Radiation of the Drosophila nannoptera species group in Mexico

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Abstract
The Drosophila nannoptera species group, a taxon of Mexican cactophilic flies, is an excellent model system to study the influence of abiotic and biotic factors on speciation, the genetic causes of ecological specialization and the evolution of unusual reproductive characters. However, the phylogenetic relationships in the nannoptera species group and its position within the virilis-repleta phylogeny have not been thoroughly investigated. Using a multilocus data set of gene coding regions of eight nuclear and three mitochondrial genes, we found that the four described nannoptera group species diverged rapidly, with very short internodes between divergence events. Phylogenetic analysis of repleta group lineages revealed that D. inca and D. canalinea are sister to all other repleta group species, whereas the annulima species D. aracataca and D. pseudotalamancana are sister to the nannoptera and bromeliae species groups. Our divergence time estimates suggest that the nannoptera species group radiated following important geological events in Central America. Our results indicate that a single evolutionary transition to asymmetric genitalia and to unusual sperm storage may have occurred during evolution of the nannoptera group.

Introduction
Species of the genus Drosophila, because of their well-defined phylogenetic relationships and diverse ecologies and life histories, provide an attractive group of model organisms for the study of evolution (Markow & O’Grady, 2007). A few taxa in Drosophilidae have evolved the ability to feed and breed in necrotic cactus, predominantly in the repleta and nannoptera species groups (Markow & O’Grady, 2007). The nannoptera species group consists of only four described species: Drosophila nannoptera (Wheeler, 1949), D. acanthoptera (Wheeler, 1949), D. wassermani (Pitnick & Heed, 1994) and D. pachea (Patterson & Wheeler, 1942). Even though all species of the nannoptera group live on columnar cacti, they exhibit diverse degrees of ecological specialization. Whereas D. nannoptera can live on a variety of host plants of the genera Stenocereus, Pachycereus, Escontria and Myrtillocactus (Heed, 1982), D. acanthoptera and D. wassermani are restricted to species in the genus Stenocereus (Heed, 1982). An even more tight ecological specialization links D. pachea to a single host plant, the senita cactus Lophocereus schottii (Engelmann, 1852), which is toxic to the other three species of the nannoptera group (Heed & Kircher, 1965; Etges et al., 1999) and which provides a particular sterol (lathosterol) absolutely required for D. pachea survival (Heed & Kircher, 1965; Lang et al., 2012).

The geographic distribution of D. pachea coincides with the distribution of senita cactus, which is restricted to the Sonoran desert in north-west mainland Mexico and to the Baja California peninsula (Fig. 1; Lindsay, 1963; Hastings et al., 1972). The Gulf of California and the Sierra Madre Occidental mountain range on the mainland separate the distribution of D. pachea from D. wassermani (Fig. 1b). The other nannoptera species are found in an overlapping region in southern Mexico (Heed, 1982; Markow & O’Grady, 2005). Drosophila nannoptera generally localizes in highlands, whereas D. wassermani is primarily found in lowlands (Heed,
Specimens of *D. acanthoptera* were also reported from Venezuela (Hunter, 1970), but this single sampling record, based on morphological characterization, remains dubious.

The nannoptera species group inhabits a zone of important geological history. About 15 million years ago (Ma), seismic activity along a volcanic arc formed the Isthmus of Panama that connected Central and South America (Montes et al., 2012a,b). The formation of the isthmus had a huge biological and climatic impact and provided a means for terrestrial fauna to move between the two continents (Webb, 1976; Leigh et al., 2014). More recently, about 6–3 Ma, the Baja California peninsula formed as a result of a series of complex geological events that caused the successive separation of landmasses from mainland Mexico (Lizarralde et al., 2007; Umhoefer, 2011). Increased desertification of North and South America in the past 10 Ma due to a global climate change and an uplift period of the Andes during the late Miocene-Pliocene (Gregory-Wodzicki, 2000; Capitanio et al., 2011) was accompanied by radiations of major succulent plant lineages (Arakaki et al., 2011). Whether these geological events might have influenced speciation within the nannoptera species group is unknown.

In addition to the ability to utilize cactus tissue as a resource, some unusual reproductive characters have evolved in the nannoptera species group. For instance, sperm gigantism was observed in *D. pachea* and *D. nannoptera*, but not in the other two members of the group, *D. acanthoptera* and *D. wassermani* (Pitnick et al., 1995). Two additional very curious reproductive characters, genital asymmetry (Vilela & Baechli, 1990; Pitnick & Heed, 1994; Lang & Orgogozo, 2012) and site of sperm storage in females (Pitnick et al., 1999), have also been reported in a few species of the nannoptera group. Whereas *D. nannoptera*, like most other Drosophilidae, has fully symmetric genitalia (Vilela & Baechli, 1990; Huber et al., 2007), the other three species of the nannoptera group possess diverse genital organs with conspicuous left-right asymmetric morphologies. *Drosophila pachea* displays an epandrial lobe size asymmetry (Pitnick & Heed, 1994; Lang & Orgogozo, 2012).

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**Fig. 1** Phylogenetic relationships of the nannoptera group and related species. (a) Phylogenetic tree generated in BEAST based on the concatenated data set with nine partitions. Bootstrap support from maximum likelihood (PhyML) analysis is presented on the left side of each node. Bayesian posterior probabilities are presented on the right side of each node for the BEAST analysis/and *BEAST* analysis, respectively. The time scale was calculated according to estimates B in Table 1. (b) Distributions of the species of the nannoptera group and of *Drosophila machalilla* and *D. bromelieae*, reproduced from Heed (1982) and Markow & O’Grady (2005).
D. acanthoptera possesses an asymmetric phallus (Vilela & Baechli, 1990), and D. wassermani has a left–right concave–convex-shaped cercus (Pitnick & Heed, 1994; Fig. 2). Furthermore, D. nannoptera, like most other Drosophila species, uses two types of organs for post-copulatory storage of sperm in females: the paired spermathecae and the single seminal receptacle. However, D. acanthoptera, D. wassermani and D. pachea use only the spermathecae (Pitnick et al., 1999). Therefore, with regard to the evolution of asymmetric genitalia and unusual sperm storage, the most parsimonious scenario would be that D. nannoptera is an out group relative to the other three species of the nannoptera group.

For the reasons mentioned previously, the nannoptera group thus represents an interesting model system to tackle a variety of important questions in evolutionary biology, such as the influence of abiotic and biotic factors on speciation, the genetic causes of ecological specialization and the evolution of reproductive characters. To address these questions and to trace back the evolution of different characters across the nannoptera species group, a reliable phylogeny of the four species and related taxa is required. Whereas relationships within the repleta group have been characterized to a great extent (Van der Linde et al., 2010; Oliveira et al., 2012), previous phylogenetic studies of the nannoptera group have led to equivocal and conflicting results (Pitnick et al., 1995, 1997, 1999; Van der Linde et al., 2010; Oliveira et al., 2012; Yassin, 2013). Some of these analyses were based on relatively few genetic loci (Pitnick et al., 1995, 1997, 1999) and others that included more loci either lacked appropriate out groups (Oliveira et al., 2012) or did not examine all members of the nannoptera species group (Van der Linde et al., 2010; Yassin, 2013). In a morphological analysis based on internal reproductive organ morphology, Heed (1982) proposed a phylogeny of the nannoptera group, with D. wassermani and D. pachea forming two sister species, which are in turn sister to D. acanthoptera, and with D. nannoptera being out group relative to the other

Fig. 2 Hypothetical character evolution of asymmetric male genitalia in the nannoptera species group. The red dot indicates the putative origin of both left–right asymmetric male genitalia and spermathecae-restricted sperm storage. Images below each species names illustrate male external genitalia of each species. Asymmetric parts were artificially coloured in red. Sperm storage organs are indicated below each species name. The scale bar is 100 μm.
three species. According to this topology, the evolution of asymmetric genitalia and the evolutionary change in sperm storage would have occurred only once, whereas sperm gigantism would have evolved twice within the nannoptera group.

The closest relatives of the nannoptera group are thought to be the bromeliae species group (Van der Linde et al., 2010; Yassin, 2013) and a newly described species, D. machalilla (Acurio et al., 2013; A. Acurio, K. Goodman, D.C. Oliveira, V. Rafael & A. Ruiz, unpublished) that was proposed to belong to a new species group, the altaia group (Acurio et al., 2013). Whereas the nannoptera and bromeliae species groups are part of the virilis-repleta group radiation (Throckmorton, 1975), the particular branching order of lineages leading to the nannoptera group has never been fully resolved (Van der Linde et al., 2010; Oliveira et al., 2012; Yassin, 2013).

Here, we address the phylogenetic relationships of the four species of the nannoptera group and related taxa. Using a multilocus data set of gene coding regions of eight nuclear and three mitochondrial genes, we found that the four species of the nannoptera group diverged rapidly. We discuss the nannoptera group radiation with respect to important geological events in Central America. Furthermore, our results allow us to propose a scenario for the evolution of reproductive traits in the nannoptera group.

Materials and methods

Taxon sampling

In addition to the four described species of the nannoptera group, we sequenced two species, D. bromeliae and D. machalilla, that are hypothesized to be close relatives of the nannoptera group based on morphology and previous phylogenetic analyses (Van der Linde et al., 2010; Oliveira et al., 2012; Yassin, 2013; A. Acurio, K. Goodman, D.C. Oliveira, V. Rafael & A. Ruiz, unpublished). We also included the annulimana group species D. aracataca and D. pseudotalamancana, and D. canalinea of the canalinea group to represent sister lineages of the repleta group. From the repleta group, we included four representative species, D. inca (inca subgroup), D. mettleri (mulleri subgroup), D. buzzatii (mulleri subgroup, buzzatii species complex) and D. mojavensis (mulleri subgroup, mojavensis species complex). Drosophila virilis and D. robusta were chosen as distant lineages of the repleta-virilis radiation (Van der Linde et al., 2010; Yassin, 2013) and D. funebris and D. grimshawi were used to root the phylogeny. The latter two species belong to different species radiations of the Drosophila subgenus (Throckmorton, 1975; Van der Linde et al., 2010; Yassin, 2013). Flies were obtained from the Drosophila Species Stock Center (Table S1), except for D. machalilla and D. inca (both collected by A. Acurio) and D. buzzatii (provided by Jean David).

DNA sequencing

Genomic DNA was obtained in a single extraction per species including 2–5 adults using the DNeasy blood and tissue kit (QIAGEN, Hilden, Germany). Partial genomic regions of eight nuclear genes (amy, amryel, boss, fkh, marf, sinA, snf, wee) and three mitochondrial genes (ND2, COI, COII) were amplified by PCR with gene-specific or degenerate primers (Liu & Beckenbach, 1992; Bonacum et al., 2001; Wang et al., 2006; Da Lage et al., 2007; Table S2). Degenerate oligonucleotides optionally contained T7 or SP6 universal primer sequences at their 5’ end (Table S2), following Bonacum et al. (2001). For PCR amplifications, we used 0.4 μM oligonucleotides, 1 μM GoTaq® DNA Polymerase (Promega, Fitchburg, WI, USA) per 35 μL reaction volume, 2 mM MgCl2 and 200 μM dNTP, and reactions were carried out using standard thermocycle conditions. PCR products were purified and Sanger-sequenced with gene-specific primers or with T7, SP6 universal primers at Cogenics (www.cogenics.com, Beckman Coulter, Brea, California, USA). Sequence data (GenBank accession numbers KF632591–KF632711; Table S3) were examined and aligned with Geneious 6.1.3 (Biomatters, Auckland, New Zealand). Additional sequence data were retrieved from GenBank (Table S3). We generated a data set that contained all the selected homologous genomic regions of all species. DNA sequences were aligned using MAFFT (Katoh et al., 2002). Nuclear loci were tested for recombination with the pairwise homoplasy index (Φw) statistics using PhilPack (Bruen et al., 2006). No evidence of recombination was detected (Table S4). A total of 121 polymorphic sites were detected within single sequences based on the presence of double peaks in sequencing chromatograms. Thirty-eight of these polymorphic sites were found in the nannoptera species group sequences. All polymorphic sites were excluded from the analysis. Furthermore, noncoding DNA sequences were removed from the data set, as well as a short region of the wee locus (sequences homologous to positions 267–306 in D. pachea wee, accession number KF632622), which was difficult to align and that did not contain any parsimony informative sites in the nannoptera species group. The extremities of each locus-specific alignment were also trimmed to be in codon frame. Alignments were realigned with the Geneious translation alignment program and either used separately or concatenated in the following order: amy-amryel-boss-fkh-marf-sinA-snf-wee-ND2-COI-COII. The number of informative sites was calculated using MEGA4 (Tamura et al., 2007).

Phylogenetic analysis

Phylogenetic analysis was performed using both maximum likelihood and Bayesian approaches. For all analyses, models of nucleotide substitution were selected using the Akaike Information Criterion as cal-

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calculated in jModelTest 2.1.3 (Posada, 2008; Darriba et al., 2012). Maximum likelihood inference was carried out on the concatenated data set in PhyML (Guindon & Gascuel, 2003) using a GTR+I+G model of nucleotide substitution. Node support was determined by performing 100 bootstrap replicates.

Two types of Bayesian analyses were carried out in BEAST v1.7.5 (Drummond et al., 2012). The first analysis was carried out on the concatenated data set using nine partitions with individual and unlinked models of nucleotide substitution (Table S5). The partitions corresponded to the eight nuclear loci (amy, amyl, boss, fhk, marf, sinA, snf and wee) plus a ninth partition for mitochondrial sequences (ND2, COI, COII). Mitochondrial genes were combined into one partition because they are located in the same order and orientation in the mitochondrial genome, and largely evolve as a single unit with little to no recombination (Ballard, 2000). Clock models were linked, and a common strict clock rate was assumed for all partitions using the Yule birth process tree prior. We also estimated a species tree using *BEAST (Heled & Drummond, 2010). For the *BEAST analysis, tree and clock models were unlinked for each partition and a relaxed exponential clock model was assumed.

For species divergence time estimates, we set priors for most recent common ancestors (MRCA) using estimates from Obbard et al. (2012) for the splits D. grimshawi – D. virilis: 13 ± 2.5 Ma and D. mojavensis – D. virilis: 10 ± 2.5 Ma (estimates A). Alternatively, calibration dates for the divergence of D. grimshawi – D. virilis: 42.9 Ma ± 8.7 (Tamura et al., 2004), D. mojavensis – D. virilis: 26 ± 6 Ma (Russo et al., 1995; Spicer & Bell, 2002; Oliveira et al., 2012), D. mojavensis – D. buzzatii: 11.3 ± 2 Ma and D. mojavensis – D. mettleri: 16.3 ± 2 Ma (Oliveira et al., 2012) were used (estimates B). Priors were assumed to follow a normal distribution with the mean and a standard deviation according to the literature estimates. Markov-Chain Monte-Carlo (MCMC) runs were performed with a chain length of 10^8 generations and were recorded every 1000 generations. Estimates were computed with Tracer version 1.5 (Rambaut & Drummond, 2009), and MCMC output analysis was carried out using TreeAnnotator (Drummond et al., 2012). The first 2000 sampled trees were discarded as the burn-in. Phylogenies were visualized and annotated with Figtree version 1.4 (Rambaut & Drummond, 2012).

Results

Phylogenetic analysis

To analyse the phylogenetic relationships of the nannoptera group, we gathered DNA sequences of partial coding gene regions of eight nuclear genes (amy, amyl, boss, fhk, marf, sinA, snf and wee) and three mitochondrial genes (ND2, COI, COII) from 17 species. The entire data set comprised 6810 aligned positions, including 4208 constant positions, 695 single variable positions and 1907 parsimony informative (28%) positions (Table S3).

Phylogenetic analysis of the concatenated, nine-partition data set was performed using a maximum likelihood approach in PhyML (Guindon & Gascuel, 2003) and by Bayesian inference in BEAST (Drummond et al., 2012). Phylogenetic relationships inferred from these analyses resulted in identical tree topologies, but with varying node support values (Fig. 1). The resulting phylogeny supports Acurio et al.’s findings (A. Acurio, K. Goodman, D.C. Oliveira, V. Rafael & A. Ruiz, unpublished) that the nannoptera species group is a sister clade of the atalaia species group, with D. machalilla being more closely related to the nannoptera clade than to D. bromeliae (bromeliae group). In our phylogeny, D. inca and D. canalinea form a lineage sister to the repleta group. Furthermore, both the Bayesian and the maximum likelihood phylogeny provided, for the first time, strong support for the monophyly of annulimana species, D. aracatara and D. pseudotalamancana, which we found to be more closely related to the nannoptera group than to the repleta group. Phylogenetic relationships within the nannoptera group were relatively well-resolved in the Bayesian analysis, but not in the maximum likelihood analysis. Our results from the Bayesian analysis were congruent with the phylogenetic relationships previously suggested based on morphological data (Heed, 1982) (Fig. 1a).

Analysis of concatenated multilocus data has recently been criticized as it poorly integrates locus-specific phylogenetic signals and can lead to false phylogenetic inferences with high statistical support (Song et al., 2012). Therefore, we also analysed our data set with *BEAST (Heled & Drummond, 2010), an extension of the BEAST package that incorporates coalescence models to estimate a species tree from multiple gene-specific phylogenies. The topology of the species tree inferred in *BEAST was similar to the phylogeny obtained with the concatenated data set (Fig. 1a, Fig. S1), except that relationships within the nannoptera group differed, with D. pachea being sister to the clade containing D. acanthoptera and the sister species pair D. nannoptera and D. wassermani (Fig. S1). However, the posterior probability for the corresponding nodes were low, suggesting that our data set might not contain enough information for species tree estimation using *BEAST. Within the nannoptera group, we observed only 120 parsimony informative sites across all genes in the data set (Table S3). The mean length of DNA sequence per nuclear locus was 477 ± 178 bp (SD), which, on average, included only 5 ± 3 (SD) parsimony informative sites among the nannoptera species group (Table S3). We wondered whether the number of informative sites per locus was too low for a
coalescent multilocus phylogenetic approach to produce a well-resolved species tree. To estimate the number of loci that would be necessary to reliably establish nanoptera group relationships using *BEAST, we produced partial data sets containing 2, 4, 6 or 8 loci and inferred the phylogeny of each data set. The average node support was calculated and the number of necessary loci was approximated by a logarithmic regression (Fig. S2). Whereas we are aware that the number of parsimony informative sites per locus is low and that the data sets are partially redundant, this analysis showed that for the entire phylogeny, the average posterior probability was 0.90 when six and eight loci were used. Support for the nodes within the nanoptera group also increased with the number of loci, but at a much lower rate. We estimated that approximately 60 loci would be required to obtain a posterior probability of 0.90 for the internal nodes within the nanoptera group.

**Divergence time estimates**

We estimated the divergence times of the nanoptera group radiation and the splits of *D. machalilla* and *D. bromelia* from the branch leading to the nanoptera species group. There is conflicting information about species divergence times in *Drosophila*. Most estimates are based on the phylogeny of Hawaiian Drosophilidae where species divergence times can be approximated based on the ages of the islands they inhabit (Price & Clague, 2002). Recently, Obbard et al. (2012) proposed a refinement of this approach to take lineage-specific variation of mutation rates into account (Obbard et al., 2012). This new approach suggested a younger age for the virilis-repleta radiation, of about 10 Ma compared to 20 Ma (see Material and methods). We computed species divergence times either based on Obbard et al. (2012) (dates A) or based on previous species divergence estimates (dates B) (Table 1).

As the calibration estimates in A were about half the ages in B, divergence time estimates of dates A were expectedly also half the age compared to estimates B.

**Table 1 Divergence time estimates.**

<table>
<thead>
<tr>
<th>Dated nodes</th>
<th>A</th>
<th>B</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>D. mojavensis</em> -- <em>D. buzzati</em></td>
<td>5.5 Ma (3.5–7.5)</td>
<td>12.2 Ma (10.0–14.5)</td>
</tr>
<tr>
<td><em>D. mojavensis</em> -- <em>D. mettleri</em></td>
<td>7.5 Ma (4.8–10.1)</td>
<td>16.7 Ma (13.7–19.6)</td>
</tr>
<tr>
<td><em>D. pachea</em> -- <em>D. wassermani</em></td>
<td>3.0 Ma (1.9–4.1)</td>
<td>6.7 Ma (5.3–8.1)</td>
</tr>
<tr>
<td><em>D. pachea</em> -- <em>D. nanoptera</em></td>
<td>3.7 Ma (2.4–5.0)</td>
<td>8.3 Ma (6.7–10.0)</td>
</tr>
<tr>
<td><em>D. pachea</em> -- <em>D. machalilla</em></td>
<td>7.5 Ma (4.9–10.2)</td>
<td>16.9 Ma (13.7–20.1)</td>
</tr>
<tr>
<td><em>D. pachea</em> -- <em>D. bromeliae</em></td>
<td>8.4 Ma (5.4–11.3)</td>
<td>17.9 Ma (15.8–20.9)</td>
</tr>
<tr>
<td><em>D. virilis</em> -- <em>D. mojavensis</em></td>
<td>10.6 Ma (7.0–14.3)</td>
<td>18.9 Ma (15.3–22.4)</td>
</tr>
</tbody>
</table>

*Estimates are the posterior means with 95% highest posterior density intervals.

We estimated that the nanoptera group lineage diverged about 3.7 Ma (dates A) or 8.3 Ma (dates B). Furthermore, the most recent split of *D. pachea* and *D. wassermani* was estimated to have occurred shortly thereafter, about 3.0 Ma (dates A) or 6.7 Ma (dates B). The *D. machalilla* lineage separated from the nanoptera group about 7.5 Ma (dates A) or 16.9 Ma (dates B) and the bromeliae group separated from the nanoptera group lineage about 8.4 Ma (dates A) or 17.9 Ma (dates B). Based on the conflicting calibration, these dates do not precisely estimate speciation events, but they put the nanoptera group radiation into an approximate time frame.

**Discussion**

**Origin of the nanoptera group**

The four species of the nanoptera group are endemic to distinct regions of Mexico (Fig. 1b). Our phylogenetic analysis uncovered three closely related out groups to the nanoptera group: *D. machalilla*, the *bromeliae* species group (comprising five species including *D. bromeliae*) and the members of the annulimana group. Members of these species groups are primarily found in South America but also in Central America (Fig. 1b) (Sturtevant, 1916; Duda, 1927; Pavan & da Cunha, 1947; Do Val & Marques, 1996; Da Silva et al., 2004; Markow & O’Grady, 2005; Acurio et al., 2013). These species distributions thus suggest that the ancestor of the four nanoptera group species may have originated from South America. Interestingly, our dating analysis estimates that the nanoptera group diverged from *D. machalilla* around 16.9 Ma (B) – 7.5 Ma (A). This time period corresponds to the closure of the Isthmus of Panama, about 15–9 Ma (Montes et al., 2012b), suggesting that the ancestor of the nanoptera group may have migrated over the newly formed isthmus from South America. Most species were found to migrate across the isthmus much later, at about 3–2 Ma (Leigh et al., 2014). However, exceptions are known such as the extinct carnivora *Cyonasua* and ground sloths, which migrated about 9 Ma from north to south and south to north, respectively (Webb, 1976). Furthermore, recent data suggest that the isthmus was already passable for stingless bees at late Eocene and early Miocene times (20–15 Ma), which migrated from South to Central America (Roubik & de Camargo, 2012). The isthmus might have faced multiple events of temporary land bridge formations and disconnections, allowing a few species to cross continents before a permanent land bridge formed about 4–3 Ma (Webb, 1976; Roubik & de Camargo, 2012; Stone, 2013).

*Drosophila* species have been extensively sampled in Mexico and Central America (Patterson & Stone, 1952), but multiple areas known as biodiversity hotspots in South America are still unexplored. An origin of the
nannoptera group in South America would suggest that yet undescribed close out group species of the nannoptera group might be present in these geographic areas.

**Radiation of the nannoptera species group**

Within the nannoptera species group, we observed only 120 parsimony informative sites in our entire data set (Table S6), whereas 850 nucleotide changes were lineage-specific. The distribution of mutations indicates that the four species of the nannoptera group diverged within a relatively short time period. Phylogenetic inference can be particularly difficult in these cases especially if node support remains low with increasing data. One interpretation of the current data is that the ancestral lineage diverged nearly simultaneously into the four described extant lineages (Walsh et al., 1999; Humphries & Winker, 2010). Such scenario might be expected from species with a large geographic range where peripatric speciation can occur in different regions. One well-studied example is the *D. simulans* species complex, where *D. mauritiana* and *D. sechellia* diverged independently on islands that were geographically separated from the cosmopolitan species *D. simulans* (Garrigan et al., 2012).

Alternatively, very short internode distances could result from a rapid succession of divergence events that could be inferred with increasing amounts of data (soft polytomy). Our current sequence data are insufficient to distinguish between a soft and a hard polytomy in the nannoptera species group. The rapid and ongoing decrease in high-throughput sequencing costs now makes it more practical to sequence and to compare whole genomes for future studies aiming at a better resolution of the nannoptera group phylogeny.

**Phylogenetic relationships and evolution of reproductive traits within the nannoptera group**

Under the hypothesis of a soft polytomy in the nannoptera species group, the phylogeny of the nannoptera group that we inferred using the concatenated data (Fig. 1a) appears to propose the most plausible scenario, despite a low node support in one of our analyses (PhyML maximum likelihood analysis). Indeed, several lines of evidence corroborate this topology. First, compared to the previous molecular phylogenetic analysis of Oliveira et al. (2012), which hypothesized different relationships for the nannoptera group, our phylogeny is based on a higher number of loci and on an increased number of relevant out group species close to the nannoptera group.

Second, our inferred topology recapitulates the species relationships presented by Heed (1982) based on internal reproductive organ anatomy and by Pitnick et al. (1999) independently based on cytochrome oxidase data. Third, it is congruent with chromosome inversions. Comparisons of polytene chromosome banding patterns revealed that *D. nannoptera* and *D. wassermani* have a homosequential ‘ancestral-like’ chromosomal organization, whereas *D. achatinoptera* and *D. pachea* are derived with three and one inversion, respectively (Ward & Heed, 1970). A fourth, polymorphic, inversion is also found in *D. pachea* and is not detected in the other species of the nannoptera group (Etges et al., 1999).

Fourth, our inferred topology is consistent with a parsimonious scenario of the evolution of the unusual reproductive characters within the nannoptera group. Genital asymmetry is found in *D. achatinoptera* (Vilela & Baechli, 1990), *D. wassermani* (Pitnick & Heed, 1994) and *D. pachea* (Pitnick & Heed, 1994; Lang & Orgogozo, 2012) (Fig. 2), whereas *D. nannoptera* (Vilela & Baechli, 1990), as well as the species *D. bromeliae*, *D. speciosa* and *D. aigupe* of the bromelaiid group and *D. machalilla* (ataiala group) have symmetric genitalia (Do Val & Marques, 1996; Da Silva et al., 2004; Acurio et al., 2013). Therefore, a single evolutionary transition to asymmetric genitalia might have occurred in the nannoptera group. Even though the asymmetry involves different male genitalia organs in each species, a common genetic and developmental process may underlie these distinct morphological asymmetries. We currently are trying to unravel the genetic factors that determine the asymmetric development of male genitalia in the three nannoptera species. In particular, we are testing whether genitalia clockwise rotation (Feuerborn, 1922; Suzanne et al., 2010) during pupal development could be the signal that triggers differential growth between the left and right parts of various organs in distinct species. Furthermore, our inferred topology is consistent with a single evolutionary change in sperm storage in the nannoptera group. After copulation, females of *D. achatinoptera*, *D. wassermani* and *D. pachea* exceptionally use only the spermathecae to store the sperm and not the seminal receptacle as is typical for Drosophilidae, including *D. nannoptera* (Pitnick et al., 1999).

Future efforts are required to examine how copulation position might affect sperm transfer in the nannoptera species group and to determine whether asymmetric male genitalia and unusual sperm storage are functionally linked. Finally, according to our inferred phylogeny, sperm gigantism would have evolved twice independently in the *nannoptera* group, which is consistent with other reported instances of rapid evolution of sperm size in Drosophila (Pitnick et al., 1995).

**Evolutionary history of the nannoptera species group**

Species divergence estimates for the virilis-repleta radiation vary greatly, from 30–20 Ma (Russo et al., 1995; Spicer & Bell, 2002; Tamura et al., 2004; Oliveira et al., 2012) to 10 Ma (Obbard et al., 2012) when adjusting
for lineage-specific mutation rates. We performed two separate estimations (Table 1) to account for two incompatible calibrations of species divergence estimates. The two sets of estimates (A and B) for the radiation of the nannoptera group approximate the lower and upper bounds of the geological time approximation of sea floor spreading of the southern Gulf of California. Formation of the Baja California peninsula started approximately 12 Ma due to changes in continental plate tectonics (Umhoefer, 2011). The peninsula itself formed along an almost north-south-directed rift, now partially covered by the Gulf of California. Landmasses separated from the continent as a result of complex geological events about 6–2.5 Ma and successively formed the peninsula (Lizarralde et al., 2007; Umhoefer, 2011). Thus, the divergence of D. wassermani and D. pachea might have been influenced by the formation of the Baja California peninsula and by the separation of these landmasses from the continent (Heed, 1982). Whether senita cacti were already present in the forming Baja peninsula and whether D. pachea or its predecessors were already feeding on senita cactus when landmasses disconnected from the continent is unknown. A phylogenetic analysis of the senita cactus and its closely related species, together with estimations of divergence times, would be helpful to try to infer the evolutionary history of the close ecological relationship between D. pachea and its host cactus. The distribution area of D. wassermani is limited to the north by the Trans-Mexican Volcanic Belt and by the Sierra Madre Occidental mountain range (Fig. S3), which originated about 17–7 and 38–25 Ma, respectively (Ferrari et al., 1999). A plausible scenario is that these mountains formed an obstacle for the ancestor of D. wassermani and D. pachea, which colonized further northern regions through the coastal lowlands of north-west Mexico (Heed, 1982). As this region successively re-arranged into the Baja California peninsula, the Gulf of California created a natural barrier and could have led to the isolation of D. pachea in the north and D. wassermani in the south (Fig. S3).

Drosophila machalilla, the most closely related outgroup of the nannoptera group, is a recently described species that was collected in traps containing Opuntia cactus, and the columnnar cactus Armatocereus cartwrightianus (Britton & Rose, 1920) was proposed to be their native host plant (Acurio et al., 2013). As all nannoptera group species also feed on columnar cacti, the MRCA of D. machalilla and the nannoptera species group was likely to be already cactophilic. Our results suggest that the major radiation of succulent plants, which occurred in the past 10 million years in North and South America (Arakaki et al., 2011), could have then contributed to shifts in cactus hosts and to speciation in the nannoptera group.

In summary, our results indicate that the four species of the nannoptera group originated within a short time period. Our approximations of species divergence times suggest that the emergence of the southern Gulf of California might have been involved in the split between D. pachea and D. wassermani. The branching order of basal repleta lineages reveals that the annulimana species D. aracataca and D. pseudechominae are the most closely related taxa to the nannoptera and bromeliae species groups. Our phylogenetic analysis suggests that evolution of asymmetric genital and unusual sperm storage have evolved only once within the nannoptera group, and that the ancestor of the nannoptera group was already feeding on columnar cacti.

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References


Supporting information

Additional Supporting Information may be found in the online version of this article:
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Table S2 Oligonucleotides used in this study.
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Figure S1 *BEAST analysis and gene-specific tree topologies.
Figure S2 Approximation of the number of loci required for a *BEAST multilocus analysis of the nemoptera group with reliable node support.
Figure S3 Hypothetical speciation scenario of Drosophila wassermani and D. pachea.

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