spider mite genome. In support of the hypothesis that spider mites have adapted to an environment characterized by very low levels of pathogen challenge, we found that compared with the *L. deliense* genome, the *T. urticae* genome displays a relative paucity of PGRP (Supplementary Fig. S10), CTLD (Supplementary Table S5a), and Dscam genes (Supplementary Fig. S9). Thus, the common ancestor of the Trombidiformes may have harbored a diverse immune gene repertoire that was selectively lost in the branch leading to the Tetranychoidea and/or the Trombiculidoidea have undergone more recent immune gene family expansions. The intermediate immune gene repertoire of the Trombiculidoidea between that of the spider mites and the chiggers (*D. tinctorium* has considerably fewer CTLD proteins than *L. deliense*) suggests that both immune-related gene losses and gains have occurred during the evolution of the Trombidiformes in response to their radically different natural histories. Indeed, in terms of the degree of exposure to pathogen diversity and abundance, the euedaphic velvet mites are likely to encounter greater infectious challenges than spider mites, and the feeding behavior of chiggers on vertebrates is likely to exacerbate this exposure further compared to their relatives that are ectoparasitic on other invertebrates only.

Conclusions

This first analysis of trombiculid mite genomes has revealed their dynamic nature relative to those of other acariform mites, including expansions in laterally transferred gene families and mobile elements. These genomes provide a foundation for fundamental experimental studies on mite immune responses, host-seeking behavior and feeding, and environmental impacts on life-cycle progression. The function of the laterally transferred terpene synthases will become a major research theme for chigger biology, as only experimental exposure of chiggers and their potential natural enemies to mite terpene extracts will be able to determine if these unique aspects of secondary metabolism have evolved to attract conspecifics or, conversely, to repel predators, parasitoids, and/or competitors. From an applied perspective, the identification of predicted allergens in the *L. deliense* genome sets the scene for immunoproteomic studies of trombiculiasis in both humans and domestic animals, with the potential for immunotherapeutic approaches to be developed as for dust mite allergy [181]. Finally, the successful development of recombinant vaccines against ticks [182] and the promising progress of recombinant vaccine development for both sheep scab [183] and poultry red mite [184] indicate that a similar approach could be explored for chiggers, which has the potential to interrupt, or at least reduce, the transmission of *O. tsutsugamushi* to humans in scrub typhus-endemic areas. Considering the high strain variability of the scrub typhus agent [185], a chigger vaccine utilizing mite salivary or gut antigens could provide a much-needed breakthrough against this intractable disease. Thus, a key priority for future research on this enigmatic mites is to obtain organ-specific proteomic data, as whole-body proteomes such as those analyzed here may overlook important proteins required for host immunomodulation or pathogen transmission that only reach high concentrations in certain anatomical compartments.

Materials and Methods

Sample collection and DNA extraction

Adult specimens of giant red velvet mites were collected within the grounds of the UK Medical Research Council Field Station at Wali Kunda, The Gambia (13°34'N, 14°55'W), in June 2010. Mites were sampled from flower beds following heavy rains in June 2010 and stored in 95% ethanol at -80°C. They were identified as *Dinothrombium tinctorium* by Joanna Małol (Wroclaw University of Environmental and Life Sciences, Poland). Approximately 5 µg of DNA was extracted from a single individual using a Genomic tip Kit (Qiagen) according to the manufacturer's instructions. Integrity of the DNA was confirmed by agarose gel electrophoresis, which showed a single band of ~20 kb.

For *L. deliense*, engorged larvae were collected from two Berdmore's ground squirrels (*Menetes berdmorei*) captured in Udon-thani Province, Thailand, in September 2015. Trapping and euthanasia of small mammals followed the CERoPath (Community Ecology of Rodents and their Pathogens in a changing environment) project protocols [186]. Chiggers were located inside the ears and the inguinal area of the squirrels and stored in absolute ethanol at -20°C. A subsample of the mites was selected and mounted in clearing medium, Berlese fluid (TCS Bioscience, UK), prior to species identification under a compound microscope. Fifty unmounted larvae were pooled and ~30 ng of genomic DNA was extracted using a DNeasy Blood & Tissue Kit (Qiagen) according to the manufacturer's instructions. The DNA

was partially degraded, but a dominant band of ~5 kb was apparent by agarose gel electrophoresis.

Library preparation and sequencing

These steps were performed at the Centre for Genomic Research at the University of Liverpool. The *D. tinctorium* DNA was used to generate two Illumina TruSeq libraries and one Nextera mate-pair library. For the former, bead-based size selection using 100 ng and 200 ng of DNA as input into the TruSeq DNA LT Sample Prep Kit with 350 and 550 bp inserts, respectively, was applied. Following eight cycles of amplification, libraries were purified using Agencourt AMPure XP beads (Beckman Coulter). Each library was quantified using a Qubit fluorimeter (Life Technologies), and the size distribution was assessed using a 2100 Bioanalyzer (Agilent). The final libraries were pooled in equimolar amounts using the Qubit and Bioanalyzer data. The quantity and quality of each pool were assessed on the Bioanalyzer and subsequently by qPCR using the Illumina Library Quantification Kit (KAPA Biosystems) on a LightCycler 480 instrument II (Roche Molecular Diagnostics) according to the manufacturer's instructions. The pool of libraries was sequenced on one lane of the HiSeq 2000 with 2 × 100 bp PE sequencing and v3 chemistry.

The *D. tinctorium* mate-pair library was constructed using the Nextera Mate Pair Kit (Illumina) with 3 kb inserts. The DNA (3 µg) was fragmented as described in the manufacturer's protocol and cleaned using a Genomic DNA Clean & Concentrator column (Zymo Research). The sample was then subjected to strand displacement and cleaned with AMPure XP beads. A 0.6% Certified Megabase Agarose gel (Bio-Rad) was used to separate the fragments, and those in the range of 2–5 kb were extracted and recovered using a Zymoclean Large Fragment DNA Recovery Kit (Zymo Research). The recovered DNA was quantified and transferred into a circularization reaction at 16°C overnight. After purification with AMPure XP beads, DNA was sonicated into ~500 bp fragments using a focused ultrasonicator (Covaris) and recovered with AMPure XP beads as before. Samples were bound to Dynabeads M-280 Streptavidin (Thermo Fisher Scientific) and all subsequent reactions (end repair, A-tailing, and adapter ligation) were bead based. Samples were amplified with 10 cycles of PCR, recovered by AMPure XP beads at a 1:1 ratio, and quantified using the Qubit dsDNA HS Assay Kit. The library was then subjected to quality control on the Bioanalyzer and LightCycler as for the TruSeq libraries above. The library was sequenced on one run of the MiSeq with 2 × 250 bp PE sequencing.

For *L. deliense*, the DNA sample was sheared to 550 bp using a Bioruptor Pico sonication device (Diagenode) and purified using an AxyPrep FragmentSelect-I Kit (Axygen). The sample was then quantified using a Qubit dsDNA HS Assay Kit on the Qubit fluorimeter, and the size distribution was ascertained on the Bioanalyzer using a High Sensitivity DNA chip (Agilent). The entire sample was used as input material for the NEB Next Ultra DNA Library Preparation Kit. Following nine PCR cycles, the library was purified using the AxyPrep kit and quantified as before by Qubit. Library size was determined on the Bioanalyzer (Agilent). The quality and quantity of the pool were assessed as described above for *D. tinctorium*. The sequencing was conducted on one lane of an Illumina MiSeq with 2 × 150 bp PE sequencing and v2 chemistry.

Assembly and annotation

For both genomes, base-calling and de-multiplexing of indexed reads was performed by bcl2fastq v. 1.8.4 (Illumina; bcl2fastq,

[RRID:SCR_015058](#)) to produce sequence data in fastq format. The raw fastq files were trimmed to remove Illumina adapter sequences using Cutadapt version 1.2.1 [187]. The option “-O 3” was set so the 3' end of any reads that matched the adapter sequence over at least 3 bp was trimmed off. The reads were further trimmed to remove low-quality bases using Sickle version 1.200 [188] with a minimum window quality score of 20. After trimming, reads shorter than 10 bp were removed. If both reads from a pair passed this filter, each was included in the R1 (forward reads) or R2 (reverse reads) file. If only one of a read pair passed this filter, it was included in the R0 (unpaired reads) file.

The Kraken taxonomic sequence classification system (v. 1.0) [189] with the standard (Mini-Kraken) database was used to assign taxonomic annotations to the reads and specifically to estimate the proportion of reads of bacterial origin in the raw data. For genome size estimations, k-mers were counted by the Jellyfish program (v. 2.0) [190], and the resultant histograms were uploaded to GenomeScope (v. 1.0) [191] to visualize the k-mer distributions.

For *D. tinctorium*, the PE reads were assembled using Abyss (v. 1.5.2) [49, 50], Allpaths-LG (v. r51279) [192] (ALLPATHS-LG, [RRID:SCR_010742](#)), SOAPdenovo2 (v. 2.04-r240) [193] (SOAPdenovo2, [RRID:SCR_014986](#)), and Discovar (v. r51454) [194]. When running Abyss, k-mer sizes were from 35 bp to 80 bp with an interval of 5; the output “k-mer size = 80 bp” was selected as this produced the optimal assembly. Allpaths-LG and Discovar specify k-mer size automatically, whereas “k-mer size = 63” was selected for SOAPdenovo2, as suggested by the developer. Discovar requires read lengths of 250 bp, so was applied only to the data generated from the mate-pair library. Assessment of the completeness of the genome assemblies was based on the percentage alignment obtained against the reads from the TruSeq 350-bp insert library using bowtie2 (v. 2.0.10) [195] and the predicted core gene content determined by CEGMA (Core Eukaryotic Genes Mapping Approach, [RRID:SCR_015055](#)) (v. 2.5) [196]. The final, optimum assembly was created by Abyss (97.4% of scaffolds >500 bp were mapped; 99.2% of key orthologs for eukaryotic genomes were present) and included all scaffolds of ≥1,000 bp, which constituted ~80% of the total length of the raw assembly. The impact of removing small scaffolds was assessed using blobtools (v0.9.19), which generates a GC-coverage plot (proportion of GC bases and node coverage; Supplementary Fig. S11b) [52]. Read coverage was calculated by mapping the reads onto the final *D. tinctorium* genome assembly. The excluded small scaffolds tended to be high coverage, high GC, and of eukaryotic origin, perhaps representing gut contents (Supplemental Fig. S11a). We also applied Redundans (v. 0.13c) [63] to identify putative redundant scaffolds in the final *D. tinctorium* assembly with the “-noscaffolding” and “-nogapclosing” options.

For *L. deliense*, a preliminary genome assembly at the contig level was performed using Velvet (v. 1.2.07) [51], with parameters of “best k-mer 99” and “-ins.length 500.” Reads derived from mammalian host genomic DNA were filtered from the preliminary genome assembly using blobtools (v0.9.19) (Fig. 3). Reads mapped on the contigs annotated as Chordata or with a node coverage of <3 were removed, while all others annotated as Arthropoda were retained. The filtered *L. deliense* reads were then reassembled using SPAdes assembler (v3.7.1) [53] (SPAdes, [RRID:SCR_000131](#)) with default settings and the length cutoff for scaffolds set at 500 bp.

To find, classify, and mask repeated sequences in the mite genome assemblies, a *de novo* repeat library was first built using RepeatModeler (v. 1.0.8) [197] (RepeatModeler, [RRID:SCR_015027](#)) with the “-database” function, followed by application of Re-

peatMasker (v. 4.0.6) [198] (RepeatMasker, RRID:SCR_012954) using default settings for *de novo* repeated sequences prediction. Then, a homology-based prediction of repeated sequences in the genome was achieved using RepeatMasker with default settings to search against the RepBase repeat library (issued on 7 August 2015). For non-interspersed repeated sequences, RepeatMasker was run with the “-noint” option, which is specific for simple repeats, microsatellites, and low-complexity repeats.

The *ab initio* gene prediction programs, including Augustus (v. 3.2.2) [55] (Augustus: Gene Prediction, RRID:SCR_008417), SNAP (v. 2013-11-29) [56], and GeneMark (v. 2.3e) [57] (GeneMark, RRID:SCR_011930), were used for *de novo* gene predictions in each genome assembly. Augustus and SNAP were trained based on the gene structures generated by CEGMA (v. 2.5) [196], whereas GeneMark [57] was self-trained with the “-BP OFF” option. The three *ab initio* gene prediction programs were run with default settings. We also generated an integrated gene set for each genome assembly using the MAKER v. 2.31.8 [54] pipeline (MAKER, RRID:SCR_005309). The MAKER pipeline runs Augustus, SNAP, and GeneMark to produce *de novo* gene predictions and integrates them with evidence-based predictions. These were generated by aligning the invertebrate RefSeq protein sequences (downloaded on 31 March 2016 from NCBI) to the masked mite genomes by BLASTX. The MAKER pipeline was run with “-RM.off” option to turn all repeat masking options off; all parameters in control files were left in their default settings. Genes identified by *de novo* prediction, which did not overlap with any genes in the integrated gene sets, were also added to the final gene set for each genome assembly if they could be annotated by InterProScan (v. 4.8) [199] (InterProScan, RRID:SCR_005829) with the InterPro superfamily database (v. 43.1) using “-appl superfamily -nocrc” options.

The Blast2GO pipeline (v. 2.5) [200] (Blast2GO, RRID:SCR_005828) was used to annotate proteins by Gene Ontology (GO) terms. In the first step, all protein sequences were searched against the nr database with BLASTP. The E-value cutoff was set at 1×10^{-6} , and the best 20 hits were used for annotation. Based on the BLAST results, the Blast2GO pipeline then predicted the functions of the sequences and assigned GO terms to the BLAST-based annotations. Metabolic pathways were constructed using the Kyoto Encyclopedia of Genes and Genomes Automatic Annotation Server [201] with the recommended eukaryote sets, all other available insects, and *I. scapularis*. The pathways in which each gene product might be involved were derived from the best KEGG Orthology (KO) hit with the bi-directional best hit method.

Phylogenetics

Protein datasets of the following arthropod genomes were used as references: *D. melanogaster* (fruit fly; OGS release: 6.11) [202], *A. mellifera* (honey bee; OGS release: 3.2) [203], *T. mercedesae* (bee mite; OGS release: v. 1.0) [99], *T. urticae* (spider mite; OGS release: 20 150 904) [61], *Stegodyphus mimosarum* (velvet spider; OGS release: 1.0) [64], *I. scapularis* (blacklegged tick; OGS release: 1.4) [204], *M. occidentalis* (predatory mite; OGS release: 1.0) [98], and *Limulus polyphemus* (Atlantic horseshoe crab; OGS release: 2.1.2) [205]. *Caenorhabditis elegans* (nematode; OGS release: WS239) [206] was used as the outgroup. For gene-family phylogenetics, we first aligned orthologous protein sequences with Mafft (v. 7.309) [207] or Kalign (v. 2.0) [208]. We manually trimmed the aligned sequences for large gene sets. The best substitution models of amino-acid substitution were determined for the alignments by ProtTest (v. 3.4) with parameters set to “-all-matrices, -all-distributions, -AIC” [209]. Then, phylogenetic trees

were constructed using maximum likelihood methods (PhyML, v. 3.1) [210] (PhyML, RRID:SCR_014629). In addition, a neighbor-joining method was used for building the distance-based trees using MEGA (v. 7.021) [211] (MEGA Software, RRID:SCR_000667).

For species-level phylogenetics, the rapid evolution of acariform mites may challenge phylogenetic analyses due to long-branch attraction [151]. Thus, we used a very strict E-value (1×10^{-50}) when performing a reciprocal BLASTP to exclude the most variant orthologous genes across all genomes tested. The reciprocal BLAST search resulted in identification of 360 highly conserved one-to-one orthologues in all 12 genomes. Each of these orthologous groups was aligned using Mafft with the “-auto” option. These alignments were trimmed by Gblocks (v. 0.91b) [212] and concatenated into unique protein super-alignments. ProtTest determined the best-fit substitution model of LG with invariant sites (0.133) and gamma distributed rates (0.878) using parameters as above prior to conducting the phylogenetic analysis with PhyML. Based on the topology defined by this phylogenetic analysis, we estimated the divergence time of each species using the Bayesian MCMC method in the PAML package (v. 4.9a) [213] (PAML, RRID:SCR_014932) with the correction of several fossil records (time expressed in million years ago): Chelicera-Mandibulata: 514–636 [214], *T. urticae*-tick-spider: 395–503 [64] (oldest Acari), *A. mellifera*-*D. melanogaster*: 238–307 [215], and nematode-arthropods: 521–581 [215].

Analysis of gene family expansions

Orthologous gene clusters of *D. tinctorium*, *L. deliense*, and the other reference genomes described above were defined based on OrthoMCL (v. 1.4) [216] (OrthoMCL DB: Ortholog Groups of Protein Sequences, RRID:SCR_007839). We used CAFE (v. 3.1) [217] to infer the gene family expansion and contraction in *D. tinctorium* and *L. deliense* against other Acariformes (*T. urticae* and *S. scabiei*). The ultrametric species tree used in the CAFE analyses was created as described for gene-family phylogenetics above.

Analysis of candidate lateral gene transfers

We used a modification of the Crisp method [6] for examination of LGTs in the two mite genomes. Each mite protein dataset was aligned with BLASTP against two databases derived from the NCBI nr database, one consisting of metazoan proteins (excluding proteins from species in the same phylum as the studied species—Arthropoda) and the other of non-metazoan proteins. The LGT index, h , was calculated by subtracting the bit-score of the best metazoan match from that of the best non-metazoan match. The genes can be classified into class C if they gained an h index ≥ 30 and a best non-metazoan bit-score of ≥ 100 . For each class C gene, its average h value (h_{orth}) and that of its paralogous genes in each OrthoMCL cluster defined above was determined. If h was ≥ 30 , the best non-metazoan bit-score was ≥ 100 , and the h_{orth} value was ≥ 30 , the gene was considered to be a class B gene. Class A genes were defined as a subset of class B genes with $h \geq 30$, a best non-metazoan bit-score of ≥ 100 , an h_{orth} value of ≥ 30 , and a best metazoan bit-score < 100 . The class A criterion that no member of the orthologous group had a metazoan match with a bit-score ≥ 100 was not applied, as it was considered too strict for large gene clusters.

Analysis of immune-related gene families

A search for mite immune-related genes was initially preformed with a BLASTP search (E-value, $< 1 \times 10^{-5}$) against each mite

protein set using immune-related genes defined by Palmer and Jiggins [86]. The identified potentially immune-related genes were then manually checked using BLASTP online at NCBI. For analysis of CTLD proteins, FASTA sequences of proteins in ORTHOMCL223, ORTHOMCL584, ORTHOMCL1095, and ORTHOMCL1094 were analyzed by SecretomeP 2.0 [94] (with the “mammalian” option), TMHMM 2.0 [95], and NetNGlyc 1.0 [96] using default settings to identify respective sequence features. Protein sequences were also submitted to InterPro [218] for domain structure analysis. The CTLDs extracted from individual protein sequences were then manually searched for the amino-acid motifs “EPN,” “WND,” and “QPD.”

Analysis of chemosensory and photoreceptor gene families

A search for *D. tinctorium* and *L. deliense* OBPs was initially preformed using TBLASTN (E -value, $<1 \times 10^{-3}$) against their genome assemblies using *D. melanogaster*, *Drosophila mojavensis*, *Anopheles gambiae*, *Bombyx mori*, *Tribolium castaneum*, *A. mellifera*, *Pediculus humanus humanus*, and *Acyrthosiphon pisum* OBPs (identified by Vieira and Rozas [102]) as queries. No OBPs were found in the *D. tinctorium* and *L. deliense* genome assemblies. Because OBPs are very divergent in terms of the amino-acid sequences within the family, and the sequence identities between the family members from the different species can be as low as 8% [102], a TBLASTN search has limited power to identify these genes. A search for OBPs was therefore performed again with BLASTP (E -value, $<1 \times 10^{-3}$) to search the automated protein predictions from the mite genome assemblies. A search for small chemosensory proteins in *D. tinctorium* and *L. deliense* was preformed using the same methods as for the OBPs. The query sequences were also based on the study of Vieira and Rozas [102], using chemosensory protein sequences from *D. melanogaster*, *D. mojavensis*, *A. gambiae*, *B. mori*, *T. castaneum*, *A. mellifera*, *P. humanus humanus*, *A. pisum*, *I. scapularis*, and *Daphnia pulex*. For odorant receptors, both TBLASTN and BLASTP searches were performed using *D. melanogaster* and *A. mellifera* sequences (identified by Nozawa and Nei [219] and Robertson and Wanner [220], respectively) as queries.

The *D. tinctorium* and *L. deliense* GR gene families were manually annotated according to TBLASTN and BLASTP searches (both with an E -value cutoff of $<1 \times 10^{-3}$) against their genome assemblies and predicted protein coding genes, respectively, using all *D. melanogaster* [221], *A. mellifera* [220], *I. scapularis* [204], *T. urticae* [101], *T. mercedesae* [99], and *M. occidentalis* [98] GRs as queries. An iterative search was also conducted with termite GRs as queries until no new genes were identified in each major subfamily or lineage. For phylogenetic analysis, *D. tinctorium* and *L. deliense* GRs were aligned with *D. melanogaster* GRs by Kalign [208] with default settings. Poorly aligned and variable N-terminal and C-terminal regions, as well as several internal regions of highly variable sequences, were excluded from the phylogenetic analysis. Other regions of potentially uncertain alignment between these highly divergent proteins were retained, as removing these regions could potentially compromise subfamily relationships. Based on the trimmed alignment, a PhyML tree was constructed using the substitution model of LG determined by ProtTest [209]. Here, the SH-like local support method was used to assess the significance of phylogenetic clustering.

The *D. tinctorium* and *L. deliense* iGluRs and IRs were manually annotated according to a TBLASTN (E -value cutoff, $<1 \times 10^{-3}$) search against the *D. tinctorium* and *L. deliense* genome assemblies using all iGluRs and IRs identified by Croset et al. [222]

across vertebrates and invertebrates, as well as those identified in the recent *T. mercedesae* genome project [99]. Iterative searches were also conducted with termite iGluRs and IRs as queries until no new genes were identified in each major subfamily or lineage. In the phylogenetic analysis, all manually annotated *D. tinctorium* and *L. deliense* IRs and iGluRs were aligned with the *D. melanogaster* IRs and iGluRs by Mafft [207] using default settings. Phylogenetic analysis proceeded as for the GR genes described above using the more conserved iGluRs to root the tree.

Reference opsin genes were collected based on the work of Nagata et al. [97]. Opsin-like sequences in *M. occidentalis*, *I. scapularis*, and *T. urticae* were obtained from the NCBI database. These opsin genes were classified by phylogenetic analysis using the neighbor-joining method with 1,000 bootstraps. The multiple alignment of the amino acid sequences was carried out using Mafft [207] with the “–auto” option. The gaps deletion of the alignment was set to 75% in MEGA7 [211].

Prediction of allergenic gene families

Allergenic protein-coding genes in the genomes of acariform mites (*D. tinctorium*, *L. deliense*, *T. urticae*, *D. farinae*, *S. scabiei*, and *E. maynei*) were predicted using a stand-alone version of Allerdictor (v. 1.0) [223]. Because the predicted proteome of *D. farinae* is not publicly available, we used protein sequences identified from a new Trinity (v. 2.4) [224] (Trinity, RRID:SCR_013048) assembly for the prediction of allergenic genes. The allergenic gene clusters were constructed using OrthoMCL (v. 1.4) [216]; individual protein sequences were submitted to Pfam (EBI, v.31.0 [225]) to be assigned to protein families. Mapped Pfam domain identifications were then searched against the AllFam database [114] to retrieve corresponding AllFam identifiers for allergen families. Venn diagrams were constructed using InteractiVenn [226].

Salivary and cement protein analysis

A total of 159 nonredundant tick cement proteins were retrieved from the NCBI database and then clustered with all *D. tinctorium* and *L. deliense* amino-acid sequences using OrthoMCL in order to identify the tick cement orthologues in the new genomes. Salivary proteins of *I. scapularis* and *T. urticae* have been previously identified using proteomic methods by Kim et al. [122] and Jonckheere et al. [128], respectively. Proteins that might be present in the saliva of *D. tinctorium* and *L. deliense* were identified by clustering all predicted protein-coding sequences in the new genomes with these *T. urticae* and *I. scapularis* salivary proteins using OrthoMCL [216]. Venn diagrams were constructed using InteractiVenn [226]. To identify putatively secreted proteins, both classic signal peptides and nonclassic internal secretion signatures were detected by SecretomeP 2.0 [94] using the “mammalian” option.

Sample preparation for proteomics

Engorged larvae of *L. deliense* (10 specimens) were pooled from infested rodent specimens (*Bandicota indica*, *Bandicota savilei*, and *Rattus tanezumi*) captured across several provinces in Thailand during the field studies of the CERoPath project [186]. The chigger samples were fixed in 70% ethanol and identified as *L. deliense* using autofluorescence microscopy [227]. For *D. tinctorium*, proteomic analysis was performed on a single ethanol-fixed individual from the same collection used for genome sequencing.

The chiggers were washed with chilled 50 mM ammonium bicarbonate. Soluble protein extracts were prepared by homogenization in 0.1% w/v Rapigest (Waters) in 50 mM ammonium bicarbonate using a polypropylene mini-pestle. This was followed by three cycles of sonication on ice (Vibra-cell 130PB sonicator at 20 Hz with microprobe; 10 seconds of sonication alternating with 30 seconds of incubation on ice). Samples were centrifuged at 13,000 × g for 10 minutes at 4°C. The supernatant was removed and retained. The *D. tinctorium* individual was homogenized using a mini-pestle in lysis buffer (4% SDS, Tris-hydrochloride, pH 7.6) and sonicated and centrifuged as above. Supernatants from both mite preparations were stored at -80°C.

Protein concentrations of the samples were determined using a Bradford protein assay (Thermo Fisher Scientific). The *L. deliense* protein extract was reduced with 3 mM dithiothreitol (Sigma) at 60°C for 10 minutes, cooled, then alkylated with 9 mM iodoacetamide (Sigma) at room temperature for 30 minutes in the dark; all steps were performed with intermittent vortex-mixing. Proteomic-grade trypsin (Sigma) was added as a protein: trypsin ratio of 50:1 and incubated at 37°C overnight. Rapigest was removed by adding trifluoroacetic acid (TFA) to a final concentration of 0.5% (v/v). Peptide samples were centrifuged at 13,000 × g for 30 minutes to remove precipitated Rapigest. The *D. tinctorium* protein extract was reduced, alkylated, and digested with trypsin using the filter-aided sample preparation approach [58]. Peptides from *D. tinctorium* were split into eight fractions using the Pierce High pH Reversed-Phase Peptide Fractionation Kit according to the manufacturer's instructions. Each digest and fraction was concentrated and desalted using C18 Stage tips (Thermo Fisher Scientific), then dried using a centrifugal vacuum concentrator (Eppendorf) and resuspended in a 0.1% (v/v) TFA, 3% (v/v) acetonitrile solution.

Mass spectrometry

Peptides were analyzed by on-line nanoflow LC using the Ultimate 3000 nano system (Dionex/Thermo Fisher Scientific). Samples were loaded onto a trap column (Acclaim PepMap 100, 2 cm × 75 μm inner diameter, C18, 3 μm, 100 Å) at 9 μL/min with an aqueous solution containing 0.1% (v/v) TFA and 2% (v/v) acetonitrile. After 3 minutes, the trap column was set in-line to an analytical column (Easy-Spray PepMap RSLC 50 cm × 75 μm inner diameter, C18, 2 μm, 100 Å) fused to a silica nano-electrospray emitter (Dionex). The column was operated at a constant temperature of 35°C, and the LC system was coupled to a Q-Exactive mass spectrometer (Thermo Fisher Scientific). Chromatography was performed with a buffer system consisting of 0.1% formic acid (buffer A) and 80% acetonitrile in 0.1% formic acid (buffer B). The peptides were separated by a linear gradient of 3.8%–50% buffer B over 90 minutes (*D. tinctorium* and *L. deliense* whole digests) or 30 minutes (*D. tinctorium* fractions) at a flow rate of 300 nL/min. The Q-Exactive was operated in data-dependent mode with survey scans acquired at a resolution of 70,000 at m/z 200. Scan range was 300 to 2,000 m/z. Up to the top 10 most abundant isotope patterns with charge states +2 to +5 from the survey scan were selected with an isolation window of 2.0 Th and fragmented by higher-energy collisional dissociation with normalized collision energies of 30. The maximum ion injection times for the survey scan and the MS/MS scans were 250 and 50 ms, respectively, and the ion target value was set to 1×10^6 for survey scans and 1×10^4 for the MS/MS scans. The MS/MS events were acquired at a resolution of 17,500. Repetitive sequencing of peptides was minimized through dynamic exclusion of the sequenced peptides for 20 seconds.

Protein identification, quantification, and enrichment analysis

Thermo RAW files were imported into Progenesis LC-MS (version 4.1, Nonlinear Dynamics). Peaks were picked by the software using default settings and filtered to include only peaks with a charge state between +2 and +7. Spectral data were converted into .mgf files with Progenesis LC-MS and exported for peptide identification using the Mascot (version 2.3.02, Matrix Science) search engine as described above. Tandem MS data were searched against a database including translated ORFs from either the *D. tinctorium* genome (DinoT_V2.aug2016, 19,258 sequences; 8386,445 residues) and a contaminant database (cRAP, GPMDB, 2012) (119 sequences; 40,423 residues) or the *L. deliense* genome (L_deliense_V2.Aug16, 15,096 sequences; 5183,596 residues), the *Rattus norvegicus* genome (UniProt, Apr16 7948 sequences; 4022,300 residues), and a contaminant database (cRAP, GPMDB, 2012) (119 sequences; 40,423 residues). The search parameters were as follows: the precursor mass tolerance was set to 10 ppm and the fragment mass tolerance was set as 0.05 Da. Two missed tryptic cleavages were permitted. Carbamidomethylation (cysteine) was set as a fixed modification and oxidation (methionine) set as a variable modification. Mascot search results were further validated using the machine-learning algorithm Percolator embedded within Mascot. The Mascot decoy database function was utilized, and the false discovery rate was <1%, while individual percolator ion scores >13 indicated identity or extensive homology (P <0.05). Mascot search results were imported into Progenesis LC-MS as XML files. Fractions were combined using the Progenesis "combine analyzed fractions" workflow. Relative protein abundance was calculated by the Hi-3 default method in Progenesis. Mass spectrometric data were deposited to the ProteomeXchange Consortium [228] via the PRIDE partner repository [229] with the dataset identifier PXD008346.

Enrichment of protein domains was assessed using Pfam (EBI, v.27.0 [225]) as previously described [230] using the gathering threshold as a cutoff. Briefly, a hypergeometric test for enrichment of Pfam domains in the observed proteome (for identifications supported by ≥2 unique peptides only) relative to the complete search database was performed using R (phyper). The Benjamini & Hochberg step-up false-discovery rate-controlling procedure was applied to the calculated P values [231], and enrichment was considered statistically significant where P <0.05.

Availability of supporting data

The datasets supporting the results presented here are available in the GigaDB repository associated with this publication [232]. Raw genomic sequence data have also been submitted to NCBI under BioProjects PRJNA377801 and PRJNA377800 for *L. deliense* and *D. tinctorium*, respectively. Proteomic data for both species are available from the PRIDE repository with the dataset identifier PXD008346.

Additional files

Additional file 1 (.pdf format) contains the following supplemental figures and tables:

Supplemental figure S1: K-mer distributions for *Leptotrombidium deliense* (A) and *Dinothrombium tinctorium* (B) plotted by GenomeScope.

Supplemental table S2: Identification of repetitive sequences in the *Dinothrombium tinctorium* and *Leptotrombidium deliense* assemblies compared with other acariform mites.

Supplemental table S3: Impact of Redundans analysis on genome statistics for *Dinothrombium tinctorium*.

Supplemental figure S2: The number of gene families shared among acariform mites (*Dinothrombium tinctorium*, *Leptotrombidium deliense*, *Tetranychus urticae* and *Sarcopes scabiei*); alongside other references including *Drosophila melanogaster*, *Apis mellifera*, *Tropilaelaps mercedesae*, *Metaseiulus occidentalis*, *Ixodes scapularis*, *Stegodyphus mimosarum* and *Caenorhabditis elegans* by the OrthoMCL classification algorithm.

Supplemental figure S3: Gene family contraction and expansion in 12 species of Ecdysozoa.

Supplemental figure S4: Phylogeny of carotenoid synthases/cyclases from trombiculid mites, spider mites, aphids and fungi.

Supplemental figure S5: Genomic scaffold of *Dinothrombium tinctorium* containing a putative lateral gene transfer adjacent to an incontrovertible mite gene.

Supplemental table S9: Microbial reads identified in the trombiculid genomic data by the Kraken taxonomic sequence classification system.

Supplemental figure S6: Peptides detected by mass spectrometry from two terpene synthases in an adult specimen of *Dinothrombium tinctorium*.

Supplemental figure S7: Phylogeny of reverse ribonuclease integrases in *Dinothrombium tinctorium* and their closest homologues in other taxa.

Supplemental figure S8: Phylogeny of Pol-like polyproteins in trombiculid mites and their closest homologues in other taxa.

Supplemental figure S9: Phylogeny of Dscam protein-coding sequences in *Dinothrombium tinctorium*, *Leptotrombidium deliense*, *Tetranychus urticae* and *Ixodes scapularis*.

Supplemental figure S10: Phylogeny of peptidoglycan recognition protein sequences in *Dinothrombium tinctorium*, *Leptotrombidium deliense*, *Tetranychus urticae* and *Ixodes scapularis* alongside homologous sequences from insects.

Supplemental table S12: Orthologous clusters of tick cement proteins in the genomes of *Dinothrombium tinctorium* and *Leptotrombidium deliense*.

Supplemental figure S11: Blob-plot of *D. tinctorium* genomic scaffolds before (A) and after (B) removal of small scaffolds (<1000 bp).

Additional file 2 (.xlsx format) contains the following supplemental tables:

Supplemental table S1: Genome assembly and gene set statistics compared with 14 other arachnids.

Supplemental table S4: Comparative sequence identity and coverage between *D. tinctorium* genomic scaffolds.

Supplemental table S5a: Changes of gene family size in *L. deliense* in comparison with three other acariform mites.

Supplemental table S5b: Changes of gene family size in *D. tinctorium* in comparison with three other acariform mites.

Supplemental table S6a: High-confidence protein identifications and abundance scores for *L. deliense* engorged larvae.

Supplemental table S6b: All protein identifications and abundance scores for *L. deliense* engorged larvae (includes host contamination).

Supplemental table S7: High-confidence protein identifications and abundance scores for a single adult *D. tinctorium* individual.

Supplemental table S8a: Gene- and scaffold-level sequence coverage of scaffolds containing terpene synthase genes for *D. tinctorium*.

Supplemental table S8b: Gene- and scaffold-level sequence coverage of scaffolds containing terpene synthase genes for *L. deliense*.

Supplemental table S10: RNA families identified in the Rfam database in 10 arthropod genomes.

Supplemental table S11a: Predicted allergenic protein clusters in the *L. deliense* genome with top BLAST hit, Pfam domains and AllFam classifications.

Supplemental table S11b: Predicted unclustered allergenic proteins in the *L. deliense* genome with top BLAST hit, Pfam domains and AllFam classifications.

Abbreviations

BLAST: Basic Local Alignment Search Tool; BUSCO: Benchmarking Universal Single-Copy Orthologues; CERoPath: Community Ecology of Rodents and their Pathogens in a changing environment; CTLD: C-type lectin domain; ERV: endogenous retrovirus; FLVCR: feline leukemia virus subgroup C receptor-related protein; GO: Gene Ontology; GR: gustatory receptor; IMD: immune deficiency; IR: ionotropic receptor; iGluR: ionotropic glutamate receptors; LGT: lateral gene transfer; Mya: million years ago; NCBI: National Center for Biotechnology Information; NEB: New England Biolabs; P: dorant-binding protein; PE: paired-end; PGPR: peptidoglycan recognition protein; qPCR: quantitative polymerase chain reaction; TFA: trifluoroacetic acid; UV: ultraviolet; XIAP: X-linked inhibitor of apoptosis protein.

Animal ethics

Wild rodents trapped during the CERoPath project [186] and used as a source of chigger material were euthanized by inhaled anesthetic overdose according to guidelines published by the American Veterinary Medical Association Council on Research [233] and the Canadian Council on Animal Care [234].

Competing interests

The authors declare that they have no competing interests.

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Author contributions

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