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Molecular evolutionary trends and biosynthesis pathways in the Oribatida revealed by the genome of *Archegozetes longisetosus*

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Original research

ABSTRACT

Oribatid mites are a speciose order of microarthropods within the Chelicerata, comprising about 11,000 described species. They are ubiquitously distributed across different microhabitats in all terrestrial ecosystems around the world and were among the first animals colonizing terrestrial habitats as decomposers and scavengers. Despite their species richness and ecological importance genomic resources are lacking for oribatids. Here, we present a 143-Mb chromosome-level genome assembly of the clonal, all-female oribatid mite species *Archegozetes longisetosus* Aoki, a model species used by numerous laboratories for the past 30 years. Comparative genomic and transcriptional analyses revealed patterns of reduced body segmentation and loss of the segmental identity gene *abd-A* within Acariformes, and unexpected expression of key eye development genes in these eyeless mites across developmental stages. Consistent with their soil dwelling lifestyle, investigation of sensory genes revealed a species-specific expansion of gustatory receptors, and evidence of horizontally transferred genes related to cell wall degradation of plant and fungal matter, both components of *Archegozetes*' diet. Oribatid mites are also noted for their biosynthesis capacities and biochemical diversity. Using biochemical and genomic data, we were able to delineate the backbone biosynthesis of monoterpenes, an important class of compounds found in the major exocrine gland system of Oribatida – the oil glands. Given the mite's strength as an experimental model, the new high-quality resources provided here will serve as the foundation for molecular research in Oribatida and will enable a broader understanding of chelicerate evolution.

Keywords soil animal; terpene synthesis; horizontal gene transfer; parthenogenesis; chemoreceptors; Hox genes; model organism; RNAseq; chromosome-level assembly; Sarcotiformes

Introduction

In the past couple of years, the number of sequenced animal genomes has increased dramatically, especially for arthropods about 500 genomes sequences are now available (Childers, 2020; Thomas *et al.*, 2020). The majority of these genomes, however, belong to the insects (e.g. flies, beetles, wasp, butterflies and bugs (Thomas *et al.*, 2020)) which compromise the most diverse, yet evolutionarily young and more derived taxa of arthropods (Regier *et al.*, 2010; Giribet and

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Edgecombe, 2019). In strong contrast, genome assemblies, many of which are incomplete or not well annotated, exist for the Chelicerata (Childers, 2020) – the other major subphylum of arthropods (Regier *et al.*, 2010; Giribet and Edgecombe, 2019). Chelicerates include sea spiders, spiders, mites and scorpions among other organisms, as well as several extinct taxa (Dunlop and Selden, 1998; Ballesteros and Sharma, 2019). Chelicerates originated as marine animals about 500 million years ago (Dunlop and Selden, 1998; Dunlop, 2010). Molecular analyses suggest that one particular group, the omnivorous and detritivores acariform mites, may have been among the first arthropods that colonized terrestrial habitats and gave rise to ancient, simple terrestrial food webs (Walter and Proctor, 1999; Dunlop and Alberti, 2008; Schaefer *et al.*, 2010).

So far, the well-annotated genomic data of chelicerates is limited to animal parasites (including human pathogens and ticks), plant parasites, and predatory mites used in pest control (Cornman *et al.*, 2010; Grbić *et al.*, 2011; Rider *et al.*, 2015; Gulia-Nuss *et al.*, 2016; Hoy *et al.*, 2016; Dong *et al.*, 2017; Dong *et al.*, 2018). Other than some lower-quality genome assemblies (Bast *et al.*, 2016), there are no resources available for free-living soil and litter inhabiting species. Such data are, however, pivotal to understanding the evolution of parasitic lifestyles from a free-living condition and to bridge the gap between early aquatic chelicerates such as horseshoe crabs, and highly derived terrestrial pest species and parasites (Klimov and OConnor, 2013; Weinstein and Kuris, 2016; Shingate *et al.*, 2020). Because the phylogeny of Chelicerata remains unresolved, additional chelicerate genomes are urgently needed for comparative analyses (Dunlop, 2010; Ballesteros and Sharma, 2019; Lozano-Fernandez *et al.*, 2019). To help address this deficit, we report here the genome assembly of the soil dwelling oribatid mite *Archezogetes longisetosus* (Aoki, 1965; **Figure 1**) (Aoki, 1965) and a comprehensive analysis in the context of developmental genes, feeding biology, horizontal gene transfer and biochemical pathway evolution of chelicerates.

Archezogetes longisetosus (hereafter referred to as *Archezogetes*) is a member of the Oribatida (Acariformes, Sarcopiformes), an order of chelicerates well-known for their exceptional biosynthesis capacities, biochemical diversity, unusual mode of reproduction, unusually high pulling strength, mechanical resistances and pivotal ecological importance (Norton and Palmer, 1991; Maraun and Scheu, 2000; Heethoff and Koerner, 2007; Maraun *et al.*, 2007; Heethoff *et al.*, 2009; Raspotnig, 2009; Brückner *et al.*, 2017b; Schmelzle and Blüthgen, 2019; Brückner *et al.*, 2020). *Archezogetes*, like all members of its family Trhypochthoniidae (**Figure 1a**), reproduce *via* thelytoky (Heethoff *et al.*, 2013). That means the all-female lineages procreate *via* automictic parthenogenesis with an inverted meiosis of the holokinetic chromosomes, resulting in clonal offspring (Palmer and Norton, 1992; Wrensch *et al.*, 1994; Heethoff *et al.*, 2006; Bergmann *et al.*, 2018). While studying a parthenogenetic species is useful for the development of genetic tools as stable germ-line modifications can be obtained from the clonal progeny without laboratory crosses, one is confronted with the technical and philosophical problems of species delineation, cryptic diversity and uncertain species distribution (Heethoff *et al.*, 2013; Oxley *et al.*, 2014). Reviewing all available data, Norton (Norton, 1994; 2007) and Heethoff *et al.* (2013) concluded that *Archezogetes* is found widely on continents and islands throughout the tropical and partly subtropical regions of the world and that it is a middle-derived oribatid mite closely related to the suborder Astigmata.

One major feature of most oribatid mites is a pair of opisthonotal oil-glands and *Archezogetes* is no exception (Sakata and Norton, 2001; Raspotnig, 2009). These are a pair of large exocrine glands, each composed of a single-cell layer invagination of the cuticle, which is the simplest possible paradigm of an animal gland (Heethoff, 2012; Brückner and Parker, 2020). The biological role of these glands was rather speculative for a long time; ideas ranged from a lubricating and osmo- or thermoregulative function (Zachvatkin, 1941; Riha, 1951; Smrž, 1992) to roles in chemical communication (Shimano *et al.*, 2002; Raspotnig, 2006; Heethoff *et al.*, 2011a). So far about 150 different gland components have been identified from oribatid mites, including mono- and sesquiterpenes, aldehydes, esters, aromatics, short-chained hydrocarbons, hydrogen cyanide (HCN) and alkaloids (Saporito *et al.*, 2007; Raspotnig, 2009;

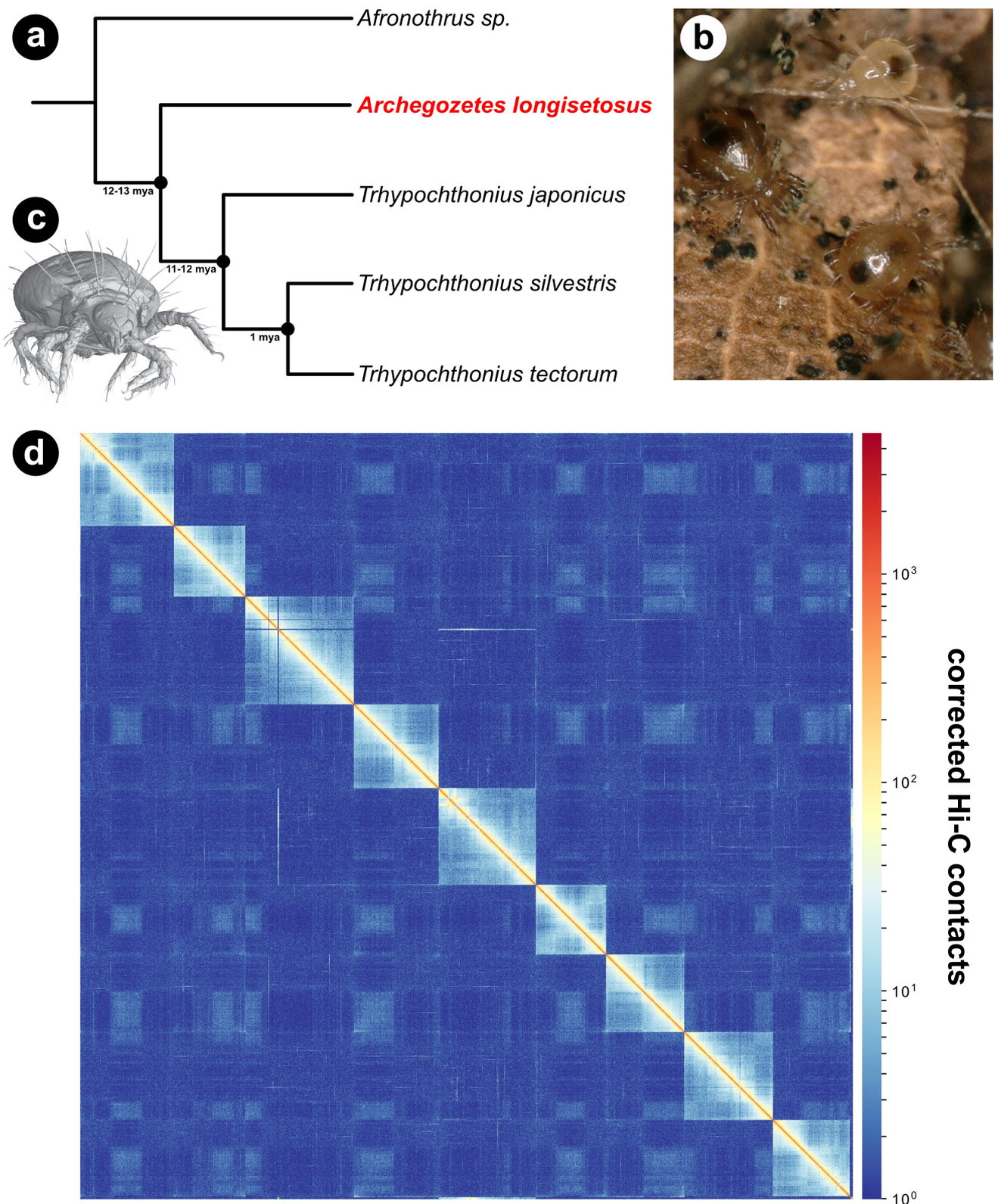


Figure 1 The mite, *Archegozetes longisetosus*, in its phylogenetic and natural environment. **a** – Species tree of selected oribatid mites of the family Trhypochthoniidae based on phylogenetic analyses and divergence time estimates (Heethöff *et al.*, 2011b). **b** – Two adults and one tritonymph of *Archegozetes* on a piece of leaf litter. The algae growing on the leaf serves as a food source for the mites. **c** – Habitus of an adult mite based on a surface rendering of a μ CT-scan reconstruction. Image courtesy of Sebastian Schmelzle. **d** – Hi-C interaction matrix maps of the nine *Archegozetes* chromosomes. The corrected contacts are indicated by the color scale on the right from red (high density) to blue (low density)

Brückner *et al.*, 2015; Brückner *et al.*, 2017b; Heethoff *et al.*, 2018). While some chemicals appear to be alarm pheromones (Shimano *et al.*, 2002; Raspotnig, 2006), most function as defensive allomones (Heethoff *et al.*, 2011a). Interestingly, alkaloids produced by oribatids mites are the ultimate source of most toxins sequestered by poison-frogs (Saporito *et al.*, 2007; Saporito *et al.*, 2009).

Terrestrial chelicerates predominately ingest fluid food. While phloem-feeding plant pests like spider mites and ecotoparasites like ticks adapted a sucking feeding mode, scorpions, spiders and others use external, pre-oral digestion before ingestion by morphologically diverse mouthparts (Cohen, 1995; Dunlop and Alberti, 2008; Bensoussan *et al.*, 2016; Gulia-Nuss *et al.*, 2016). Exceptions from this are Opiliones and sarcoptiform mites i.e., oribatid and astigmatid mites, all of which ingest solid food (Norton, 2007; Shultz, 2007; Heethoff and Norton, 2009). In general, oribatids feed on a wide range of different resources and show a low degree of dietary specialization (Brückner *et al.*, 2018b). The typical food spectrum of Oribatida, includes leaf-litter, algae, fungi, lichens, nematodes, and small dead arthropods such as collembolans (Riha, 1951; Schneider *et al.*, 2004a; Schneider *et al.*, 2004b; Schneider and Maraun, 2005; Heidemann *et al.*, 2011). In laboratory feeding trials, oribatid mites tend to prefer dark pigmented fungi, but also fatty acid-rich plant-based food (Schneider and Maraun, 2005; Brückner *et al.*, 2018b). Additionally, stable-isotope analyses of ¹⁵N and ¹³C suggested that Oribatida are primary- and secondary decomposers feeding on dead plant material and fungi, respectively (Schneider *et al.*, 2004a; Maraun *et al.*, 2011). The reasons for these preferences are still unknown, but they raise the question of how oribatid mites are able to enzymatically digest the cell walls of plants and fungi (Schneider *et al.*, 2004b; Smrž and Čatská, 2010; Brückner *et al.*, 2018a; Brückner *et al.*, 2018b).

Early studies on *Archezogozetes* and other mites found evidence for cellulase, chitinase and trehalase activity which was later attributed to symbiotic gut bacteria (Zinkler, 1971; Luxton, 1979; Haq, 1993; Siepel and de Ruiter-Dijkman, 1993; Smrž, 2000; Smrž and Norton, 2004; Smrž and Čatská, 2010). While such bacterial symbionts are a possible explanation, genomic data of other soil organisms and plant-feeding arthropods suggest a high frequency of horizontal transfer of bacterial and fungal genes enabling the digestion of cell walls (Grbić *et al.*, 2011; Mayer *et al.*, 2011; Wybouw *et al.*, 2016; Wu *et al.*, 2017; Wybouw *et al.*, 2018; McKenna *et al.*, 2019). For instance, an in-depth analysis of the spider mite *Tetranychus urticae* revealed a massive incorporation of microbial genes into the mite's genome (Grbić *et al.*, 2011; Wybouw *et al.*, 2018). Horizontal gene transfer appears to be a common mechanism for soil organisms, including mites, to acquire novel metabolic enzymes (Hoffmann *et al.*, 1998; Grbić *et al.*, 2011; Mayer *et al.*, 2011; Faddeeva-Vakhrusheva *et al.*, 2016; Wu *et al.*, 2017; Dong *et al.*, 2018), and hence seems very likely for *Archezogozetes* and other oribatid mite species that feed on plant or fungal matter.

Archezogozetes has been established as a laboratory model organism for three decades, having been used in studies, ranging from ecology, morphology, development and eco-toxicology to physiology and biochemistry (Barnett and Thomas, 2012; 2013b; 2013a; Heethoff *et al.*, 2013; Brückner *et al.*, 2017a; Barnett and Thomas, 2018; Brückner *et al.*, 2020). As such, *Archezogozetes* is among the few experimentally tractable soil organisms and by far the best-studied oribatid mite species (Thomas, 2002; Barnett and Thomas, 2012; Heethoff *et al.*, 2013). Since it meets the most desirable requirements for model organisms (Thomas, 2002), that is a rapid development under laboratory conditions, a dedicated laboratory strain was named *Archezogozetes longisetosus* **ran** in reference to its founder **Roy A. Norton** (Heethoff *et al.*, 2013, Figure 1b-c). Their large number of offspring enables mass cultures of hundreds of thousands of individuals, and their cuticular transparency during juvenile stages, and weak sclerotization as adults are general assets of an amenable model system (Heethoff and Raspotnig, 2012; Heethoff *et al.*, 2013; Brückner *et al.*, 2016; Brückner *et al.*, 2018c). In the past 10 years, *Archezogozetes* also received attention as a model system for chemical ecology (Raspotnig *et al.*, 2011; Heethoff and Raspotnig, 2012; Heethoff and Rall, 2015; Brückner *et al.*, 2016; Brückner and Heethoff, 2018; Thiel *et al.*, 2018; Brückner *et al.*, 2020). Some of these studies focusing

on the *Archezogozetes* gland revealed basic insights into the chemical ecology and biochemical capabilities of arthropods (Heethoff and Rall, 2015; Thiel *et al.*, 2018; Brückner *et al.*, 2020). Hence, *Archezogozetes* is poised to become a genetically tractable model to study the molecular basis of gland and metabolic biology.

The aim and focus of the current study were three-fold – to provide well-annotated, high-quality genomic and transcriptomic resources for *Archezogozetes longisetosus* (**Figure 1**), to reveal possible horizontal gene transfers that could further explain the feeding biology of oribatids, and to present *Archezogozetes* as a research model for biochemical pathway evolution. Through a combination of comparative genomic and detailed computational analyses, we were able to generate a comprehensive genome of *Archezogozetes* and provide it as an open resource for genomic, developmental and evolutionary research. We further identified candidate horizontal gene transfer events from bacteria and fungi that are mainly related to carbohydrate metabolism and cellulose digestion, features correlated with the mite feeding biology. We also used the genomic data together with stable-isotope labeling experiments and mass spectrometric investigation to delineate the biosynthesis pathway of monoterpenes in oribatid mites.

Results and discussion

Archezogozetes longisetosus genome assembly

Archezogozetes longisetosus (**Figure 1**) has a diploid chromosome number (2n) of 18 (Heethoff *et al.*, 2006), most likely comprising 9 autosomal pairs, the typical number of nearly all studied oribatid mite species (**Figure 1d**) (Norton *et al.*, 1993). There are no distinct sex chromosomes in *Archezogozetes*; this appears to be ancestral in the Acariformes and persisted in the Oribatida (Norton *et al.*, 1993; Wrensch *et al.*, 1994; Heethoff *et al.*, 2006). Even though some XX:XO and XX:XY genetic systems have been described in the closely related Astigmata, the sex determination mechanism in oribatids, including *Archezogozetes*, remains unknown (Oliver Jr, 1983; Norton *et al.*, 1993; Wrensch *et al.*, 1994; Heethoff *et al.*, 2006; Heethoff *et al.*, 2013).

To provide genetic resources, we sequenced and assembled the genome using both Illumina short-read and Nanopore MinION long-read sequencing approaches followed by scaffolding with Hi-C technology (**Figure 1d**; **Table 1**; see also “**Materials and Methods**”). Analyses of the *k*-mer frequency distribution of short reads (**Table 1**; **Supplementary Figure S1**) resulted in an estimated genome size range of 135-180 Mb, encompassing the final HiC assembled size of 143 Mb (**Table 1**; see also “**Materials and Methods**”). Compared to genome assemblies of other acariform mites, the assembled genome size of *Archezogozetes* is on the large end of the spectrum (with *Opiella nova* and *O. subpectinata* representing notable exceptions) (Brandt *et al.*, 2021), but is smaller than that of most mesostigmatid mites, ticks and spiders that on average range from ~250 Mb up to 2.5 Gb (Grbić *et al.*, 2011; Bast *et al.*, 2016; Gulia-Nuss *et al.*, 2016; Hoy *et al.*, 2016; Dong *et al.*, 2017; Schwager *et al.*, 2017; Dong *et al.*, 2018). In the context of arthropods in general, *Archezogozetes*' genome (**Table 1**) is among the smaller ones and shares this feature with other arthropod model species like *Tetranychus urticae* (hereafter called ‘spider mite’), *Drosophila*, clonal raider ant and red flour beetle (Consortium, 2008; Grbić *et al.*, 2011; Oxley *et al.*, 2014; dos Santos *et al.*, 2015). Even though we surface-washed the mites and only used specimens with empty alimentary tracts for sequencing, we removed a total 532 sequences (438 long-read contigs and 94 Hi-C scaffolds) with high bacterial or fugal homology making up approximately 9 Mb of contamination in the final assembly (see **supplementary Table S1**). The final filtered Hi-C genome assembly was composed of 164 scaffolds, the majority of which is composed of nine pseudochromosomes with an N₅₀ contiguity of 16.25 Mb (**Table 1** and **Figure 1d**).

Genome scaffolding and the analysis of genome structure in arthropods by all-vs-all chromosome conformation capture (Hi-C) is a relatively new field that has grown in recent years (Richards, 2019). Like other arthropods, the *Archezogozetes* genome is organized into chromosome territories, compartments, and sub-compartment structures (**Figure 1d**). Classic

microscopy and fluorescent staining previously revealed that *Archezogozetes* has a diploid set of 18 (2n) highly condensed chromosomes, and we recovered nine pseudo-chromosomes in our haploid assembly (Heethoff *et al.*, 2006). Even though Hi-C is often not able to resolve inter-chromosomal interactions and long range-contacts (Quinodoz *et al.*, 2022), we discovered multiple such contacts between different chromosomes, for instance chromosome 1 is interacting with parts of each chromosome except chromosome 5 (**Figure 1d**). One hypothesis which could explain this pattern is that the *Archezogozetes* genome is highly compact, and chromatin is densely packed in the nucleus (see also the genome of the tomato russet mite; Greenhalgh *et al.*, 2020).

The official gene set and annotation of *Archezogozetes*

We generated the official gene set (OGS) for *Archezogozetes* by an automated, multi-stage process combining *ab initio* and evidenced-based (RNAseq reads, transcriptomic data and curated protein sequences) gene prediction approaches (see “**Materials and Methods**”) yielding 24,538 gene models. In comparison to other mites and ticks as well as insects, this is well within the range of the numbers discovered in other Chelicerata so far (**Figure 2a**). Chelicerates with a large OGS, however, usually possess larger genomes (1-7 Gb), which suggests that *Archezogozetes* may have a relatively dense distribution of protein-coding genes in its genome. On the other hand, ticks can have giga-base sized genomes, but only a rather small number of gene models, probably due to high repetitive content (Palmer *et al.*, 1994; Van Zee *et al.*, 2007; Gulia-Nuss *et al.*, 2016; Barrero *et al.*, 2017). Lacking more high-quality genomic resources of mites, it is thus not clear whether the OGS of *Archezogozetes* is the rule, or rather the exception within the Oribatida.

To compare if *Archezogozetes*’ OGS is similar to predicted genes of other oribatid mites as well as Prostigmata and Astigmata, we first clustered genes by ortholog inference (OrthoFinder; Emms and Kelly (2015)), removed species-specific genes and constructed a presence-absence matrix of orthogroups to ordinate the data using non-metric multidimension scaling (NMDS, **Figure 2b**). Ordination revealed that the OGS of *Archezogozetes* is well nested with other oribatid mites and clearly separated from their closest relative the astigmatid mites as well as prostigmatid mites (**Figure 2b**). As a first step in annotating the OGS, we ran KOALA (KEGG Orthology And Links Annotation) to functionally characterize the genes (Kanehisa *et al.*, 2016). In total, 9,719 (39.6%) of all genes received annotation and about two thirds of all genes were assigned either as genes related to genetic information processing (34%) or metabolic genes (30%), while the remaining genes fell into different KEGG categories (**Figure 2c**). To further annotate the genome, we followed the general workflow of funannotate with some modifications (Palmer and Stajich, 2017, see “**Materials and Methods**”).

Overall, we found 16,685 genes (68%) of the OGS with homology to previously published

Table 1 *Archezogozetes longisetosus* genome metrics.

Feature	Value
Estimated genome size	135-180 Mb
Assembly size	143 Mb
Coverage based on assembly (short/long)	200x (short), 60x (long)
# scaffolds	164
N ₅₀ (scaffolds)	16.25 Mb
Mean scaffold length	8.73 Mb
GC content	30.90%
# gene models	24,538

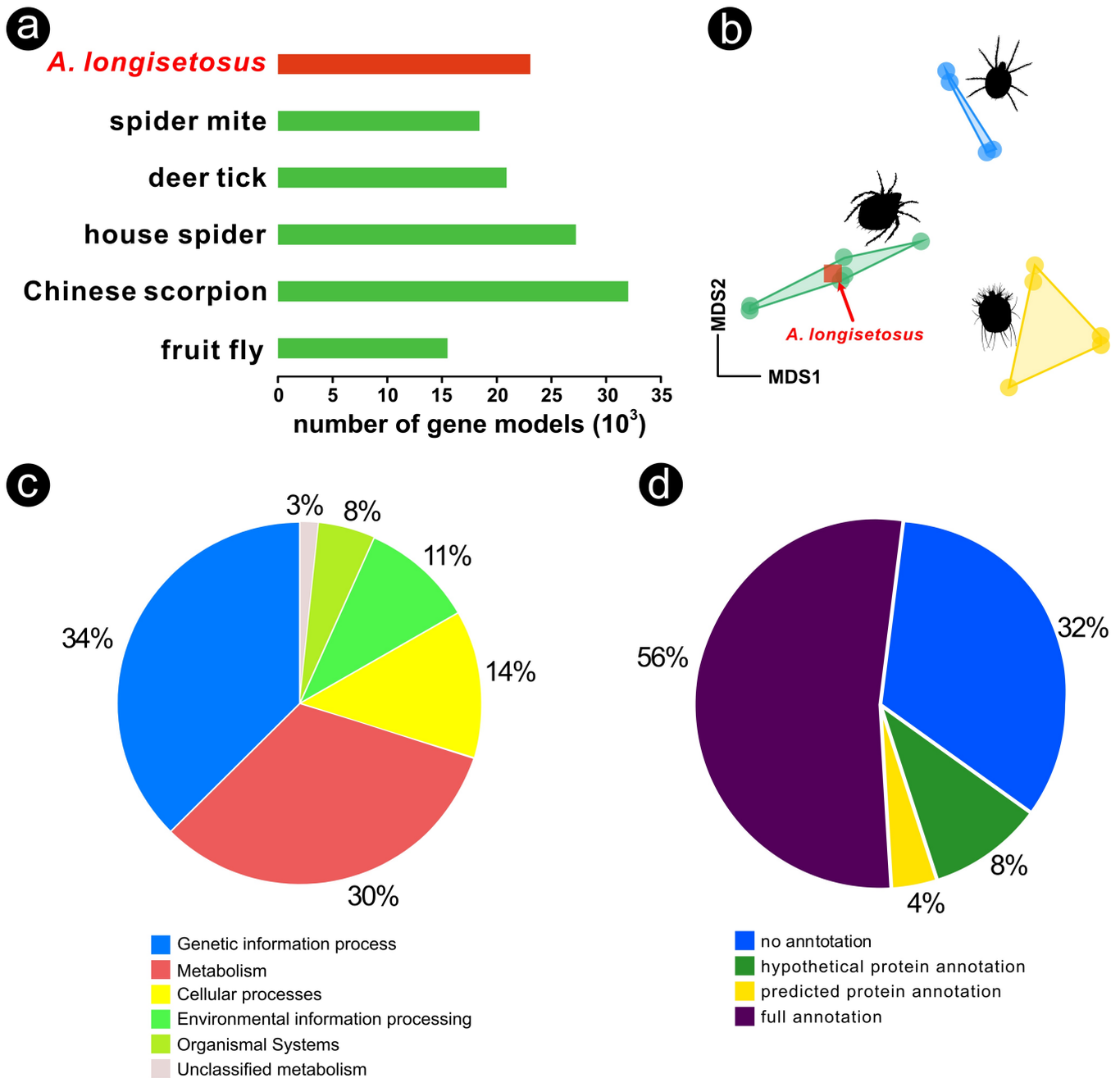


Figure 2 Comparisons and annotations of the official gene set (OGS) of *Archegozetes longisetosus*. **a** – Number of gene models of the mites compared to other mites, chelicerates and the fruit fly (Grbić *et al.*, 2011; Cao *et al.*, 2013; dos Santos *et al.*, 2015; Gulia-Nuss *et al.*, 2016; Schwager *et al.*, 2017). **b** – Non-linear multidimensional scaling plot (NMDS) of clustered orthogroups based on the OGS or predicted proteins of several mite species. *Archegozetes longisetosus* is marked a square, nested within Oribatida. Prostigmata are depicted in blue, Astigmata in yellow and Oribatida in green. **c** – Pie chart showing the percentage composition of genes of the *Archegozetes* annotated to different broad biological categories by GhostKOALA. **d** – Pie chart describing the overall annotation of the OGS of the mite.

sequences (**Figure 2d**). For over half of all genes (56%), we were able to assign a full annotation, 4% of all genes only showed homology to bioinformatically predicted proteins of other species, while 8% of all genes only showed homology to hypothetical proteins (**Figure 2d**). As only a few high-quality, annotated mite genomes are available and the two-spotted spider mite is the sole species with any experimentally confirmed gene models, it is not surprising that we were only able to confidently annotate about 60% of all genes of the OGS (**Figure 2d**).

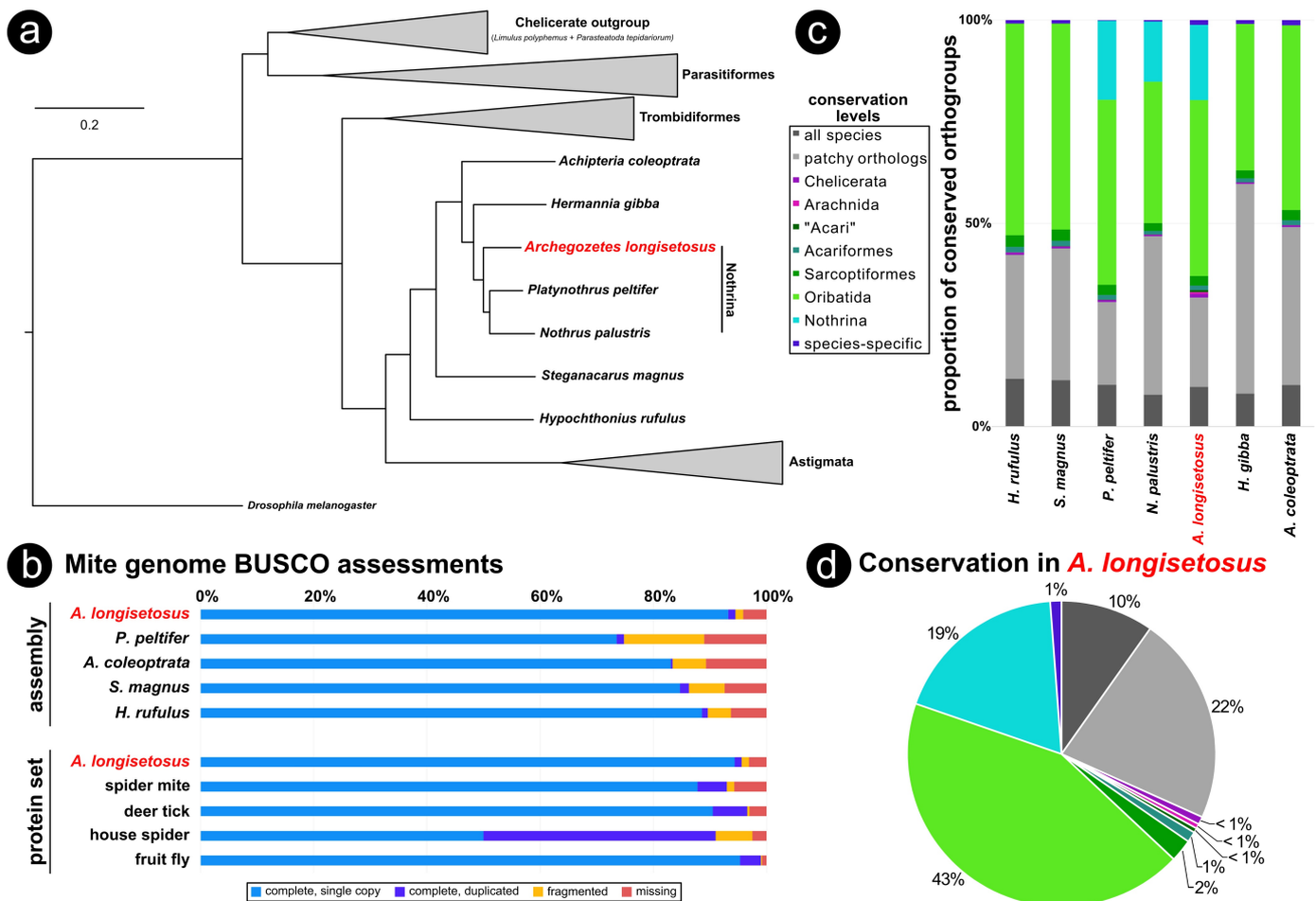


Figure 3 Orthology comparison and phylogenetic placement of *Archegozetes longisetosus* among other chelicerates. **a** – Maximum likelihood phylogeny based on concatenation of 1,121 orthologs showing the mites phylogenetic position within the Oribatida (all nodes have 100% support; branch length unit is substitutions per site). For the fully expanded tree see supplementary Figure S2. **b** – BUSCO-assessment of the *Archegozetes* genome assembly and protein set for both ortholog presence and copy number compared to other oribatid mites and selected model species, respectively. **c** – Comparisons of protein-coding genes in seven oribatid mite species (for a full comparison of all species see supplementary Figure S3) with *Archegozetes* highlighted in red. The bar charts show the proportion of orthogroup conservation with each species (see insert legend) based on OrthoFinder clustering. **d** – Detailed pie chart depicting the conservation levels of orthogroup in *Archegozetes*.

Orthology and comparative genomics of chelicerates

To further access the protein-coding genes of the mite, we compared the OGS to other chelicerates. Both concatenated maximum likelihood and coalescent species-tree phylogenomic approaches based on 1,121 orthologs placed *Archegozetes*, as expected, within the Nothrina (Pachl *et al.*, 2012; Heethoff *et al.*, 2013) with strong support and recovered previously found oribatid clade topologies (**Figure 3a**). Our analysis placed the Astigmata as a sister group of Oribatida and not nested within oribatids as suggested based on life-history, chemical defensive secretions, morphology and several molecular studies (Norton, 1994; 1998; Sakata and Norton, 2001; Alberti and Michalik, 2004; Maraun *et al.*, 2004; Liana and Witaliński, 2005; Domes *et al.*, 2007; Dabert *et al.*, 2010; Koller *et al.*, 2012; Pepato and Klimov, 2015; Klimov *et al.*, 2018; Li and Xue, 2019). The relationship of Oribatida and Astigmata has been challenging to resolve for the past decades and several studies using different set of genes, ultra-conserved elements or transcriptomic data reconstructed discordant phylogenies, some of which are similar to ours (Maraun *et al.*, 2004; Domes *et al.*, 2007; Dabert *et al.*, 2010; Pepato and Klimov, 2015; Klimov *et al.*, 2018; Li and Xue, 2019; Lozano-Fernandez *et al.*,

2019; Van Dam *et al.*, 2019). Overall, the oribatid-astigmatid relationship remains unresolved and a broader taxon sampling, especially of more basal Astigmata, will be necessary (Norton, 1994; 1998; Domes *et al.*, 2007; Klimov *et al.*, 2018; Lozano-Fernandez *et al.*, 2019; Van Dam *et al.*, 2019). We recovered Trombidiformes (Prostigmata and Sphaerolichida) as sister group of the Sarcoptiformes (Oribatida and Astigmata) constituting the Acariformes (**Figure 3a**). Neither the maximum likelihood phylogeny (**Figure 3a**), nor the coalescence-based phylogeny (**Supplementary Figure S2**) reconstructed the Acari (i.e., Acariformes and Parasitiformes) as a monophyletic clade. Even though there is morphological, ultrastructural and molecular evidence for a biphyletic Acari, as we recovered here, this relationship and larger-scale chelicerate relationships remain unclear (Alberti, 1984; 1991; Dabert, 2006; Dunlop and Alberti, 2008; Jeyaprakash and Hoy, 2009; Li and Xue, 2019; Lozano-Fernandez *et al.*, 2019; Van Dam *et al.*, 2019).

To further assess the quality and homology of both the genome assembly (**Table 1**) and the OGS (**Figure 2**), we used the 1066 arthropod Benchmarking Universal Single-Copy Ortholog (BUSCO) genes data set (Simão *et al.*, 2015). Nearly all BUSCO genes were present in the *Archezogozetes* assembly and OGS (94.6% and 95.4%, respectively; **Figure 3b**). Compared to other genomes sequenced so far, the *Archezogozetes* genome has the highest completeness among oribatid mites and the OGS completeness is on par to the well curated genomes of other chelicerate species and *Drosophila melanogaster* (**Figure 3b**). This result is not surprising because the *Archezogozetes* genome was assembled from long-read and short-read data followed by scaffolding with Hi-C technology, while most other oribatid mite genomes were solely short reads sequenced on older Illumina platforms (Bast *et al.*, 2016). The fraction of duplicated BUSCO genes in *Archezogozetes* (1.2%) was smaller than the spider mite and deer tick (Grbić *et al.*, 2011; Gulia-Nuss *et al.*, 2016), but very low compared to the house spider (**Figure 3c**), whose genome underwent an ancient whole-genome duplication (Schwager *et al.*, 2017).

Overall, the high quality of both the genome assembly and OGS of *Archezogozetes* is unprecedented as compared to genomic resources of other oribatid mites. We next categorized all protein models from the OGS by conservation level based on a global clustering orthology analysis (OrthoFinder; Emms and Kelly, 2015) of 23 species (**Figure 3c**; **supplementary Figure S3**) representing Acariformes, Parasitiformes, several other chelicerates and the fly *Drosophila*. As for most other species (Siepel *et al.*, 2005; Thomas *et al.*, 2020), about a third of all orthogroups was highly conserved (**Figure 3c**) across the arthropods, being either in all species (10%; **Figure 3d**) or in most (22%; **Figure 3d**). Only 1% of all *Archezogozetes* orthogroups did not show homology and were species specific (**Figure 3c and d**). Only a low proportion (**Figure 3c**) of orthogroups was conserved across the higher taxonomic levels (all <1% in *Archezogozetes*; **Figure 3d**), which is in line with previous studies that included prostigmatid and mesostigmatid mites (Hoy *et al.*, 2016; Dong *et al.*, 2017; Dong *et al.*, 2018). Interestingly, there was a large proportion of orthogroups conserved across all Oribatida (43% in *Archezogozetes*; **Figure 3d**) and also about 19% of orthogroups in *Archezogozetes* were shared only with other Nothrina (**Figure 3d**). A fairly large percentage of these orthogroups may contain potentially novel genes that await experimental verification and functional analyses (Emms and Kelly, 2015; Nagy *et al.*, 2020; Thomas *et al.*, 2020). Especially the lack of homology within the Sarcoptiformes (2-3%; **Figure 3c**) may explain the controversial placement of Astigmata as a sistergroup of Oribatida that we recovered (**Figure 3a**). This grouping is likely caused by a long-branch attraction artifact and the sister relationship was incorrectly inferred (Dabert, 2006; Domes *et al.*, 2007; Dabert *et al.*, 2010; Pepato and Klimov, 2015; Klimov *et al.*, 2018), because orthogroup clustering could not detect enough homology between oribatids and the Astigmata so far sequenced, which are highly derived. Hence, a broad taxon sampling of basal astigmatid mite genomes seems necessary to resolve Oribatida-Astigmata relationship (Norton, 1994; 1998; Pepato and Klimov, 2015; Li and Xue, 2019; Van Dam *et al.*, 2019).

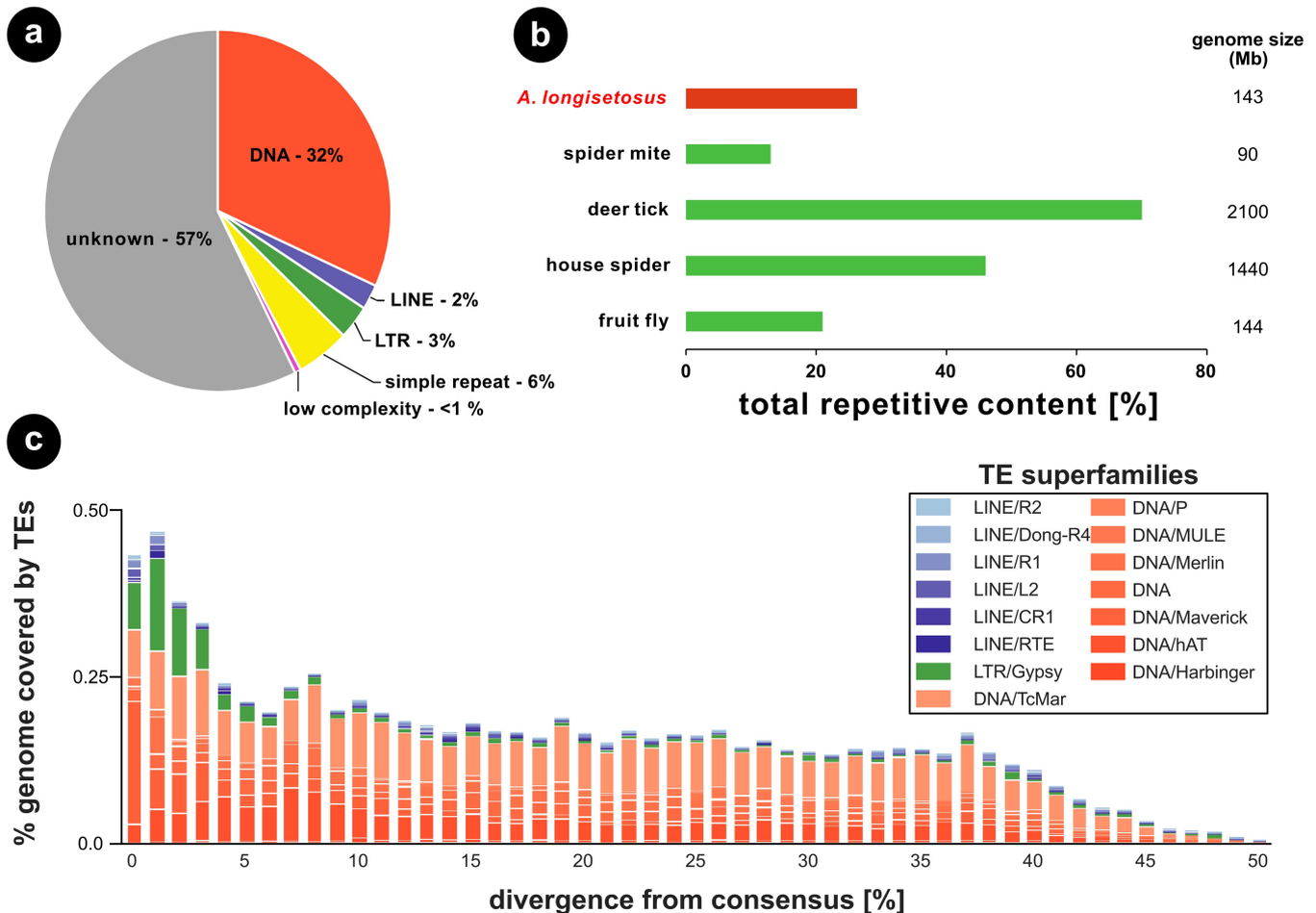


Figure 4 Comparison of repeat content estimations and transposable element (TE) landscape of *Archegozetes longisetosus*. **a** – Repetitive element categories of *Archegozetes* based on the results from RepeatModeler and MITETracker. LINE= long interspersed nuclear element, LTR= long terminal repeat. **b** – Comparison of total repetitive content among *Archegozetes*, other model chelicerates and the fly. All values are from the respective genome paper of the species, except for the fly. **c** – Repeat divergence plot showing TE activity through time for the major TE superfamilies of *Archegozetes*. Transposable elements with a low divergence from the consensus were recently active, while TEs diverging from the consensus depicted older activities (x-axis).

Repeat content analysis and transposable elements (TEs)

For clonal species like *Archegozetes*, reproducing in the absence of recombination, it has been hypothesized that a reduced efficacy of selection could result in an accumulation of deleterious mutations and repeats in the genome (Muller, 1964; Arkhipova and Meselson, 2000; Nuzhdin and Petrov, 2003; Schön *et al.*, 2009; Barton, 2010; Charlesworth, 2012). There is, however, no evidence for such an accumulation in oribatids or other arthropods (Bast *et al.*, 2016). The genome of *Archegozetes* repeat content was predicted to be 26% of the genome (Figure 4). Generally, we found that most of the repetitive content in *Archegozetes* could not be classified (57%; Figure 4a). The high proportion of unknown repeats likely corresponds to novel predicted repetitive content, because of limited repeat annotation of mites in common repeat databases such as RepBase (Bast *et al.*, 2016). Regarding the two major classes of repeat content, DNA transposons made up about 32% of total repeats, while only 5% represented retrotransposons (Figure 4a). About 6% of total repetitive content comprised simple and low complexity repeats (Figure 4a). Overall, the total repetitive content (26%, Figure 4b) seems to be within a normal range for chelicerates and arthropods.

The repeat content found in other oribatid mites was lower (Bast *et al.*, 2016), but recent

studies suggest that sequencing technology, read depth and assembly quality are paramount to the capacity of identifying repeat content and TEs (Bourque *et al.*, 2018; Panfilio *et al.*, 2019). Hence, it is very likely the current genomic data for other Oribatida underestimate the actual total repetitive content. More low-coverage, long-read sequencing could reduce the assembly fragmentation and likely reveal a higher proportion of repeats, closer to the actual repetitiveness of oribatid genomes (Panfilio *et al.*, 2019).

Different classes of transposable elements (TEs) are characterized by the mechanism they use to spread within genomes and are known to influence population dynamics differently (Finnegan, 1989; Bourque *et al.*, 2018; Crescente *et al.*, 2018). We therefore analyzed the evolutionary history of TE activity in *Archezogozetes* in more detail (**Figure 4c**). The main TE superfamilies were DNA transposons (**Figure 4a and c**), which seems to be a common pattern of oribatid mite genomes. For *Archezogozetes*, they appear to have accumulated in the genome for a long time (i.e. they are more divergent from the consensus; (Waterston *et al.*, 2002)) with Tc1/mariner – a superfamily of interspersed repeats DNA transposons (Bourque *et al.*, 2018) – being the most abundant one (**Figure 4c**). Interestingly, we found an increase in TE activity with 0-4% sequence divergence range, indicating a recent burst (**Figure 4c**). This burst contained an enrichment of DNA Mavericks, which are the largest and most complex DNA transposons with homology to viral proteins (Bourque *et al.*, 2018), but also several of retrotransposons. Among these, is the Long Terminal Repeat (LTR) gypsy retroelement (**Figure 4c**), which is closely related to retroviruses (Bourque *et al.*, 2018). Like retroviruses, it encodes genes equivalent to *gag*, *pol* and *env*, but relatively little is known about how it inserts its DNA into the host genome (Dej *et al.*, 1998; Havecker *et al.*, 2004). So far, it is unknown what these TEs do in *Archezogozetes*, but the recent burst in TE abundance might suggest that some changes in the genome might have happened since the became a laboratory model nearly 30 years ago (Heethoff *et al.*, 2013).

The *Archezogozetes* Hox cluster

The Hox genes are a group of highly conserved transcription factor-encoding genes that are used to pattern the antero-posterior axis in bilaterian metazoans (Holland and Hogan, 1988; Hrycaj and Wellik, 2016). Ancestrally, arthropods likely had ten Hox genes arranged in a cluster (Hughes and Kaufman, 2002). During arthropod development, the Hox genes specify the identities of the body segments, and mutations in Hox genes usually result in the transformation of segmental identities (Hughes and Kaufman, 2002). The importance of Hox genes in development of metazoans makes knowledge of their duplication and disappearances important for understanding their role in the evolution of body plans (Hughes and Kaufman, 2002).

Mites largely lack overt, external signs of segmentation, other than the serially arranged appendages of the prosoma (Dunlop and Lamsdell, 2017). Signs of segmentation in the posterior body tagma, the opisthosoma, do exist in adult members of Endeostigmata (van der Hammen, 1970). However, these segmental boundaries are largely present only in the dorsal opisthosoma, making it difficult to assess how these correspond to the ventral somites (van der Hammen, 1970; Dunlop and Lamsdell, 2017). Developmental genetic studies of the spider mite and *Archezogozetes* suggest that acariform mites only pattern two segments in the posterior body region, during embryogenesis (Grbić *et al.*, 2011; Barnett and Thomas, 2012; 2013b; 2018). This stands in stark contrast to other studied chelicerate embryos. For example, during embryogenesis the spider *Parasteatoda tepidariorum* patterns twelve opisthosomal segments (Schwager *et al.*, 2015) and the opilionid *Phalangium opilio* patterns seven (Sharma *et al.*, 2012). Furthermore, a member of Parasitiformes, the tick *Rhipicephalus microplus*, appears to pattern eight opisthosomal segments during embryogenesis (Santos *et al.*, 2013).

Parallel to the observation of segmental reduction in the spider mite, genomic evidence suggests that this acariform mite has lost two of its Hox genes, *i.e.*, *Hox3* and *abdominal-A* (*abd-A*) (Grbić *et al.*, 2011). Interestingly, orthologs of *abd-A* in other studied arthropods pattern the

posterior segments as well. A genomic comparison of arthropod Hox clusters has also shown a correlation between independent losses of *abd-A* and a reduction in posterior segmentation (Pace *et al.*, 2016). To investigate whether the loss of segmentation in *Archezogozetes* is also due to an absence in *abd-A*, we annotated its Hox cluster, paying close attention to the region between the Hox genes *Ultrabithorax* (*Ubx*) and *Abdominal-B* (*Abd-B*), which is usually where this gene resides in other arthropods (Hughes and Kaufman, 2002). Our results suggest that the *Archezogozetes* Hox genes are clustered in a contiguous sequence (HiC scaffold 3, total size ~12.36 Mbp) in the same order as suggested for the ancestral arthropod (Heethoff and Rall, 2015). Furthermore, we found no sequences suggestive of an *abd-A* ortholog in *Archezogozetes* (**Figure 5a**). These data also support the findings of a previous PCR survey that retrieved no *abd-A* ortholog in *Archezogozetes* (Cook *et al.*, 2001). Genomic evidence from the Parasitiformes *Ixodes scapularis* and *Metaseiulus occidentalis* reveal that these taxa maintain orthologs of all ten Hox genes, however in *M. occidentalis* these genes are not clustered as they are in *I. scapularis* (Gulia-Nuss *et al.*, 2016; Hoy *et al.*, 2016).

Taken together, these observations suggest that the last common ancestor of acariform mites likely lost its *abdominal-A* gene as well as experienced a reduction in opisthosomal segmentation (**Figure 5b**). Alternatively, these shared losses of *abd-A* may be due to convergence due to similar selective pressures favoring a reduction in body size. The dorsal, external segmentation of endeostigmatid mites does not necessarily contradict the hypothesis of a loss of *abd-A* at the base of the acariform mites. As Hox genes are usually deployed after the genetic establishment of segments in arthropods (Hughes and Kaufman, 2002), the opisthosomal segments in endeostigmatid mites may still develop in the absence of *abd-A*. However, this hypothesis needs further testing with observations of segmental gene expression in endeostigmatids as well as additional acariform species.

Life-stage specific RNA expression patterns

Developmental and gene expression data from *Archezogozetes* embryos (**Figure 5 d and e**) have elucidated many of the potential mechanisms driving the morphogenesis of many developmental peculiarities. These peculiarities include the suppression of the fourth pair of walking legs during embryogenesis as well as the reduction of opisthosomal segmentation (Telford and Thomas, 1998; Thomas, 2002; Barnett and Thomas, 2012; 2013a; 2013b; 2018). In typical acariform mites, embryogenesis ends with the first instar, the prelarva, which usually remains within the egg chorion, as in *Archezogozetes*. Hatching releases the second instar, the larva, which is followed by three nymphal instars (proto-, deutero- and tritonymph) and the adult, for a total of six instars. (Heethoff *et al.*, 2007). Thus far, methodological limitations have made it difficult to examine how mite segmentation and limb development progress throughout these instars.

To this end, we used RNAseq to calculate the transcripts per million (tpm) values of genes known to be, or suspected to be, involved in limb development and segmentation throughout the six different instars of *Archezogozetes*. Prior to comparing these tpm values, gene orthology was confirmed *via* phylogenetic analyses (**supplementary Figures S4-S11**; see **Table S2** for phylogenetic statistics and **Table S3** for tpm values). Regarding the total number of genes expressed across the different life stages, we found that earlier instars generally expressed a higher number of genes (**Figure 5c**). While most expressed genes were shared across all instars, more transcripts were shared between the eggs and the larvae and among all five juvenile instars. Additionally, we found that earlier instars expressed a larger number of stage-specific genes as compared to later instars and adults (**Figure 5c**).

Gene expression, scanning electron microscopy (SEM) and time-lapse data have revealed that the development of the fourth pair of walking legs in *Archezogozetes* is suppressed until after the larval instar (Telford and Thomas, 1998; Barnett and Thomas, 2012; 2018). The resulting larva is thus hexapodal (see also embryo in **Figure 5e**), which constitutes a putative synapomorphy of Acari, if they are monophyletic (Dunlop and Albetri, 2008). In arthropods,

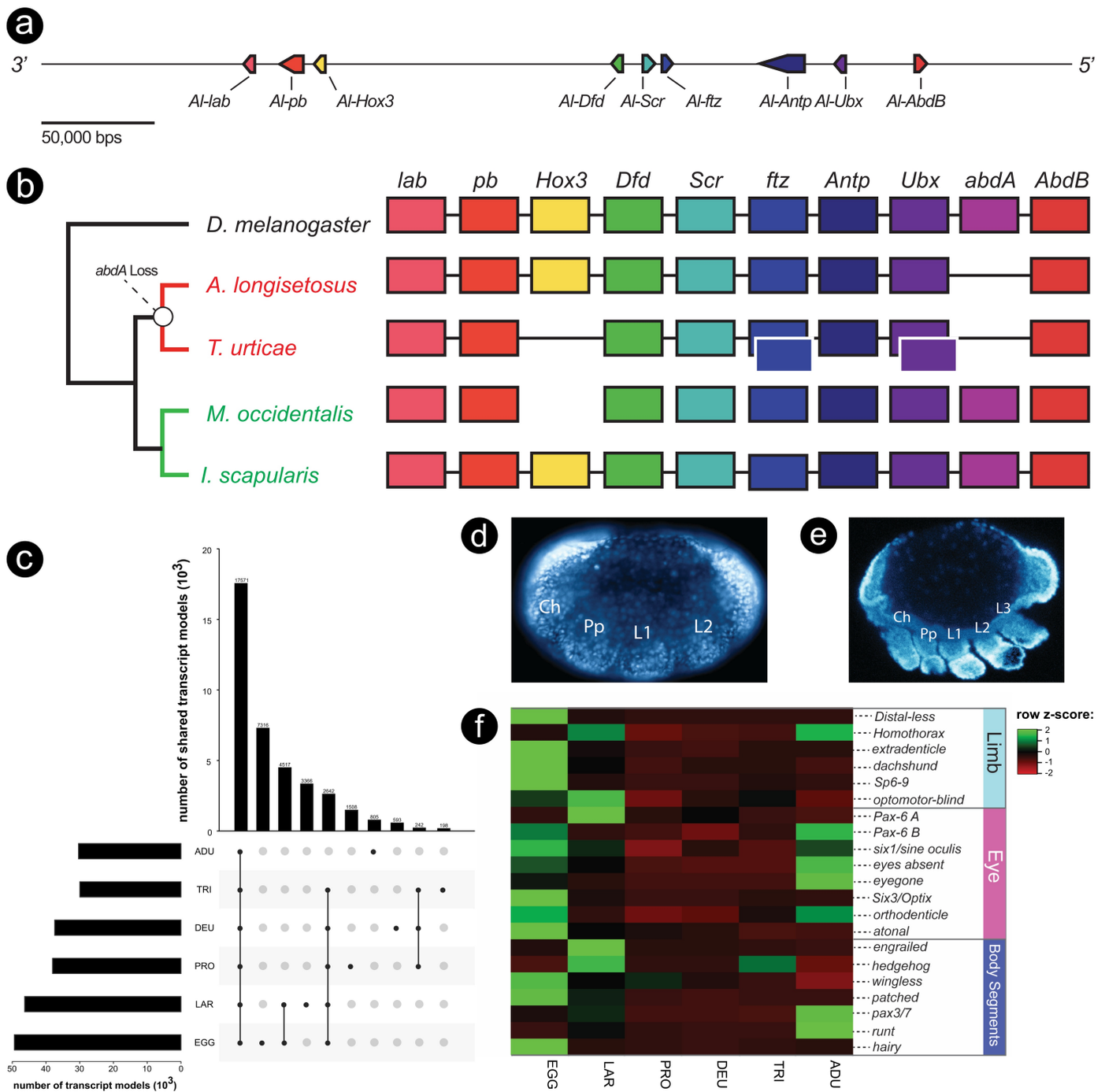


Figure 5 The genomic organization of the Hox genes and life-stage specific expression patterns of developmental genes in *Archegozetes longisetosus*. **a** – Schematic of the genomic region enclosing the *Archegozetes* Hox cluster. The genomic organization of the Hox cluster is collinear, as it is in many arthropod taxa, however an abdominal-A ortholog is absent. Arrowed boxes denote the direction of transcription. The scale bar represents 50,000 base pairs. **b** – A comparison of the Hox cluster organization of reported members of Acari with the fruit fly *Drosophila melanogaster* as the outgroup. The last common ancestor of the parasitiform mites *M. occidentalis* and *I. scapularis* likely had an intact Hox cluster (green branches and labels), whereas abdominal-A was likely lost in the last common ancestor of acariform mites, as represented by *Archegozetes* and *T. urticae* (red branches and labels). Boxes with white borders represent duplicated Hox genes. Lines through the boxes indicate an intact Hox cluster. See text for further details. **c** – Number of transcripts shared across the different life stages of *Archegozetes*. The barplot panel on the left shows the numbers of transcripts in each stage. Exemplars of **(d)** early and **(e)** mid- germ-band embryos. Ch= chelicera; L1-3= walking legs 1-3; Pp= pedipalp. Embryos are stained with the nuclear dye DAPI and oriented with the anterior to the left of the page. **f** – Non-clustered heatmap showing the relative expression (row z-score based on tpm) patterns of putative limb, eye, and body segmentation genes throughout the embryonic, larval instars, and adult stages of *Archegozetes*. See **supplementary Table S3** for average tpm values. Life stages (for **c** and **f**): EGG= egg; LAR= larva; PRO= protonymph; DEU= deutonymph; TRI= tritonymph; ADU= adult.

the development of the limbs is generally accomplished *via* the activity of highly conserved regulatory genes, termed the “limb gap genes.” These genes are expressed along their proximo-distal axes to establish the specific identities of the limb podomeres. The limb gap genes include *extradenticle* (*exd*) and *homothorax* (*hth*), which act together to specify the proximal limb podomeres, *dachshund* (*dac*), which specifies the medial podomeres, and *Distal-less* (*Dll*) which specifies the distal-most podomeres. It was previously shown that the deployment of these genes in the anterior appendages of *Archegozetes*, *i.e.*, the chelicerae, pedipalps and first three pairs of walking legs (**Figure 5d** and **e**), is similar to that of other chelicerate taxa (Barnett and Thomas, 2013a; Schwager *et al.*, 2015; Sharma *et al.*, 2015). However, in the anlagen of the fourth pair of walking legs, only the proximal-specifying genes, *exd* and *hth*, are expressed (Barnett and Thomas, 2013a).

Whether the limb gap genes are re-deployed during the transition from the prelarval to larval instars in order to activate the development of the fourth pair of walking legs remains an open question. We therefore compared the average tpm values of verified limb gap genes (*i.e.*, *Al-Dll*, *Al-Hth*, *Al-exd*, and *Al-dac* (Barnett and Thomas, 2013a)) in embryos and at each instar stage (**Figure 5f**). We also compared the tpm values of the *Archegozetes* orthologs of *Sp6-9* and *optomotor blind*, genes shown to be involved in limb formation in spiders (Königsmann *et al.*, 2017; Heingård *et al.*, 2019). We hypothesized that limb development genes would show high expression in the larval stage leading to the development of the octopodal protonymph. We did observe an increase in the tpm averages of *Al-hth* as well as *Al-optomotor-blind*, however the aforementioned limb gap gene expression levels were similar between these instars (**Figure 5f**). Taken together, these genes may not be up-regulated for the formation of the fourth pair of walking legs between these two instars.

Chelicerate embryos segment their bodies through a “short/intermediate germ” mechanism, whereby the anterior (prosomal) segments are specified asynchronously (Schwager *et al.*, 2015). This usually occurs well before the sequential addition of posterior segments from a posterior growth zone. Based on neontological and paleontological data, chelicerate arthropods may have ancestrally had an opisthosoma comprised of 12 or more segments (Dunlop and Selden, 1998; Dunlop, 2010; Dunlop and Lamsdell, 2017). Embryonic expression data for the segment polarity genes, those genes that delineate the boundaries of the final body segments, have shown that in most studied chelicerate embryos opisthosomal segments are delineated during embryogenesis (Schwager *et al.*, 2015; Dunlop and Lamsdell, 2017). However, as discussed above, expression data in *Archegozetes* embryos suggest that only two opisthosomal segments are patterned during embryogenesis (Barnett and Thomas, 2012; 2018); this indicates that mites have significantly reduced their number of opisthosomal segments either by loss or by fusion. Further complicating this is the observation that many mites add segments as they progress through the larval instars, a phenomenon known as anamorphic growth (Dunlop and Lamsdell, 2017).

To determine by what genetic process *Archegozetes* may add segments during post-embryonic ontogeny, we assessed the expression of known chelicerate and arthropod segmentation genes in each instar transcriptome (**Figure 5f**) (Schwager *et al.*, 2015). We observed an up-regulation of the segmentation genes *hedgehog* and *engrailed* in the larvae, as well as the slight up-regulation of *patched* and *pax3/7*. Furthermore, the segmentation gene *wingless* was slightly up-regulated in the protonymph, as well as a slight up-regulation of *hedgehog* in the tritonymph. Lastly, we found that transcripts of the genes *pax3/7* and *runt* were up-regulated in adults. These results suggest that *Archegozetes* does pattern body segments during the progression through the its instars similar to other Chelicerata (Dunlop and Lamsdell, 2017).

Another peculiarity of *Archegozetes* is that these mites lack eyes (see more details below). Eye loss has been documented in other arachnid clades, including independently in other members of Acari (Evans, 1992; Walter and Proctor, 1999), and it has been recently demonstrated that a species of whip spider has reduced its eyes by reducing the expression of retinal determination genes that are shared throughout arthropods (Gainett *et al.*, 2020). We sought to determine if eye loss in *Archegozetes* also is associated with the reduced expression of these

genes (see also analysis of photoreceptor genes below). The genes, which have been shown to be expressed in the developing eyes of spiders and whip scorpions, include *Pax-6*, *six1/sine oculis (so)*, *eyes absent (eya)*, *Eyegone*, *Six3/Optix*, and *atonal* (Samadi *et al.*, 2015; Schomburg *et al.*, 2015; Gainett *et al.*, 2020). We also followed the expression of *Al-orthodenticle*, a gene previously shown to be expressed in the ocular segment of *Archezogozetes* (Telford and Thomas, 1998). Surprisingly, all of these genes, excluding the *Pax-6* isoform A and *eyegone*, are indeed expressed during embryogenesis (**Figure 5f**). Aside from the larval expression of the *Pax-6* isoform A during the larval stage, these eye-development genes remain quiescent until the adult stage, where all but *Pax-6* isoform A, *six3* and *atonal* are up-regulated (**Figure 5f**). These results are exceedingly surprising, given the conserved role of genes in retinal patterning. They suggest a novel role for these genes, or alternatively, these expression patterns could be the result of early expression of a retinal determination pathway followed by negative regulation by other genes to suppress eye development.

Photoreceptor and chemosensory system of *Archezogozetes longisetosus*

Unlike insects and crustaceans, chelicerates do not have compound eyes – with horseshoe crabs being an exception. Generally, mites are eyeless or possess one or two pairs of simple ocelli (Patten, 1887; Exner, 1989; Alberti and Coons, 1999; Harzsch *et al.*, 2006; Alberti and Moreno-Twose, 2012). Ocelli are common in Prostigmata and Endeostigmata, among Acariformes, as well Opilioacarida – the most likely sister group to the Parasitiformes – but are absent in most Oribatida, Astigmata, Mesostigmata and ticks (Walter and Proctor, 1998; Walter and Proctor, 1999; Norton and Fuangarworn, 2015; Norton and Franklin, 2018). This suggests that the presence of eyes might be an ancestral condition for both Acariformes and Parasitiformes, while more derived mites rely largely on chemical communication systems (Alberti and Coons, 1999).

In oribatid mites, detailed morphological and ultrastructural investigations have suggested that setiform sensilla are the most obvious sensory structures (**Figure 6a**) (Alberti, 1998; Alberti and Coons, 1999; Walter and Proctor, 1999). The trichobothria are very complex, highly modified (e.g., filiform, ciliate, pectinate, variously thickened or clubbed) no-pore setae which are anchored in a cup-like base and likely serve as mechanosensory structures. In contrast, the setal shafts of solenidia and eupathidia (**Figure 6a**) both possess pores (Alberti, 1998; Alberti and Coons, 1999; Walter and Proctor, 1999). Solenidia have transverse rows of small pores visible under a light microscope and likely function in olfaction, while the eupathidia have one or several terminal pores and likely are used as contact/gustatory sensilla (**Figure 6a**) (Alberti, 1998; Alberti and Coons, 1999). Previous work demonstrated that oribatid mites indeed use olfactory signals in the context of chemical communication and food selection (Shimano *et al.*, 2002; Raspotnig, 2006; Heethoff *et al.*, 2011a; Heethoff and Raspotnig, 2012; Brückner *et al.*, 2018a; Brückner *et al.*, 2018b).

Interestingly, detailed morphological and ultrastructural studies showed that light-sensitive organs exist in some Palaeosomata and Enarthronota (probably true eyes) as well as in Brachypylina (the secondary lenticulus), representing lower and highly derived oribatid mites, respectively (Alberti and Coons, 1999; Alberti and Moreno-Twose, 2012; Norton and Fuangarworn, 2015; Norton and Franklin, 2018). *Archezogozetes* and most other oribatids, however, are eyeless, yet there is scattered experimental and some anecdotal evidence that even these mites show some response to light and seem to avoid it ('negative phototropism' or 'negative phototaxis') (Trägårdh, 1933; Madge, 1965; Woodring, 1966; Walter and Proctor, 1999). Hence, we mined the genome of *Archezogozetes* for potential photoreceptor genes and found one gene of the *all-trans retinal peropsin* class and one gene related to the spider mite *rhodopsin-7-like* gene (**Figure 6b**). *Peropsin-like* genes are also present in other eyeless ticks. In jumping spiders, they encode for nonvisual, photosensitive pigments, while *rhodopsin-7* may be involved in basic insect circadian photoreception (Koyanagi *et al.*, 2008; Nagata *et al.*, 2010; Shen *et al.*, 2011; Eriksson *et al.*, 2013; Senthilan and Helfrich-Förster, 2016; Senthilan

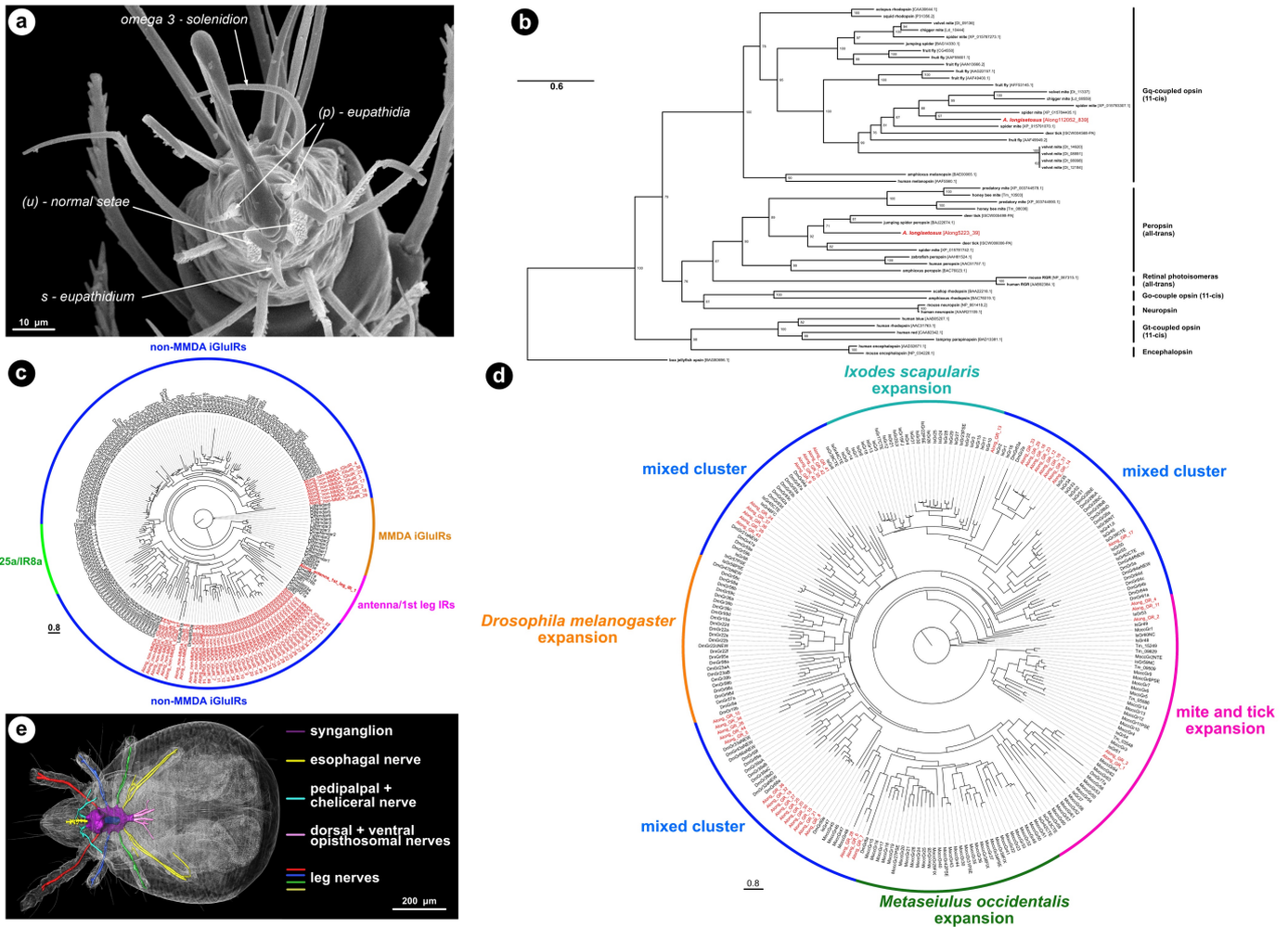


Figure 6 The sensory systems of *Archemozetes longisetosus* and phylogenetic analysis of selected photoreceptor and chemosensory genes. **a** – Scanning electron micrograph (SEM) showing the end of tarsus on *Archemozetes*' first leg. Images shows normal setae, but also modified chemosensory setae, namely eupathidia, both paired (p) and single (s), as well as an omega-3 solenidium. SEM picture courtesy of Michael Heethoff. **b** – Phylogeny and classification of opsin genes across the Metazoa, including those of several Chelicerata. The tree was constructed using a maximum likelihood approach (LG+F+R4 model) and rooted with a jelly fish opsin. *Archemozetes* sequences are depicted in red, *Drosophila* in turquoise; branch length unit is substitutions per site. **c** – Maximum likelihood phylogeny of ionotropic receptors and ionotropic glutamate receptors (LG+F+R6 model) of *Archemozetes* (Along), *Dinothrombium* (Dt), *Leptoithrombidium* (Ld), *Tetranychus* (Tu) and *Drosophila* (Dmel). IR25a/IR8a and antenna/1st leg IRs contain genes with known chemosensory function in *Drosophila*. The tree was rooted to the middle point; *Archemozetes* sequences are depicted in red, *Drosophila* in turquoise; branch length unit is substitutions per site. Bootstrap values can be found in the supplementary Figure S13. **d** – Maximum likelihood phylogenetic tree of gustatory receptors (JTT+F+R6 model) of *Archemozetes* (Along), *Ixodes* (Is), *Tropilaelaps* (Tm), *Metaseiulus* (Mocc) and *Drosophila* (Dmel). The tree was rooted to the middle point; *Archemozetes* sequences are depicted in red, *Drosophila* in turquoise; branch length unit is substitutions per site. Bootstrap values can be found in the supplementary Figure S14. **e** – Combined image of volume rendering (grey) and reconstructed nervous system of *Archemozetes* in dorsal view. Color-code corresponds to different parts of the nervous system, as depicted in the legend. The blue structure in the middle of the synganglion is the part of the esophagus which penetrates the synganglion. Scale bar: 200 μ m. Image courtesy of Sebastian Schmelzle based on data in (Hartmann *et al.*, 2016).

et al., 2019). Taken together, this might suggest that eyeless species like *Archemozetes* use *peropsin*- and *rhodopsin-7-like* genes for reproductive and diapause behaviors, or to maintain their circadian rhythm, as well as negative phototaxis.

The main sensory modality soil mites use is chemical communication *via* olfaction (Alberti, 1998; Alberti and Coons, 1999; Walter and Proctor, 1999; Raspotnig, 2006; Shen *et al.*, 2011; Brückner *et al.*, 2018a; Brückner *et al.*, 2018b). In contrast to insects, but similar to crustaceans and Myriapoda, mites do not have the full repertoire of chemosensory classes, they are missing odorant receptors and odorant-binding proteins (Table 2) (Maraun *et al.*, 2007; Raspotnig,

2009; Sánchez-Gracia *et al.*, 2009; Sánchez-Gracia *et al.*, 2011; Vieira and Rozas, 2011; Hoy *et al.*, 2016; Dong *et al.*, 2017; Dong *et al.*, 2018). Although chemosensory protein (CSP) encoding genes are absent in most mite genomes, we identified one gene encoding for such a protein in *Archezogozetes* and one CSP has been previously found in the deer tick (Table 2). Hence, *Archezogozetes* should primarily rely on gustatory receptors (GRs) and ionotropic receptors (IRs). Both the number of GRs (44 genes; **Figure 6d**) and IRs (1 gene; **Figure 6c**) was very well within the range of most mites and ticks and there was no evidence for any massive chemoreceptor expansion like in the spider mite (**Table 2**) (Ngoc *et al.*, 2016). This was surprising because *Archezogozetes*, like other acariform mites have many multiporous solenidia, present on all legs and the palp, but appear to only have a limited number of chemoreceptors.

Canonical ionotropic glutamate receptors (iGluRs) are glutamate-gated ion channels with no direct role in chemosensation, which come in two major subtypes: either NMDA iGluRs which are sensitive to N-methyl-D-aspartic acid (NMDA) or non-NMDA iGluRs. The latter group – at least in *Drosophila* – seems to have essential functions in synaptic transmission in the nervous system and have been associated with sleep and vision (Benton *et al.*, 2009; Sánchez-Gracia *et al.*, 2009; Croset *et al.*, 2010; Sánchez-Gracia *et al.*, 2011; Ngoc *et al.*, 2016). None of the IRs we found in the *Archezogozetes* genome belonged to the NMDA iGluRs and all, but one were classified as non-NMDA iGluRs (**Figure 6c**). Nothing is known about their functions in mites. It is, however, likely that they perform similar tasks in synaptic transmission in the brain and musculature. In *Drosophila* a specific set of chemosensory IRs, which do not bind glutamate, respond to acids and amines (*IR25a*), but also to temperature (*IR21a*, *IR93a*). For *Archezogozetes* we found one IR, similar to *IR21a* and *IR93a* of *Drosophila*, which fell into the antenna/1st leg IRs category (**Table 2; Figure 6c**) (Rytz *et al.*, 2013; Knecht *et al.*, 2016; Budelli *et al.*, 2019). This is consistent with an assumed limited contribution of IRs to the perception of chemical cues. Furthermore, it is so far unclear whether this specific IR is expressed in the first pair of legs (**Figure 6a and c**) in *Archezogozetes*, but similar genes seem to be expressed in the legs of other mite species (Dong *et al.*, 2017; Dong *et al.*, 2018), which could suggest a similar function as in the fruit fly.

GRs are multifunctional proteins and at least in insects they are responsible for the perception of taste, heat or volatile molecules (Montell, 2009). In *Archezogozetes* we found 44 GRs, 20 of which had full RNAseq support across all life stages, yet none of them appeared to belong to a species-specific expansion of the GR gene family (**Figure 6d**). Generally, it is unclear if GRs in *Archezogozetes* and other mites have similar functions as in insects, but the GR gene family is heavily expanded in many acariform mites and also is present in ticks (**Table 2**), suggesting an important biological role (Gulia-Nuss *et al.*, 2016; Hoy *et al.*, 2016; Ngoc *et al.*, 2016; Barrero *et al.*, 2017; Dong *et al.*, 2017; Dong *et al.*, 2018). This is supported by experimental evidence which suggested that ticks and other mites, including *Archezogozetes*, use chemical cues to find their host, communicate or discriminate food (Yunker *et al.*, 1992; Kuwahara, 2004; Raspotnig, 2006; Bunnell *et al.*, 2011; Gulia-Nuss *et al.*, 2016; Barrero *et al.*,

Table 2 Comparison of chemosensory receptor repertoires between *Archezogozetes longisetosus* and other arthropods. GR= gustatory receptor, OR= odorant receptor, IR= ionotropic receptor, OBP= odorant binding protein, CSP= chemosensory protein.

	Chemosensory receptors				
	GR	OR	IR	OBP	CSP
<i>Archezogozete longisetosus</i>	44	0	1	0	1
spider mite	689	0	4	0	0
deer tick	60	0	22	0	1
house spider	634	0	108	4	0
fruit fly	73	62	66	51	4

2017; Brückner *et al.*, 2018a; Brückner *et al.*, 2018b).

In general, not much is known about the nervous and sensory system of oribatid mites, or about sensory integration or the neuronal bases of their behavior (Alberti, 1998; Alberti and Coons, 1999; Norton, 2007). Modern methods like Synchrotron X-ray microtomography (SR μ CT) recently made it possible to investigate the organization and development of the nervous systems of oribatid mites (**Figure 6e**; (Hartmann *et al.*, 2016)). We here provide the first genomic resource for the investigation of the photo- and chemosensory systems of Oribatida (**Figure 6b-d**). In addition, we give a conservative estimate of olfactory gene numbers based on automated detection algorithm which should be refined by manual curation in the future (**Table 2**; see ‘**Materials & Methods**’). For instance, manual curation of olfactory genes revealed additional chemosensory receptor gene models in the spider mite (Ngoc *et al.*, 2016).

Horizontal gene transfer event sheds light on oribatid feeding biology

Horizontal gene transfer (HGT) is common among mites and other soil organisms (Grbić *et al.*, 2011; Mayer *et al.*, 2011; Faddeeva-Vakhrusheva *et al.*, 2016; Wu *et al.*, 2017; Dong *et al.*, 2018; Wybouw *et al.*, 2018). In some cases, genes that had been horizontally transferred now have pivotal biological functions. For instance, terpene and carotenoid biosynthesis genes in trombidiid and tetranychid mites show high homology with bacterial (terpene synthase) or fungal (carotenoid cyclase/synthase/desaturase) genes, suggesting horizontal gene transfer from microbial donors (Altincicek *et al.*, 2012; Dong *et al.*, 2018). At least the carotenoid biosynthesis genes in spider mites still code for functional enzymes and equip these phytophages with the ability to *de novo* synthesize carotenoids, which can induce diapause in these animals (Altincicek *et al.*, 2012; Bryon *et al.*, 2017).

Soil microarthropods like collembolans show numbers of horizontally transferred genes that are among the highest found in metazoan genomes, exceeded only by nematodes living in decaying organic matter (Crisp *et al.*, 2015; Faddeeva-Vakhrusheva *et al.*, 2016; Wu *et al.*, 2017). Interestingly, many HGT genes found in collembolans are involved in carbohydrate metabolism and were especially enriched for enzyme families like glycoside hydrolases, carbohydrate esterases or glycosyltransferases (Faddeeva-Vakhrusheva *et al.*, 2016; Wu *et al.*, 2017). All three enzyme families are involved in the degradation of plant and fungal cell walls (Latgé, 2007; Gilbert, 2010). Hence, it has been hypothesized that cell-wall degrading enzymes acquired by HGT are beneficial for soil organisms as it allowed such animals to access important food source in a habitat that is highly biased towards polysaccharide-rich resources (Mitreva *et al.*, 2009; Faddeeva-Vakhrusheva *et al.*, 2016; Faddeeva-Vakhrusheva *et al.*, 2017; Wu *et al.*, 2017).

To assess the degree of HGT in *Archezogetes* we first used blobtools (v1.0) (Laetsch and Blaxter, 2017) to generate a GC proportion vs read coverage plot of our long-read genome assembly, in order to remove contigs of bacterial origin (**Figure 7a**; 438 contigs). After Hi-C scaffolding we removed another 94 scaffolds of bacterial origin amounting to a total of ~ 9 Mb of contamination. Of the remaining scaffolds, candidate HGTs were identified using the Alien Index (Flot *et al.*, 2013; Thorpe *et al.*, 2018), where HGTs are those genes with blast homology (bit score) closer to non-metazoan than metazoan sequences (**supplementary Table S4**). We further filtered these HGT candidates to remove those that overlapped predicted repeats by $\geq 50\%$, resulting in 748 genes. As HGT become integrated into the host genome, they begin to mirror features of the host genome, including changes in GC content and introduction of introns (Lawrence, 1997). Comparing the GC content of the HGT candidates showed two distinct peaks, one at 53.9% and the other at 34.2%, slightly higher than the remaining *Archezogetes* genes, GC content of 31.8% (**Figure 7b**). Of the 429 HGT genes that shared similar GC content to the host genome, 78.8% had at least one intron (**Table S4**). In a final step, we used the gene expression data (RNAseq) to filter the list of all putative HGT genes and only retained candidates that were expressed in any life stage of *Archezogetes* ($n = 399$ HGT genes).

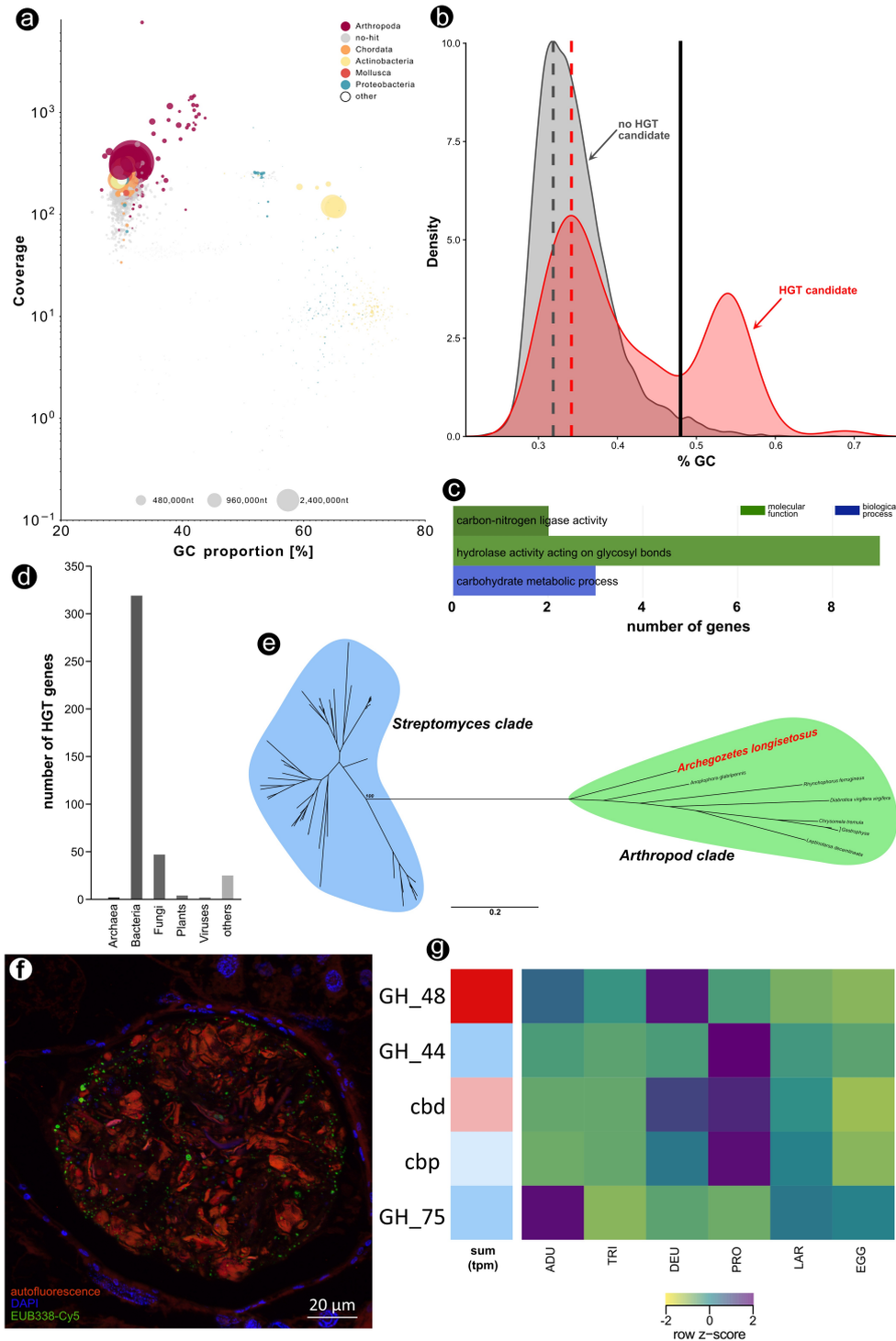


Figure 7 Horizontal gene transfer (HGT) and implications for the feeding biology of *Archegozetes longisetosus*. **a** – Blob-plot of the long-read genome assembly contigs plotting the read coverage against GC proportion [%]. Contigs are colored according to the taxonomic order of their best Megablast hit to the NCBI nucleotide database. Size of circle corresponds to the nucleotides per contigs. **b** – Comparison of the GC content of HGT and non-HGT genes. HGT genes shifted towards the host genome GC content indicate integration within the host genome while the higher GC content HGT genes might be the product of relatively recent HGT events. **c** – Enrichment of functional categories (GO terms) describing the molecular functions and biological processes related to the HGT candidate genes. **d** – Taxonomic origin of HGT. The category “others” includes mostly protozoan donor genes among other Eukaryotes. **e** – Unrooted maximum-likelihood tree (LG+F+R4 model) of glycoside hydrolase family 48 members (GH_48) from *Streptomyces* bacteria and HGT genes from other arthropods as well as *Archegozetes*. Bootstrap values and the full tree can be found in the supplementary Figure S15. The scale bar denotes substitutions per site. **f** – Fluorescence *in situ* hybridization (FISH) micrograph of a food bolus in the mites’ alimentary tract. The food material (wheat grass power) is enclosed in a peritrophic membrane and there is a high bacterial prevalence in the food bolus. Image courtesy of Benjamin Weiss and Martin Kaltenpoth. **g** – RNAseq support of HGT candidates related to cell wall degrading enzymes. The first block (single column) shows the overall RNA expression (tpm) of the HGT in all life stages; red denotes high total expression, while blue depicts low total expression. The second block (six columns) shows the expression (row z-score based on tpm) of the same HGT candidates across the different life stages of *Archegozetes*. Abbreviations: GH_48= glycoside hydrolase family 48, GH_44= glycoside hydrolase family 44, cbd= cellulose-binding domain, cbp= cellulose-binding protein, GH_75= glycoside hydrolase family 75.

The majority of HGT candidates were of bacterial origin (79.9%), followed by genes likely acquired from fungi (11.8%), while transfer from Archaea, plants, virus, and other sources was comparatively low (**Figure 7d**). This composition of HGT taxonomic origin is different from genes found in collembolans, which appear to have acquired more genes of fungal and protist origin (Faddeeva-Vakhrusheva *et al.*, 2016; Faddeeva-Vakhrusheva *et al.*, 2017; Wu *et al.*, 2017). Subsequently, we performed an over-representation analysis of GO terms associated with these genes. We found an over-representation of genes with GO terms related to carbon-nitrogen ligase activity and hydrolase activity on glycosyl bonds (molecular function; **Figure 7c**) as well as carbohydrate metabolism (biological process; **Figure 7c**). This provides a first line of evidence that *Archezogozetes* possess HGT related to plant- and fungal cell wall degradation similar to collembolans, yet mite and collembolan genes did not appear to be direct homologs. For instance, genes related to cell wall degrading enzymes in mites appear to be acquired by HGT from *Streptomyces* bacteria (**Figure 7**) while they are of fungal origin in collembolans (Faddeeva-Vakhrusheva *et al.*, 2016; Faddeeva-Vakhrusheva *et al.*, 2017; Wu *et al.*, 2017).

As mentioned previously, oribatid mites are among the few Chelicerata that ingest solid food and are primary- and secondary decomposers feeding on dead plant material and fungi (Cohen, 1995; Norton, 2007; Shultz, 2007; Dunlop and Alberti, 2008; Heethoff and Norton, 2009; Maraun *et al.*, 2011). It was argued for decades that the enzymes necessary to break down these polysaccharide-rich resources originate from the mite's gut microbes (Stefaniak, 1976; 1981; Smrž, 1992; Siepel and de Ruyter-Dijkman, 1993; Smrž, 2000; Smrž and Norton, 2004; Smrž and Čatská, 2010). Microbes might be mixed with the food in the ventriculus and digest it while passing through the alimentary tract as food boli enclosed in a peritrophic membrane (see **Figure 7f** for an example) (Stefaniak, 1976; 1981). However, screening the HGT candidate list for potential cell-wall degrading enzymes and mapping their overall and life-stage specific expression in *Archezogozetes* using the RNAseq reads, revealed at least five HGT genes related to polysaccharide breakdown (**Figure 7g**). We found that specifically members of the *glycoside hydrolases family 48* and *cellulose-binding domain* genes showed high expression in most life stages - the egg being an obvious exception (**Figure 7g**). Moreover, the majority of these genes were flanked by a predicted metazoan gene, suggesting host transcriptional regulation (**Table S4**).

In a last step we blasted the highly expressed HGT candidates (**Figure 7g**) against the non-redundant protein sequence database, aligned the sequences with the highest alignment score and performed a phylogenetic maximum likelihood analysis. For the highest expressed HGT related to cell-wall-degrading enzymes (*glycoside hydrolases family 48 gene*), we recovered that the *Archezogozetes* sequences was well nested within a clade of *GH 48* sequences from herbivores beetles (McKenna *et al.*, 2019), which appear to be related to similar genes from various *Streptomyces* (**Figure 7e**) and we reconstructed similar phylogenies for other highly expressed HGT candidates (**supplementary Figure S12**). All the sequences of beetle *glycoside hydrolases family 48* members (**Figure 7e**) were included in recent studies arguing for a convergent horizontal transfer of bacterial and fungal genes that enabled the digestion of lignocellulose from plant cell walls in herbivores beetles (McKenna *et al.*, 2016; McKenna *et al.*, 2019). They showed that phytophagous beetles likely acquired all genes of the *GH 48 family* from Actinobacteria (including *Streptomyces*) (McKenna *et al.*, 2019) and our phylogenetic analysis (**Figure 7e**) revealed the same pattern as well as a highly similar tree topology (compare to Fig 3B in (McKenna *et al.*, 2019)).

Overall, our findings indicate that genes encoding for enzymes in *Archezogozetes* capable of degrading plant and fungal cell walls were likely horizontally transferred from bacteria (likely *Streptomyces*). Bacterial symbionts and commensal living in the mites' gut are still likely to contribute to the breakdown of food (**Figure 7f**). Yet, the high expression of genes encoding cell-wall degrading enzymes (**Figure 7g**) as well as the evolutionary analyses of such genes (**Figure 7e**) suggest that *Archezogozetes* – and potentially many other oribatid mites – are able to exploit polysaccharide-rich resources like dead plant material or chitinous fungi

without microbial aid. Enzymological and microscopical investigation of *Archezogetes* have suggested that certain digestive enzymes (chitinase and cellulase) are only active when the mites consume a particular type of food (e.g. algae, fungi or filter paper) (Smrž and Norton, 2004). These results were interpreted as evidence that these enzymes are directly derived from the consumed food source (Smrž and Norton, 2004). By contrast, we argue that this instead confirms our findings of HGT: upon consumption of food containing either chitin or cellulose, gene expression of polysaccharide-degrading enzymes starts, and proteins can readily be detected. Further enzymological studies have placed oribatid mites in feeding guilds based on carbohydrase activity and also found highly similar enzyme activity between samples of mites from different times and locations (Luxton, 1972; 1979; 1981; 1982; Siepel and de Ruiter-Dijkman, 1993). Future functional studies can disentangle the contribution of the host and microbes to cell wall digestion and novel metabolic roles of the HGTs identified here.

Biosynthesis of monoterpenes – a common chemical defense compound class across oribatid mite

Oribatid and astigmatid mites are characterized by a highly diverse spectrum of natural compounds that are produced by and stored in so-called oil glands (for an example see **Figure 8a**) (Raspotnig, 2009; Raspotnig *et al.*, 2011; Heethoff *et al.*, 2016). These paired glands are located in the opisthosoma (i.e., the posterior part of chelicerate arthropods, analogous to the abdomen of insects) and are composed of a single-cell layer invagination of the cuticle (**Figure 8f**). As previously mentioned, mites use chemicals produced by these glands to protect themselves against environmental antagonists (predators or microbes) or use them as pheromones (Shimano *et al.*, 2002; Raspotnig, 2006; 2009; Heethoff *et al.*, 2011a; Heethoff and Raspotnig, 2012; Brückner *et al.*, 2015; Heethoff and Rall, 2015). The monoterpene aldehyde citral – a stereoisomeric mixture of geranial ((*E*)-3,7-dimethylocta-2,6-dienal) and neral ((*Z*)-3,7-dimethylocta-2,6-dienal) – and its derivatives are widely detected compounds in glandular secretions of oribatids and astigmatids (Sakata *et al.*, 1995; Sakata, 1997; Kuwahara *et al.*, 2001; Sakata and Norton, 2001; Sakata and Norton, 2003; Kuwahara, 2004; Raspotnig *et al.*, 2004; Koller *et al.*, 2012). These monoterpenes have been called “astigmatid compounds” (Sakata and Norton, 2001) as they characterize the biochemical evolutionary lineage of major oribatid mite taxa (Mixonomata and Desmonomata) and almost all investigated astigmatid mites (Alberti, 1984; Sakata, 1997; Sakata and Norton, 2001; Kuwahara, 2004; Raspotnig, 2009).

The chemical cocktail released by *Archezogetes* consists of a blend of 10 compounds (**Figure 8a**) including two terpenes (approx. 45%) – neral and neryl formate – six hydrocarbons (approx. 15%) and two aromatic compounds (approx. 40%) (Sakata and Norton, 2003; Brückner and Heethoff, 2017). The hydrocarbons likely serve as solvents, while the terpenes and aromatics are bioactive compounds used in chemical alarm and defense (Shimano *et al.*, 2002; Sakata and Norton, 2003; Raspotnig, 2006; Heethoff *et al.*, 2011a). Recently, it was shown that *Archezogetes* synthesizes the two aromatic compounds using a polyketide-like head-to-tail condensation of (poly)- β -carbonyls *via* a horizontally acquired putative polyketide synthase (Brückner *et al.*, 2020). Studies in Astigmata found that the monoterpenes of these mites appeared to be made *de novo* from (poly)- β -carbonyls as well and one study identified a novel geraniol dehydrogenase (GeDH), unrelated to those of bacteria, in *Carpoglyphus lactis* (Morita *et al.*, 2004; Noge *et al.*, 2005; Noge *et al.*, 2008). To learn about the biosynthesis of astigmatid compounds in *Archezogetes* and demonstrate the mite’s applicability as research model for biochemical pathway evolution, we used the novel genomic resources presented in this study.

First, we delineated the basic biochemical reactions likely to happen in the *Archezogetes* gland through a stable-isotope labeling experiment. We supplemented the diet of the mite with food containing 25% heavy $^{13}\text{C}_6$ D-glucose and 10% antibiotics (a combination of three different antibiotics was fed, because this mixture is able to eliminate nearly all qPCR and

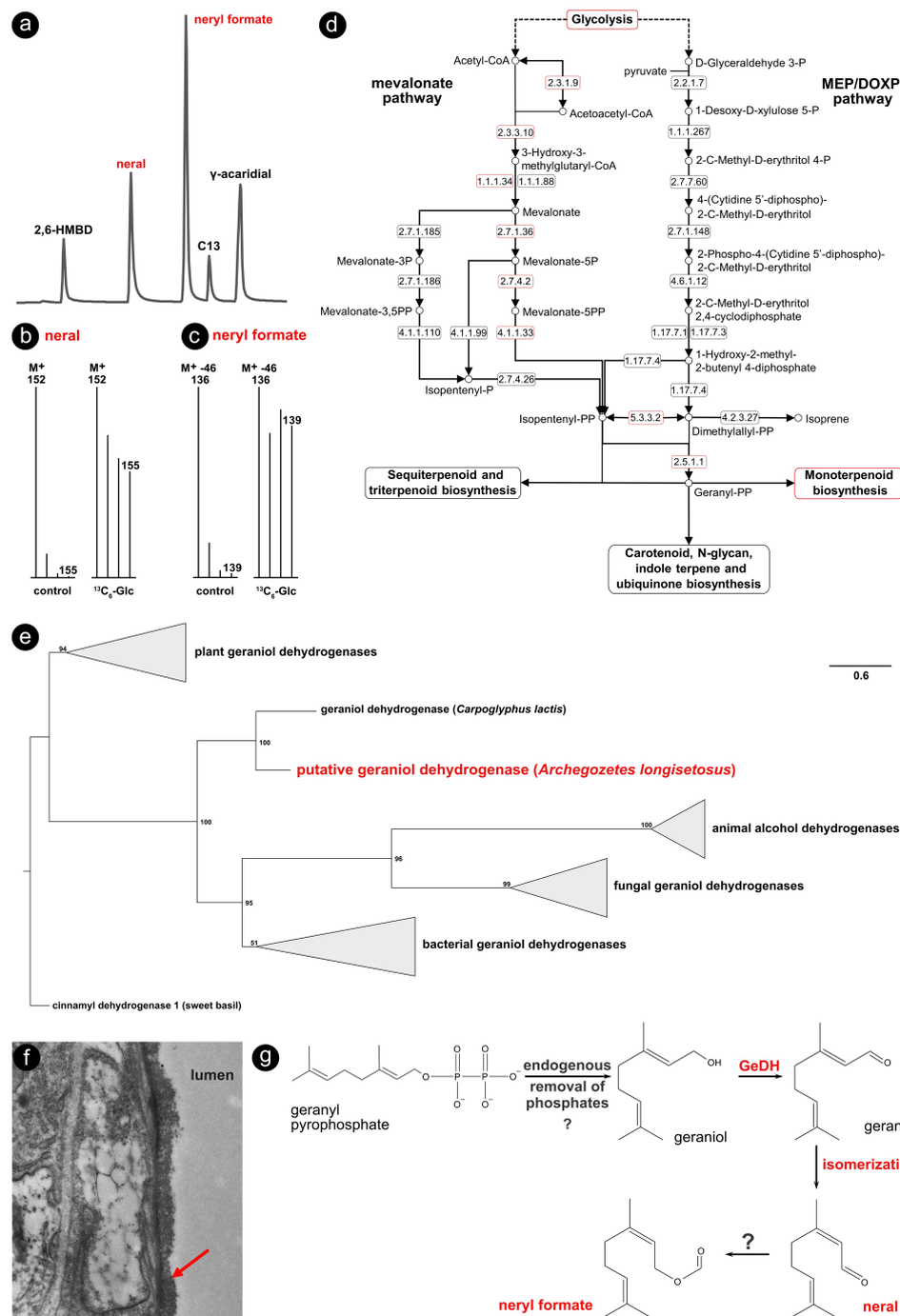


Figure 8 Reconstruction of the biosynthetic pathway leading to monoterpenes in *Arcegozetes longisetosus*. **a** – Representative gas chromatogram of the mite’ gland content; in order of retention time: 2-hydroxy-6-methyl-benzaldehyde (2,6-HMBD), neral ((*Z*)-3,7-dimethylocta-2,6-dienal) neryl formate ((*Z*)-3,7-dimethyl-2,6-octadienyl formate), tridecane, 3-hydroxybenzene-1,2-dicarbaldehyde (γ -acaridial). Further alkanes/alkenes (pentadec-7-ene, pentadecane, heptadeca-6,9-diene, heptadec-8-ene, heptadecane) are not shown. Monoterpenes are marked in red. **b and c** – Representative mass spectra of neral (**b**) and neryl formate (**c**) extracted from defensive glands of mites fed with unlabeled wheatgrass powder (control), or wheatgrass infused with $^{13}\text{C}_6$ -labelled glucose recorded in single-ion mode. The mass spectra for neral (**b**) shows the M^+ -ion series, while the spectra for neryl formate (**c**) show the diagnostic ion series at $[\text{M}-46]^+$. Mites fed with the $^{13}\text{C}_6$ glucose infused wheatgrass showed enriched ions. **d** – KEGG reference pathway map for terpenoid backbone biosynthesis. Mapping genes from the *Arcegozetes* genome encoding for pathway enzymes (labeled in red) revealed that the mite can produce geranyl pyrophosphate (GPP) via the mevalonate pathway from precursors provided by glycolysis. Enzymes names correspond to EC numbers: 2.3.1.9= acetyl-CoA C-acetyltransferase; 2.3.3.10= hydroxymethylglutaryl-CoA synthase; 1.1.1.34= hydroxymethylglutaryl-CoA reductase; 2.7.1.36= mevalonate kinase; 2.7.4.2= phosphomevalonate kinase; 4.1.1.33= diphosphomevalonate decarboxylase; 5.3.3.2= isopentenyl-diphosphate delta-isomerase; 2.5.1.1= farnesyl diphosphate synthase. **e** – Maximum-likelihood tree (LG+F+R4 model) based on an alignment of plant, fungal and bacterial geraniol dehydrogenases, animal alcohol dehydrogenase and two mite (*Carpoglyphus lactis* and *Arcegozetes*) geraniol dehydrogenases (GeDH). Bootstrap values (based on 1000 replicates) are indicated along branches and the scale bar denotes substitutions per site. The tree was rooted by the outgroup cinnamyl dehydrogenase from sweet basil. **f** – Ultrastructure of the gland-tissue of *Arcegozetes*, as observed by transmission electron microscopy (TEM). Red error shows the border between the gland cell and the glandular lumen. TEM picture courtesy of Michael Heethoff. **g** – Proposed biochemical pathway scenario leading to neral and neryl formate in *Arcegozetes* starting with GPP from the terpenoid backbone biosynthesis.

FISH detectable bacteria found on the food and in the alimentary tract (Brückner *et al.*, 2020)). To examine the incorporation of heavy $^{13}\text{C}_6$ D-glucose and its metabolic products into neral (**Figure 8b**) and neryl formate (**Figure 8c**), we compared selected fragment ions (M^+ and M^+-46 , respectively) using single ion mass spectrometry. Both neral and neryl formate showed consistent enrichment in their M^+ to $[\text{M}+4]^+$ and $[\text{M}-46]^+$ to $[\text{M}-46+4]^+$ -ion series, indicating that *Archezogozetes* used glycolysis breakdown products of $^{13}\text{C}_6$ D-glucose for the biosynthesis of their monoterpenes. We then used the OGS mapped to KEGG metabolic pathways (Kanehisa *et al.*, 2007) to reconstruct the backbone synthesis of terpenes in *Archezogozetes* (**Figure 8d**). We found mite genes, which suggest that *Archezogozetes* synthesizes geranyl pyrophosphate (GPP) – the input substrate for further monoterpene synthesis – *via* the mevalonate pathway using the Mevalonate-5P to Isopentenyl-PP route (**Figure 8d**). The Mevalonate-5P pathway is used in most higher eukaryotes as compared to the Mevalonate-3P pathway in Archaea and the MEP/DOXP pathway in bacteria, some plants and apicomplexan protists (Trapp and Croteau, 2001; Eisenreich *et al.*, 2004; Breitmaier, 2006; Degenhardt *et al.*, 2009; Miziorko, 2011; Oldfield and Lin, 2012). This likely excludes any horizontal gene transfer of mevalonate pathway genes as *Archezogozetes* uses enzymes similar to those of other animals.

The biosynthesis of monoterpenes not only depends on very widespread enzymes, but also requires more specific enzymes downstream of GPP (Trapp and Croteau, 2001; Breitmaier, 2006; Degenhardt *et al.*, 2009). For instance, *Carpoglyphus lactis* expresses a unique geraniol dehydrogenase (GeDH) – catalyzing the oxidation of geraniol to geranial – different from all previously characterized geraniol-related and alcohol dehydrogenases (ADHs) of animals and plants (Noge *et al.*, 2008). We used the functionally validated *Carpoglyphus*-GeDH (Noge *et al.*, 2008), blasted its sequence against the *Archezogozetes* OGS and found a homologous sequence. We used both mite sequences in an alignment with plant, fungal and bacterial GeDHs and animal ADHs and constructed a maximum likelihood phylogeny (**Figure 8e**). Similar to the previous analysis including only *Carpoglyphus*-GeDH, we found that the *Al-GeDH* represent a new class of geraniol dehydrogenases different from those in plants, fungi or bacteria and not nested within animal ADHs (**Figure 8e**). This is why we hypothesize that *Al-GeDH* is a novel expansion of the geraniol dehydrogenases gene family and has not been acquired by horizontal gene transfer, like other biosynthesis and digestive enzymes in *Archezogozetes* (**Figure 7**; (Brückner *et al.*, 2020)).

Based on our mass spectrometry data of stable isotopes and genomic analysis, we propose that the following biochemical pathway leading to monoterpenes is of oribatid mites (**Figure 8f and g**): geraniol is likely to be synthesized from GPP – the universal precursor of all monoterpenes – either enzymatically by a geraniol synthase (GES) or a diphosphate phosphatase (DPP), but possibly also endogenously by dephosphorylation of GPP (Oswald *et al.*, 2007; Zhou *et al.*, 2014; Liu *et al.*, 2015; Beran *et al.*, 2019). For *Archezogozetes*, we could not find any *GES* or specific *DPP* in the OGS, thus geraniol might be formed from GPP *via* endogenous dephosphorylation, but further research is required to verify or falsify this hypothesis. Subsequently, geraniol is oxidized to geranial by the pervious described *GeDH* (**Figure 8e**) and readily isomerized to neral. Trace amounts of geranial have been found in *Archezogozetes* and it is common among other oribatid and astigmatid mites, supporting this idea (Kuwahara, 2004; Raspotnig *et al.*, 2004; Raspotnig *et al.*, 2008; Koller *et al.*, 2012). Also, there is no evidence that geraniol is converted into nerol, or that neral is formed directly *via* oxidation of nerol (Morita *et al.*, 2004; Noge *et al.*, 2005; Noge *et al.*, 2008). The most parsimonious explanation for neryl formate synthesis would be an esterification of the corresponding terpene alcohol nerol. There is, however, no evidence of nerol in the traces of any oribatid or astigmatid mite species (Kuwahara, 2004; Raspotnig, 2009; Raspotnig *et al.*, 2011). Aliphatic non-terpene formates in Astigmata are synthesized by dehomologation and generation of a one-carbon–shorter primary alcohol from an aldehyde *via* hydrolysis of formate in a biological Baeyer–Villiger oxidation catalyzed by a novel, uncharacterized enzyme (Shimizu *et al.*, 2017). A similar reaction to synthesize terpene formates is unlikely, as the terpenoid backbone would be shortened by one-carbon and this does not happen in any possible

scenario. The discovery of this Baeyer–Villiger oxidation mechanism, however, highlights the probability that there are many very unusual reactions that remain to be discovered in oribatid mites (Brückner *et al.*, 2020).

Conclusion

The integrated genomic and transcriptomic resources presented here for *Archegozetes longisetosus* allowed a number of insights into the molecular evolution and basic biology of decomposer soil mites. Our analysis of an oribatid mite genome also provides the foundation for experimental studies building on the long history of *Archegozetes* as a chelicerate model organism, which now enters the molecular genetics era (Aoki, 1965; Palmer and Norton, 1992; Norton *et al.*, 1993; Heethoff *et al.*, 2013). This includes the study of biochemical pathways, biochemistry, neuroethological bases of food searching behavior, and environmental impacts on genomes of complex, clonal organisms.

Our evolutionary comparisons across the Chelicerata revealed interesting patterns of genome evolution and how horizontal gene transfer might have shaped the feeding mode of soil mites. We also showed how oribatid glandular biology and chemical ecology are reflected in the genome. The community of researchers studying the fundamental biology of oribatid, and other free-living, non-parasitic mites is growing. We think that providing these genomic and transcriptomic resources can foster a community effort to eventually allowing for basic molecular research on these mites.

Key priorities for a future community research effort include i) sequencing organ-specific transcriptomic data, ii) developing tools for genetic interrogation (RNAi or CRISPR/CAS9), iii) establishing reporter lineages with germ-line stable modifications (e.g. GAL4/UAS misexpression systems), and iv) constructing an whole-animal single-cell RNAseq expression atlas. Please do not hesitate to contact the corresponding author if you want to start your own culture of *Archegozetes*. We will be happy to provide you with starter specimens for free and share rearing protocols with you.

Material and methods

Mite husbandry

The lineage ‘ran’ (Heethoff *et al.*, 2013) of the pantropical, parthenogenetic oribatid mite *Archegozetes longisetosus* was used in this study. Stock cultures were established in 2015 from an already existing line and fed with wheat grass (*Triticum* sp.) powder from Naturya. Cultures were maintained at 20–24 °C and 90% relative humidity. Sterilized water and 3–5 mg wheat grass were provided three times each week.

DNA extraction and Illumina sequencing

For the short-read library, DNA was extracted from ~200 mites that were taken from the stock culture, starved for 24 h to avoid possible contamination from food in the gut, subsequently washed with 1% SDS for 10 s. For extraction of living specimens, we used the Quick-DNA Miniprep Plus Kit (Zymo Research) according to the manufacturer’s protocol. Amounts and quality of DNA were assessed with Qubit dsDNA HS Kit (Thermo Fisher) and with NanoDrop One (Thermo Fisher) with target OD 260/280 and OD 260/230 ratios of 1.8 and 2.0–2.2, respectively. Extracted DNA was shipped to Omega Bioservices (Norcross, GA, USA) on dry ice for library preparation and sequencing. DNA library preparation followed the KAPA HyperPrep Kit (Roche) protocol (150 bp insert size), and 200 million reads were sequenced as 150bp paired-end on a HighSeq4000 (Illumina) platform.

High-molecular weight DNA isolation and Nanopore sequencing

Genomic DNA was isolated from ~300-500 mites starved for 24 h using QIAGEN Blood & Cell Culture DNA Mini Kit. Briefly, mites were flash frozen in liquid nitrogen and homogenized with a pestle in 1 ml of buffer G2 supplemented with RNase A and Proteinase K at final concentrations of 200 ng/ μ l and 1 μ g/ μ l, respectively. Lysates were incubated at 50°C for 2 h, cleared by centrifugation at 5 krpm for 5 min at room temperature and applied to Genomic tip G/20 equilibrated with buffer QBT. Columns were washed with 4 ml of buffer QC and genomic DNA was eluted with 2 ml of buffer QF. DNA was precipitated with isopropanol, washed with 70% EtOH and resuspended in 50 μ l of buffer EB. DNA was quantified with Qubit dsDNA HS Kit (Thermo Fisher) and the absence of contaminants was confirmed with NanoDrop One (Thermo Fisher) with target OD 260/280 and OD 260/230 ratios of 1.8 and 2.0-2.2, respectively. DNA integrity was assessed using Genomic DNA ScreenTape kit for TapeStation (Agilent Technologies).

Libraries for nanopore sequencing were prepared from 1 μ g of genomic DNA using 1D Genomic DNA by Ligation Kit (Oxford Nanopore) following manufacturer's instructions. Briefly, unfragmented DNA was repaired and dA tailed with a combination of NEBNext FFPE Repair Mix (New England Biolabs) and NEBNext End repair/dA-tailing Module (New England Biolabs). DNA fragments were purified with Agencourt AMPure XP beads (Beckman Coulter) and Oxford Nanopore sequencing adapters were ligated using NEBNext Quick T4 DNA Ligase (New England Biolabs). Following AMPure XP bead cleanup, ~500 ng of the library was combined with 37.5 μ L of SQB sequencing buffer and 25.5 μ l of loading beads in the final volume of 75 μ l and loaded on a MinION Spot-ON Flow Cell version R9.4 (Oxford Nanopore). Two flow cells were run on MinION device controlled by MinKNOW software version 3.1.13 for 48 hours each with local basecalling turned off generating 9.7 and 5.1 GB of sequence data. Post run basecalling was performed with Guppy Basecalling Software, version 3.4.5 (Oxford Nanopore). After filtering low quality reads ($Q < 7$), the combined output of the two runs was 13.69 GB and 4.7 million reads.

Hi-C library preparation

A sequencing library was constructed from approximately 150 mg of mites, or ~1000 individuals, using the Proximo Hi-C kit following specific instructions for insects (Phase Genomics). Library quantity and quality was assessed using Qubit dsDNA High Sensitivity Assay (Thermo Fisher) and bioanalyzer (Agilent), respectively. Two hundred million 150bp paired-end reads were sequenced on a NextSeq 2000 platform (Illumina) by the Millard and Muriel Jacobs Genetics and Genomics Laboratory at California Institute of Technology.

Genome assembly and contamination filtering

Read quality was assessed using FastQC v0.11.8 (Andrews, 2010). Illumina adapters, low-quality nucleotide bases (phred score below 15) from the 3' and 5' ends and reads shorter than 50 bp were removed using cutadapt v1.18 (Martin, 2011). From the filtered reads, *in silico* genome size estimates were calculated using *k-mer* based tools kmergenie v.1.7048 (Chikhi and Medvedev, 2014), GenomeScope v1.0 (Vurture *et al.*, 2017), and findGSE v0.1.0 R package (Sun *et al.*, 2018). The latter two required a *k-mer* histogram computed by jellyfish v2.2.10 (Marçais and Kingsford, 2011) with *k-mer* size of 21. The long-read genome was assembled using 4.7 million reads from two MinION runs (60x coverage) using Canu v1.8 with default settings and setting the expected genome size to 200 Mb (Koren *et al.*, 2017). To improve assembly quality, paired end Illumina reads were mapped to the genome with BWA aligner (Li and Durbin, 2009) using BWA-MEM algorithm and polished with Pilon v. 1.23 with '—changes' and '—fix all' options (Walker *et al.*, 2014). Assembled contigs identified as bacterial and fungal contaminants based on divergent GC content from most *Archeogozetes*

contigs, high coverage and blast homology to the nt database (downloaded February 2019, Evalue $1e^{-25}$) were removed using Blobtools v1.0 (Laetsch and Blaxter, 2017).

Duplicate contigs, or haplotigs, of the long-read assembly were removed using `purge_haplotigs` v.1.1 (Roach *et al.*, 2018). The short-read data was mapped to the long-read assembly using `minimap2` (Li, 2018) and then filtered based on read coverage (parameters: `-l 25 -m 230 -h 500`). The Hi-C sequencing data was then mapped to the purged assembly with BWA (Li and Durbin, 2009). The Hi-C contact map was generated and corrected from the mapped reads using `hicexplorer` (Ramírez *et al.*, 2018) with a bin size of 10000 and z-score thresholds of -1.5 and 3 for correction. The contact map was then used by `HiCAssembler` (Renschler *et al.*, 2019) along with the purged assembly for scaffolding (parameters: `-min_scaffold_length 300000 -bin_size 20000 -num_iterations -3 split_positions_file split.bed`). The utility tool `plotScaffoldInteractive` was used to identify coordinates of misassemblies to manually split in the assembly process (`split.bed`).

Identification, classification, and masking of repetitive element

Repetitive elements in the genome *Archegozetes* were identified using a species-specific library generated with RepeatModeler v 1.0.11 (Smit and Hubley, 2008; Bao *et al.*, 2015) and MITE tracker (Crescente *et al.*, 2018) and annotated by RepeatClassifier, a utility of the RepeatModeler software that uses the RepBase database (version Dfam_Consensus-20181026). Unclassified repeat families from both programs were run through CENSOR v 4.2.29 (Kohany *et al.*, 2006) executable `cursor.ncbi` against the invertebrate library v 19.03 to provide further annotation. Predicted repeats were removed if they had significant blast homology (E-value $1e^{-5}$) to genuine proteins in the NCBI nr database and/or a local database of arthropod genomes (*Drosophila melanogaster*, *Tribolium castaneum*, *Tetranychus urticae*, *Leptotrombidium deliense*, *Dinothrombium tinctorium*, *Sarcoptes scabiei*, *Euroglyphus maynei*, *Galendromus occidentalis*, *Dermatophagoides pteronyssinus*). Unclassified repeats with blast homology to known TEs were retained whereas those with no blast homology were removed (Petersen *et al.*, 2019). The remaining repeat families were combined with the Arthropoda sequences in RepBase and clustered using `vsearch` v 2.7.1 (`-iddef 1 -id 0.8 -strand both`; (Rognes *et al.*, 2016)). The filtered repeat library was used to soft mask the *A. longisetosus* Hi-C assembly using RepeatMasker v 4.07 (Smit *et al.*, 1996-2010). A summary of the masked repeat content was generated using the “`buildSummary.pl`” script, the Kimura sequence divergence calculated using the “`calcDivergenceFromAlign.pl`” script and the repeat landscape visualized using the “`createRepeatLandscape.pl`” script, all utilities of RepeatMasker.

Gene prediction and annotation

Both *ab initio* and reference-based tools were used for gene prediction of the long-read assembly using modified steps of the funannotate pipeline (Palmer and Stajich, 2017). The *ab initio* tool GeneMark-ES v4.33 (Ter-Hovhannisyan *et al.*, 2008) was used along with reference based tools BRAKER v2.1.2 (Bruna *et al.*, 2020) using RNAseq reads discussed below and PASA v 2.3.3 (Haas *et al.*, 2008) using genome-guided transcriptome assembly from Trinity described below. Lastly, *Tetranychus urticae* gene models from the NCBI database (GCF_000239435.1) were aligned to the contigs using GeMoMa (Keilwagen *et al.*, 2019). All gene predictions were combined in EvidenceModeler (Haas *et al.*, 2008) with the following weights: GeMoMa = 1, PASA = 10, other BRAKER = 1, and GeneMark = 1. Predicted tRNAs using tRNAscan-SE v 2.0.3 (Chan and Lowe, 2019) were combined with the gene predictions in the final gene feature format (GFF) file and filtered for overlap using bedtools (Quinlan and Hall, 2010) `intersect` tool (Quinlan and Hall, 2010). After constructing the Hi-C assembly, gene predictions above were assigned to the new coordinates using a combination of LiftOff (Shumate and Salzberg, 2021) and GeMoMa (Keilwagen *et al.*, 2019), both with default settings. The new assembly was also reanalyzed with PASA using the transcriptome as described above to recover incomplete

open-reading frames from the lift over procedure. The gene predictions were combined with EvidenceModeler with the following weights: GeMoMa= 5, Liftoff= 5, and PASA = 10.

The predicted genes were searched against the NCBI nr (February 2019) (Pruitt *et al.*, 2005), SwissProt (February 2019) (Bairoch and Apweiler, 2000), a custom-made Chelicerata database including genomes of *Tetranychus urticae*, *Leptotrombidium deliense*, *Dinotrombidium tinctorium*, *Sarcoptes scabiei*, *Euroglyphus maynei*, *Galendromus occidentalis*, *Metaseiulus occidentalis*, *Dermatophagoides pteronyssinus*, *Trichonephila clavipes*, *Stegodyphus mimosarum*, *Centruroides sculpturatus*, *Ixodes scapularis* and *Parasteatoda tepidariorum* (all downloaded Feb 2019), PFAM (v 32, August 2018) (Bateman *et al.*, 2004), merops (v 12, October 2017) (Rawlings *et al.*, 2010) and CAZY (v 7, August 2018) (Cantarel *et al.*, 2009) databases. The results of the hmm-based (Eddy, 2011) PFAM and CAZY searches were filtered using cath-tools v 0.16.2 (<https://cath-tools.readthedocs.io/en/>; E-value $1e^{-5}$) and the blast-based searches were filtered by the top hit (E-value $1e^{-5}$ threshold). Predicted genes were also assigned to orthologous groups using eggNOG-mapper (Huerta-Cepas *et al.*, 2017). Gene annotation was prioritized by the SwissProt hit if the E-value $< 1e^{-10}$ followed by NCBI annotation, the custom Chelicerata database and if no homology was recovered, then the gene was annotated as, “hypothetical protein”.

Analysis of the official gene set (OGS)

To allow the OGS to be used as resources for functional studies, we assigned functional categories based on Gene Ontology (GO) and the Kyoto Encyclopedia of Genes and Genomes (KEGG) (Kanehisa and Goto, 2000; Consortium, 2004). GO terms for the respective genes models of the OGS were assigned based on the gene id with highest homology from the SwissProt database or NCBI nr database (Bairoch and Apweiler, 2000; Pruitt *et al.*, 2005). A custom database of GO terms was created with makeOrgPackage function in the R package AnnotationForge v1.26.0 (Carlson and Pagès, 2019). Over-representation analysis of GO terms was tested using the enrichGO function in the R package clusterProfiler v3.12.0 (Yu *et al.*, 2012) with a hypergeometric distribution and a Fisher’s Exact test. P-values were adjusted for multiple comparisons using false discovery rate correction (Benjamini and Hochberg, 1995). Resulting enriched GO terms were processed with GO slim (Consortium, 2019) and the final list of over represented GO terms was used to plot the number of genes in a respective category.

KEGG orthology terms were assigned from single-directional best hit BLAST searches of each gene model on the KEGG Automatic Annotation Server (Moriya *et al.*, 2007). Additionally, we ran GhostKOALA (Kanehisa *et al.*, 2016) (GHOSTX searches for KEGG Orthology And Links Annotation) to obtain KEGG orthology terms. Compared to conventional BLAST searches, GhostKOALA is about 100 times more efficient than BLAST to remote homologs by using suffix arrays (Suzuki *et al.*, 2014).

Orthology and phylogenomic analyses

Orthologs of *A. longisetosus*, other species within Acari, Chelicerata and the fruit fly *Drosophila* were identified using OrthoFinder v 2.3.3 (-M msa -A mafft -T fasttree; (Emms and Kelly, 2015)). Prior to running OrthoFinder, isoform variants were removed from the gene predictions using CD-Hit (Fu *et al.*, 2012). Trees of orthogroups with at least 80% of taxa present ($n=4,553$) were constructed using fasttree v 2.1.10 (Price *et al.*, 2010), trimmed with TrimAl v 1.4.1 (-keepheader -fasta -gappyout; (Capella-Gutiérrez *et al.*, 2009)) and paralogs pruned using phylotreepruner v 1.0 (min_number_of_taxa =18, bootstrap_cutoff= 0.7, longest sequence for a given orthogroup=u; (Kocot *et al.*, 2013)). Alignments shorter than 100 amino acids were removed, leaving 1,121 orthogroups.

For the maximum likelihood analysis, the trimmed and pruned alignments were concatenated into a supermatrix using FasConCat v1.04 (Kück and Meusemann, 2010) composed of 377,532 amino acids and the best substitution models determined using PartitionFinder v 2.1.1 (Lanfear *et al.*, 2016). The maximum likelihood consensus phylogeny from the supermatrix and

partition scheme was constructed using IQ-tree and 1,000 ultrafast bootstrap replicates (Nguyen *et al.*, 2015). For the coalescence species tree reconstruction, gene trees were generated using IQ-tree v 1.6.12 on the trimmed alignments of the 1,121 filtered orthogroups and processed using ASTRAL v 5.6.3 (Zhang *et al.*, 2018). Branch lengths are presented in coalescent units (differences in the 1,121 gene trees) and the node values reflect the local posterior probabilities.

RNA sequencing and transcriptome assembly

For RNA extraction, about 200 mites of all life stages were taken from stock culture and subsequently washed with 1% SDS for 10 s. RNA was extracted from living specimens using the Quick-RNA MiniPrep Kit (Zymo Research) according to the manufacturer's protocol. Quantity and quality of RNA were assessed using a Qubit fluorometer and NanoDrop One (Thermo Fisher Scientific), respectively.

Extracted RNA was shipped to Omega Bioservices (Norcross, GA, USA) on dry ice for library preparation and sequencing. Whole animal RNA was used for poly-A selection, cDNA synthesis and library preparation following the Illumina TruSeq mRNA Stranded Kit protocol. The library was sequenced with 100 million 150 bp paired-end on a HighSeq4000 platform. For the genome-guided assembly of the transcriptome a bam-file was created from the long-read genome using STAR (Dobin *et al.*, 2013). RNAseq reads were *in silico* normalized and subsequently used together with the bam-file to assemble the transcripts using Trinity v2.8.4 (Grabherr *et al.*, 2011; Haas *et al.*, 2013), yielding an assembly with a total length of 162.8 Mb, an N_{50} = 2994 bp and a BUSCO score (Simão *et al.*, 2015) of C:96.3% [S:36.5%,D:59.8%], F:1.3%, M:2.4%.

Life-stage specific RNAseq

For life-stage specific RNAseq, we collected 15 specimens per life stage from the stock culture that were split into three replicates of five individuals. Whole animals (for all stages but eggs) were flash frozen in 50 µl TRIzol using a mixture of dry ice and ethanol (100%) and stored at -80° . RNA was extracted using a combination of the TRIzol RNA isolation protocol (Life Technologies) and RNeasy Mini Kit (Qiagen) (Kitchen *et al.*, 2015). The TRIzol protocol was used for initial steps up to and including the chloroform extraction. Following tissue homogenization, an additional centrifugation step was performed at $12,000 \times g$ for 10 min to remove tissue debris. After the chloroform extraction, the aqueous layer was combined with an equal volume of ethanol and the RNeasy Mini Kit was used to perform washes following the manufacturer's protocol. Eggs were crushed using pipette tips and directly stored in a mixture of cell lysis buffer and murine RNase Inhibitor (New England Biolab).

We used the NEBNext® Single Cell/Low Input RNA Library Prep Kit for Illumina® together with NEBNext® Multiplex Oligos for Illumina® (New England Biolab) for library preparation, including reverse transcription of poly(A) RNA, amplification full-length cDNA, fragmentation, ligation and final library amplification according to the manufacturer's protocol. We performed cDNA amplification for 16 (18 for egg samples) PCR cycles and final library amplification 8 PCR cycles. In total, we constructed 18 libraries (three for each life stage). The quality and concentration of the resulting libraries were assessed using the Qubit High Sensitivity dsDNA kit (Thermo Scientific) and Agilent Bioanalyzer High Sensitivity DNA assay. Libraries were sequenced on an Illumina HiSeq2500 platform (single-end with read lengths of 50 bp) with ~18 million reads per library.

Illumina sequencing reads were pseudoaligned to the bulk transcriptome and quantified (100 bootstrap samples) with kallisto 0.46.0 (Bray *et al.*, 2016) using default options for single-end reads. Fragment length sizes were extracted from the Agilent Bioanalyzer runs. For life-stage specific differential expression analysis, kallisto quantified RNAseq data was processed with sleuth 0.30.0 (Pimentel *et al.*, 2017) using Likelihood Ratio tests in R 3.6.1 (RCoreTeam, 2019). The average transcripts per million (tpm) values for each target transcript were extracted from the sleuth object (see R script) and used with the Heatmapper tool (Babicki

et al., 2016) to produce an unclustered heatmap showing relative expression levels. UpSetR (Conway *et al.*, 2017) was used to compare the number of unique and shared expressed genes across life stages.

Identification of horizontal gene transfer events

To detect HGTs, we used the published tool “.Lateral_gene_transfer_predictor.py” (Thorpe *et al.*, 2018) to calculate the Alien Index described by (Gladyshev *et al.*, 2008) and (Flot *et al.*, 2013). All predicted genes were compared to the NCBI nr database as previously described (Thorpe *et al.*, 2018). Results to Arthropoda (tax id 6656) were ignored in the downstream calculations. The HGT candidates were filtered for contamination identified by both Blobtools (Laetsch and Blaxter, 2017) and the Alien Index (AI > 30 and >70% percent identity to a non-metazon sequence). The candidates were further filtered for > 50% overlap with predicted repeats using the bedtools *intersect* tool with the RepeatMasker gff file and expression from any developmental stage. Introns were scored manually from visualization in IGV genome browser (Robinson *et al.*, 2011) and GC content for all predicted genes was calculated using the bedtools *nuc* tool.

Analysis of chemosensory and photoreceptor gene families

The search and analysis chemosensory genes largely followed the procedure outlined by Dong *et al.* (Dong *et al.*, 2018) with slight modifications. First, the *Archezogozetes* official gene set (OGS) was searched using BLASTP (E-value, $<1 \times 10^{-3}$) against the following queries for the different chemosensory gene families. The OGS was queried against i) *D. melanogaster*, *D. mojavensis*, *Anopheles gambiae*, *Bombyx mori*, *T. castaneum*, *Apis mellifera*, *Pediculus humanus humanus*, and *Acyrtosiphon pisum* odorant binding proteins (OBPs) (Vieira and Rozas, 2011); ii) *D. melanogaster*, *D. mojavensis*, *A. gambiae*, *B. mori*, *T. castaneum*, *A. mellifera*, *P. humanus humanus*, *A. pisum*, *I. scapularis*, and *Daphnia pulex* small chemosensory proteins (CSP) (Niimura and Nei, 2005; Robertson and Wanner, 2006; Vieira and Rozas, 2011); iii) *D. melanogaster* and *A. mellifera* odorant receptors (Niimura and Nei, 2005; Robertson and Wanner, 2006); iv) *D. melanogaster*, *A. mellifera*, *I. scapularis*, *T. urticae*, *T. mercedesae*, and *M. occidentalis* gustatory receptors (GRs) (Robertson *et al.*, 2003; Robertson and Wanner, 2006; Gulia-Nuss *et al.*, 2016; Hoy *et al.*, 2016; Ngoc *et al.*, 2016; Dong *et al.*, 2017); v) a comprehensive list of iGluRs and IRs across vertebrates and invertebrates (Croset *et al.*, 2010), as well as those identified in the *T. mercedesae*, *D. tinctorium* and *L. deliense* genome projects (Dong *et al.*, 2017; Dong *et al.*, 2018). Second, all candidate *Archezogozetes* sequences were reciprocally blasted (BLASTP, E-value $<1 \times 10^{-3}$) against the NCBI database (Pruitt *et al.*, 2005) and all sequences that did not hit one of the respective receptors or transmembrane proteins were removed from the list. Third, for phylogenetic analysis of IRs and GRs from *Archezogozetes* were aligned with IRs from *D. melanogaster*, *T. urticae*, *D. tinctorium* and *L. deliense* and GRs from iv) *D. melanogaster*, *T. mercedesae*, *I. scapularis*, and *M. occidentalis*, respectively, using MAFFT (v 7.012b) with default settings (Katoh and Standley, 2013). Poorly aligned and variable terminal regions, as well as several internal regions of highly variable sequences were excluded from the phylogenetic analysis. Fourth, maximum likelihood trees were constructed with the IQ-TREE pipeline (v 1.6.12) with automated model selection using 1,000 ultrafast bootstrap runs (Nguyen *et al.*, 2015).

Reference opsin genes and opsin-like sequences were obtained from Dong *et al.* (Dong *et al.*, 2018) and used to query the *Archezogozetes* OGS using BLASTP (E-value, $<1 \times 10^{-5}$). Subsequently, candidates sequenced were reciprocally blasted against NCBI using the same settings and only retained if they hit an opsin or opsin-like gene. The *Archezogozetes* candidates were aligned with the query sequence list using MAFFT (v 7.012b) with default settings (Katoh and Standley, 2013). This opsin gene alignment phylogenetically analyzed using the IQ-TREE pipeline (v 1.6.12) with automated model selection and 1,000 ultrafast bootstrap runs (Nguyen *et al.*, 2015).

Gene family phylogenies

We used the following workflow to analyse genes related to Figure 5 (hox and developmental genes), Figure 7 (cell wall-degrading enzyme encoding genes) and Figure 8 (alcohol and geraniol dehydrogenases genes). Generally, protein orthologs were retrieved from NCBI (Pruitt *et al.*, 2005), and aligned using MUSCLE (Edgar, 2004) or MAFFT (v 7.012b) (Kato and Standley, 2013) and ends were manually inspected and trimmed. The resulting final protein sequence alignments used to construct a maximum likelihood (ML) phylogenetic tree with either i) PhyML with Smart Model Selection (Guindon *et al.*, 2010; Lefort *et al.*, 2017) or ii) the IQ-TREE pipeline with automated model selection (Nguyen *et al.*, 2015). The ML trees were constructed using either 1,000 ultrafast bootstrap runs (IQ-TREE) or approximate-likelihood ratio test (PhyML) was used to assess node support.

Feeding experiments with labelled precursors and chemical analysis (GC/MS)

Stable isotope incorporation experiments were carried out as previously described (Brückner *et al.*, 2020). Briefly, mites were fed with wheat grass containing a 10% (w/w) mixture of three antibiotics (amoxicillin, streptomycin and tetracycline) and additionally, we added 25% (w/w) of the stable isotope-labelled precursors [¹³C₆] D-glucose (Cambridge Isotope Laboratories, Inc.) as well as a control with untreated wheat grass. Cultures were maintained for one generation and glands of adult specimens were extracted one week after eclosion by submersing groups of 15 individuals in 50 µl hexane for 5 min, which is a well-established method to obtain oil gland compounds from mites (Raspotnig *et al.*, 2008; Brückner and Heethoff, 2016; 2017; Brückner *et al.*, 2017b).

Crude hexane extracts (2-5 µl) were analysed with a GCMS-QP2020 gas chromatography–mass spectrometry (GCMS) system from Shimadzu equipped with a ZB-5MS capillary column (0.25 mm x 30m, 0.25 µm film thickness) from Phenomenex. Helium was used as a carrier gas with a flow rate of 2.14 ml/min, with splitless injection and a temperature ramp was set to increase from 50 °C (5 min) to 210 °C at a rate of 6 °C/min, followed by 35 °C/min up to 320 °C (for 5 min). Electron ionization mass spectra were recorded at 70 eV and characteristic fragment ions were monitored in single ion mode. The temperatures of the ion source and transfer line were 230 °C and 320 °C, respectively.

Abbreviations

¹³C= stable isotope of carbon with the isotopic mass 13; ¹⁵N= stable isotope of nitrogen with the isotopic mass 15; abd-A = abdominal A; Abd-B= Abdominal-B; ADHs= alcohol dehydrogenases; ADU= adult; BLAST= basic local alignment search tool; bp= base pairs; BUSCO: C= complete genes, S= single copy genes, D= doublet genes, F= fragmented genes, M= missing genes; BUSCO= Benchmarking Universal Single-Copy Orthologs; Ch= chelicera; CRISPR/CAS9= clustered regularly interspaced short palindromic repeats/ CRISPR associated protein 9; CSP= chemosensory protein; dac= dachshund; DAPI= 4',6-diamidino-2-phenylindole; DEU= deutonymph; Dll= Distal-less; EGG= egg; EtOH= ethanol; exd= extradenticle; eya= eyes absent; FISH= fluorescence in situ hybridization; Gb= gigabase pairs, GeDH= geraniol dehydrogenase; GES= geraniol synthase; GFF= general feature format; GO= Gene Ontology; GPP= geranyl pyrophosphate; GR= gustatory receptor; HCN= hydrogen cyanide; HGT= horizontal gene transfer; Hi-C= all-*versus*-all chromosome conformation capture sequencing; hth= homothorax; iGluR= ionotropic glutamate receptors; IR= ionotropic receptor; kb= kilobase pairs, KEGG= Kyoto Encyclopedia of Genes and Genomes; KOALA= KEGG Orthology And Links Annotation; L1-3= walking legs 1-3; LAR= larva; LTR= Long Terminal Repeat; Mb = megabase pairs, MEP/DOXP pathway= 2-C-methyl-D-erythritol 4-phosphate/1-deoxy-D-xylulose 5-phosphate pathway; NCBI= National Center for Biotechnology Information; NMDA= N-methyl-D-aspartic acid; NMDS= non-metric

multidimension scaling; OBP= odorant binding protein; OD= optical density; OGS= official gene set; OR= odorant receptor; PCR= polymerase chain reaction; PFAM= Protein Families; Pp= pedipalp; PRO= protonymph; qPCR= quantitative polymerase chain reaction; ran= Roy A. Norton; RNAi= RNA interference; RNAseq= RNA sequencing; SEM= scanning electron microscopy; so= *six1/sine oculis*; SR μ CT= Synchrotron X-ray microtomography; TEM= transmission electron microscopy; TEs= transposable elements; tpm= transcripts per million; TRI= tritonymph; Ubx= Ultrabithorax

Declarations

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Ethics statement

There are no legal restrictions on working with mites.

Authors contributions

AB had the initial idea for the study; AB, AAB, PB and SAK designed research; IAA performed long-read and Hi-C sequencing and assembled the long-read genome; AB performed all other experimental work; AAB analyzed hox and life-stage specific expression data; AB analyzed chemical data; SAK and AB performed bioinformatic analyses; AB wrote the first draft of the manuscript with input from AAB, PB, and SAK; SAK revised the manuscript. All authors gave final approval for publication.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no conflict of interest.

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Data availability

Genomic and transcriptomic data generated for his project can be found on NCBI under the accession numbers PRJNA683935 and PRJNA683999. All other data that support the findings of this study have been deposited at <https://doi.org/10.22002/D1.1877> (Brückner, 2021). Mite specimens to start an own laboratory culture can be requested from the corresponding author.

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