Quantifying the extent of morphological homoplasy: A phylogenetic analysis of 490 characters in *Drosophila*

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Homoplasy is a fundamental phenomenon in evolutionary biology but an appraisal of its extent at the morphological level is still lacking. Here, we analyzed the evolution of 490 morphological characters conceptualized among 56 drosophilid species. We found that two thirds of morphological changes were homoplastic and that the level of homoplasy depended on the stage of development and the type of the organ, with the adult terminalia being the least homoplastic. In spite of its predominance at the character change level, homoplasy accounts for only ~13% of between species similarities in pairwise comparisons. These results provide empirical insights on the limits of morphological changes and the frequency of recurrent evolution.

**KEY WORDS:** Character conceptualization, character state coding, developmental hourglass, genitalia, maximum parsimony, recurrent evolution.

**Impact Summary**

Is morphological evolution limited, with life being constrained to a small area in the space of all possible forms? Or is it endless, as has been postulated by Darwin 150 years ago? The repeated origin of similar traits in phylogenetically distant lineages, known as homoplasy, challenges Darwin’s view but we still lack empirical appreciation of the extent of this phenomenon. Analysis of the evolution of a large set of morphological traits in different developmental stages among 56 fly species revealed that two thirds of morphological changes were homoplastic. Homoplasy was also more frequent in juvenile stages than in adults who showed the highest morphological diversity. Although these findings support the prevalence of homoplasy, they also show that opportunities for the origin of new forms are still higher than it has recently been suggested.

“Although new and important modifications may not arise from reversion and analogous variation, such modifications will add to the beautiful and harmonious diversity of nature.”

(Darwin 1859)

Homoplasy, that is, the independent origin of similar character states between distant taxa, is widespread in the living world (Wake et al. 2011), but an appraisal of its extent and underlying mechanisms remains lacking. Some authors argue that homoplasy is so ubiquitous that it is evidence for the limitation and predictability of evolution (Conway Morris 2008; McGhee 2011; Blount et al. 2018). Others, however, caution against this view due to possible experimental and character selection biases (Powell and Mariscal 2015; Stayton 2015). Indeed, whereas long lists of homoplastic examples have been compiled (e.g., Martin and Orgogozo 2013), equivalent lists of nonhomoplastic, that is, apomorphic, states have rarely been made. An account of the diversity of as many character states as possible in clades with well-defined phylogenies is therefore strongly needed.

Characters are qualities attributed to delimited structures. For molecular data, the structures are nucleotides or amino acid sequences.
acid residues at a well-defined spatial position in a sequence, whereas the qualities are the biochemical compositions of these nucleotides and amino acids. Categorical values with no intermediates could be attributed to these qualities, that is, 4 for nucleotides and 20 for amino acids. For morphological traits, on the other hand, such character conceptualization and categorical coding are difficult (Vogt et al. 2010). In addition, morphological traits are usually stored in the context of phylogenetic analyses wherein autapomorphic states, that is, unique, novel states existing in only a single taxon or species, are often intentionally omitted when parsimony, the predominant approach, is used for tree reconstruction (Bryant 1995; Lewis 2001). This omission could lead to strong biases in estimating homoplasy since homoplasy is time-independent, whereas synapomorphies, that is, shared, commonly inherited derived states, reflect a particular case of evolutionary stasis wherein a state is maintained with minimal changes for long periods of time.

Here, we address the question of morphological homoplasy in Drosophila. One hundred years of genetics and developmental biology research have made this fly one of the best-understood animals at the morphological level. Annotated genome sequences for multiple species are available and links between genetic mutations and morphological aberrations are curated in accessible online databases such as Flybase (www.flybase.org). Besides the availability of unique genetic toolkits, these resources have made the genus ideal for studies aiming to trace morphological homoplasy between species to their molecular underpinnings (e.g., Wittkopp et al. 2003b; Prud’homme et al. 2006; Kagesawa et al. 2008; Tanaka et al. 2009; Frankel et al. 2012; Signor et al. 2016; Yassin et al. 2016a, b). From two major taxonomic references, we conceptualized 490 morphological characters among 56 drosophilid species. By analyzing the evolution of these traits on a molecularly inferred phylogeny, we were able to quantify the extent of morphological homoplasy in this important clade.

Materials and Methods

TAXON SAMPLING

We selected 56 drosophilid species for which molecular sequences were available in GenBank and full morphological descriptions could be obtained from two major taxonomic books, Okada’s (1968) Systematic Study of the Early Stages of Drosophilidae and Bächli et al.’s (2004) The Drosophilidae (Diptera) of Fennoscandia and Denmark (Table S1). These books represent two of the most explicit standardized and illustrated descriptions for drosophilids juveniles and adults. We selected species from the main drosophilid clades, that is, the subfamily Steganininae with its two tribes the Steganini and the Gitonini and the subfamily Drosophilinae with its two tribes the Colocasiomyini and the Drosophilini, following Yassin’s (2013) classification scheme (see O’Grady and DeSalle 2018 for the current status of drosophilids phylogenetics). The 56 species belonged to 15 genera, including four Drosophila subgenera (namely Sophophora, Dorsilopha, Drosophilus, and Siphlopla). More than one member represented some species groups, e.g., the melanogaster, obscura, and quinaria groups, hence encompassing shallow and profound phylogenetic depths. Some species were treated as “name holder,” having composite DNA sequences or grouping juvenile data from Okada (1968) and adult data from Bächli et al. (2004), such as Leucophera sp, from knowledge or presumption of monophyletic relationships (Table S1; Fig. 1). For morphological data, only 21 species had data from both taxonomic monographs (Table S1; Fig. 1).

MOLECULAR PHYLOGENETIC TREE

In order to define the phylogenetic relationships of species that were not included in Yassin’s (2013) family-wide phylogenetic revision, we obtained molecular sequences from GenBank for the 56 taxa for one mitochondrial (COI) and four nuclear genes (28S rRNA, Adh, Amyrel, and Gpdh; Table S1). For each gene, DNA sequences were aligned using the Muscle program (Edgar 2004) with default parameters as implemented by the MEGA7 software package (Kumar et al. 2016). The alignments were then concatenated in a single nexus file (Supporting Information S1). MEGA7 was also used to infer the best DNA substitution model for each gene. For the five genes, the GTR+G+I model had the lowest Akaike Information Criterion (AIC) value and thus was chosen for phylogenetic inference.

Phylogenetic analysis was conducted using MrBayes ver. 3.2. (Ronquist et al. 2012). We used Yassin’s (2013) phylogenetic classification as a topological constraint prior given its larger taxonomic and gene sampling. Because the GTR+G+I model was suggested for all genes, we tested unpartitioned model versus partitioned model strategies using stepping-stone sampling (Xie et al. 2011) implemented in MrBayes. For each strategy, we conducted two simultaneous runs of 1,000,000 generations with sampling every 100 generations, and considered a value of average SD of ≤0.01 an appropriate indicator for convergence between the two runs. Unpartitioned model had a higher marginal likelihood value than partitioned model (–42,712 vs. –42,804). We also used stepping-stone sampling to test for various clock models using the same run conditions of the unpartitioned data. Relaxed clock models had better marginal likelihood values (TK02 = –42,657 and IGR = –42,658) than the strict (–42,684) and no clock (–42,712). Consequently, we chose the unpartitioned, relaxed clock under the TK02 model. For the trees to be used in subsequent homoplasy analyses in this paper, we performed two simultaneous runs of 2,000,000 generations with sampling every 100 generations for the unpartitioned matrix under the TK02 relaxed clock model (Supporting Information S1). However, we
Figure 1. Bayesian consensus phylogenetic relationships of the 56 sampled drosophilid species (left) and the extent of coded characters (black) and missing data (white) (right). Values above nodes refer to posterior probability estimates (in percent). Asterisks (*) indicate clades that have been a priori constrained (see text).

stopped the runs after 1,004,000 generations since an average SD of 0.006 was attained. We then estimated the consensus tree after a burning period of 25% of the 20,086 sampled trees (Fig. 1). In subsequent analyses, we rerooted the consensus and sampled trees by considering the subfamily Steganinae a sister outgroup to the remaining species following the general consensus in drosophilid systematics (Yassin 2013; O’Grady and DeSalle 2018).

MORPHOLOGICAL CHARACTER CONCEPTUALIZATION

The taxonomic description of specimens consists of a laconic style wherein the name of an anatomical structure is followed by a quality. The same quality (e.g., color) could be attributed to multiple structures (e.g., head, legs, etc.). The same structure (e.g., the aedeagus) could hold multiple qualities (e.g., size, shape, texture, etc.). The same structure bearing the same quality but at different developmental stage (e.g., the number of teeth of the mouth hook of the cephalopharyngeal skeleton in the three larval instars) is conceptualized as a separate character for each stage. Subtle differences in the same quality could lead to the conceptualization of multiple characters. For example, “pleura yellowish” or “pleura with three dark stripes” stem from the same quality, that is, color, but each describes a different character, that is, pigmentation and color pattern, respectively.

MORPHOLOGICAL CHARACTER STATE CODING

Coding refers to how different values of the same quality of a character in each species could be categorized so that diversity between species and state transformation during evolution could be inferred. We opted for discrete coding, because of its long tradition in the phylogenetic literature and its ability to summarize different types of descriptions such as binary, verbal, and numerical (Supporting Information S2 and S3). Below is how we proceeded for coding the different types of traits:
Figure 2. Coding numerical and color traits. (A–D) An example for coding wing length. (A) Definition of the wing length character. (B) Histogram showing the distribution of wing length among the sampled taxa. (C) Hierarchical cluster analysis of wing length values. (D) NBClust estimation of 3 as the optimal number of clusters, delimited with dotted lines in (C). (E) Each of the 60 colors present in the literature was translated into its RGB values. (F) PCA of RGB values followed by cluster analyses on the first two PC axes identified six clusters or states (delimited with dotted lines).
(1) Numerical descriptions: Numerical values such as lengths, widths, counts (e.g., bristles), and indices were directly obtained for each taxon. When the range over multiple specimens was recorded, we used the average between the extremes as the summarizing value. We then used the NBClust package in R (Charrad et al. 2014) to estimate the optimal number of clusters for each character using Euclidean distance and Ward’s method. Values belonging to the same cluster were then attributed the same code in the data matrix. Figure 2A–D shows an example for the coding of a single numerical character (wing length).

(2) Verbal descriptions: Qualities such as “large,” “fusiform,” “concave,” “divergent,” and so on were directly coded when only few distinct values were given (e.g., “concave” vs. “convex”). When illustrations were present, we used ImageJ package (Abramoff et al. 2004) to estimate lengths, angles, and areas for characters for which qualities were slightly ambiguous (e.g., “elliptical” vs. “ovoid,” “subequal” vs. “slightly equal,” etc.). The numerical values obtained from ImageJ were then analyzed using NBClust in R to define the number of optimal states.

(3) Color descriptions: We found 60 verbal descriptions of colors in the two books such as “yellow,” “slightly grayish,” “brownish black,” “reddish brown,” and so on. We translated these descriptions into corresponding RGB values (Table S2, Fig. 2E). Since color perception is highly subjective, each of us produced an independent RGB table for the same descriptions. We then asked a colleague with an experience in Drosophila pigmentation to select for the most likely values in cases of disagreement. We conducted a Principal Component Analysis (PCA) on the RGB values, and projection of the two first axes identified a triangular distribution (Fig. 2F) corresponding to the three main Drosophila pigments, that is, black and brown melanin, yellow NADA sclerotin, and white NBCAD sclerotin (Wittkopp et al. 2003a). We analyzed each of these axes using NBClust and identified six states for colors that were used for all color characters.

(4) Pattern descriptions: Patterns describe the spatial distribution of a quality within a structure. We encountered several pattern descriptions such as trichomes on the larval abdomen, tracheae of the anterior spiracles of the pupa, tubercules on the posterior spiracles of the puparia, pilosity on the genitalia, pigmentation of adult tergites, and so on. For larval trichomes, we followed Okada’s (1968) categories. For pupal tracheae and tubercules, we followed a reductive coding, wherein each structure was reduced to its components and qualities of each component coded separately (e.g., tracheae were separated into basals, pseudobasals, centrals, pseudocentrals, and marginal). For pigmentation patterns on adult tergites, a trait that has intensively been studied in Drosophila evo-devo research (Massey and Wittkopp 2016), we divided each tergite into distinct spatial regions (Figs. S1–S7) and coded the colors as described above.

Missing data corresponded to either values that were not given for all taxa or those that are attributed to structures not present in all taxa. For example, steganine eggs lack filaments (Character 7). They were then attributed a missing data sign (‘?’) in all characters describing filaments number and shape in other species whose eggs have filaments (Characters 8–10).

The biological significance of our statistical delimitation remains to be clarified since the genetic basis of only a few morphological changes in drosophilids have yet been determined in evo-devo research (Table 1). From this literature, we found that our approach has led to “state lumping”, that is, coding genetically different states into similar categories, for the number of hypandrial bristles (Character 394). This character was statistically coded as having three states with state 0 bearing 0 to 3 pairs. However, the transition from 1 pair to 0 bristles between a pair of Drosophila species has a strong genetic component (Nagy et al. 2018). Consequently, we used the biological information to recode 0 bristles as a fourth state for this character.

QUANTIFYING ENSEMBLE HOMOPLASY
Several comparative phylogenetic measurements and tests for homoplasy have recently been developed (Speed and Arbuckle 2017). However, for discrete data, traditional cladistics measurements such as the ensemble consistency (CI) and homoplasy (HI) indices (Kluge and Farris 1969) and the retention indices (RI, Farris 1989) remain the only developed statistics (Speed and Arbuckle 2017). We thus used PAUP* version 4.0a (Swofford 2002) to estimate these indices for the whole data set, using the Describe Tree command under both the ACCTRAN and DELTRAN models and all characters were unordered. To account for topological uncertainties, the indices averaged over the 20,086 sampled trees from the two simultaneous MrBayes runs (Supporting Information S3). Given the strong topological constraints we applied (see above), we averaged indices over all sampled trees and not only on those retained after burnin.

COUNTING HOMOPLASTIC STATES
We used the PAUP* to produce the list of character changes (DescribeTrees/chgList) as well as to infer ancestral states (DescribeTrees/internal) on the consensus Bayesian tree under the ACCTRAN and DELTRAN models. We then parsed the output file to count for each character state the number of times it has been derived and shared between taxa. This led to five possible categories: (1) nonhomoplastic root states, (2) homoplastic root states (i.e., root states that have been secondarily derived), (3)
Table 1. Examples of morphological characters studied in Drosophila evo-devo studies.

| No. | Character                                      | References                                      | This study |
|-----|-----------------------------------------------|-------------------------------------------------|------------|
| 1   | Egg filaments                                 | (Kagesawa et al. 2008; Osterfield et al. 2015)  | Yes        |
| 2   | Egg dorsal ridge                              | (Niepielko and Yakoby 2014)                     | Yes        |
| 3   | Larval hooklets                               | (Frankel et al. 2012)                           | Yes        |
| 4   | Puparium color                                | (Ahmed-Braimah and Sweigart 2015)               | Yes        |
| 5   | Head shape                                    | (Arif et al. 2013a)                             | Yes        |
| 6   | Dorsocentral bristles                         | (Marcellini and Simpson 2006)                  | Yes        |
| 7   | Protarsal bristles (sex combs, brushes)       | (Tanaka et al. 2009; Atallah et al. 2014b; Rice et al. 2018) | Yes |
| 8   | Mesofemoral bristles                          | (Arif et al. 2013b)                             | NO         |
| 9   | Wing color patterns                           | (Prud’homme et al. 2006; Werner et al. 2012; Arnoult et al. 2013) | Yes |
| 10  | Abdomen tergal pigmentation                   | (Wittkopp et al. 2003b; Jeong et al. 2008; Rebeiz et al. 2009; Signor et al. 2016; Yassin et al. 2016a,b; Grover et al. 2018) | Yes |
| 11  | Male abdominal muscles                        | (Orgogozo et al. 2007)                          | NO         |
| 12  | Epandrial posterior lobe                      | (Glassford et al. 2015; Tanaka et al. 2015)    | Yes        |
| 13  | Hypandrial bristles                           | (Nagy et al. 2018)                              | Yes        |
| 14  | Aedeagal basal shape                          | (Peluffo et al. 2015)                           | Yes        |
| 15  | Ovariole number                               | (Orgogozo et al. 2006; Green and Extavour 2012) | No         |
| 16  | Oviscapt shape                                | (Atallah et al. 2014a)                          | Yes        |

homoplastic derived states, (4) synapomorphic states (i.e., states that have been derived once and are shared by at least two terminal taxa), and (5) autapomorphic states (i.e., states that have been derived once and are specific to a single terminal taxon).

TESTING THE EFFECTS OF DEVELOPMENTAL STAGES AND BODY ORGANIZATION

A character consistency index \( c_i \) of 1 indicates that all states are apomorphic, that is, they have been derived once. While a \( c_i < 1 \) is a sign of homoplasy in the character, \( c_i < 1 \) values cannot be compared between characters because the lower boundary of \( c_i \) depends on the number of character states (e.g., being 0.5 for a binary character). The \( r_i \) was proposed to correct for this limitation. A character retention index \( r_i \) value of 1 also indicates apomorphy, but unlike the \( c_i \), \( r_i \) cannot be calculated for autapomorphic traits, that is, characters with derived states appearing in only one terminal taxon. Consequently, an \( r_i \) of 1 indicates only synapomorphies, that is, commonly inherited derived states among a group of taxa. For each character, we estimated both \( c_i \) and \( r_i \) using the PAUP* DescribeTrees/diagnose command on the 20,086 sampled Bayesian trees. Averaged values for each character were then used to test for differences in the amount of homoplasy among developmental stages and body parts. For this, characters were classified according to six developmental stages: egg, larva 1, larva 2, larva 3, pupa, and adult, with the adult characters being further classified into five body parts: head, thorax, abdomen, male terminalia, and female terminalia. We also tested the effect of autapomorphies by conducting statistical tests on \( r_i \) without autapomorphies or after attributing an \( r_i \) value of 1 to autapomorphic characters (hereafter denoted \( r_{ic} \)). We conducted all statistical analyses and model testing using R (www.R-project.org).

DISTINGUISHING BETWEEN HOMOPLASTIC MECHANISMS

Traditionally, homoplasy is thought to result from three major mechanisms: parallelism, convergence, and reversal. There are confusions on the distinction between parallelism and convergence, with the former meaning similar initial conditions of the homoplastic state between the compared taxa and the latter referring to distinct origins (reviewed in Pearce 2012 and Stayton 2015). Some authors consider the “origin” in developmental terms, for example, changes in similar or different genes for parallelism or convergence, respectively. Others consider the “origin” as the ancestral placement on the morphospace regardless of the underlying developmental mechanism as proposed by Jablonski (cited in Pearce 2012). We follow here Jablonski’s definition for its operationality in phylogenetic analyses.
We distinguished six possible changes for character spaces in a pair of species depending on the degree of sameness, similarity, and difference of the ancestral and present states in the two species (Fig. 3). We then used PAUP* to infer under the ACCTRAN and DELTRAN models the ancestral states at each internal node. Using a customized perl script we obtained for each taxon for each trait its current state, its original state, and the internal node at which the state changed (Supporting Information S4). For each trait, we conducted all possible pairwise comparisons between species with no missing data and estimated the frequency of each change category (Supporting Information S4).

Unlike parallelism and convergence, reversal does not imply comparisons between taxa, but a comparison between the state present in a taxon with the states present in its ancestors, and consequently this may lead to any of the possible between-species mechanisms shown in Figure 3. Using a perl script (Supporting Information S5), we compared for each trait the current state of each taxon with that of its ancestors and counted incidences were a reversal (e.g., $0 \rightarrow 1 \rightarrow 0$) was found. Because the path of each taxon to the root had to be manually entered in the script from PAUP* node labeled cladogram, we conducted these analyses on a single tree, that is, the consensus Bayesian phylogeny.

**Results**

**AT LEAST 487 MORPHOLOGICAL CHARACTERS DIFFER BETWEEN DROSOPHILID SPECIES**

Thorough reading of the descriptions of the 56 species led to the conceptualization of 490 characters from drosophilids eggs ($N = 12$), larvae1 ($N = 13$), larvae 2 ($N = 23$), larvae 3 ($N = 26$), pupae ($N = 46$), and adults ($N = 370$). Adult characters were also classified into traits on the head ($N = 78$), thorax ($N = 90$), abdomen ($N = 40$), male terminalia ($N = 150$), and female terminalia ($N = 12$). The total number of characters states was 1479, ranging from 1 to 8 with an average of three states per character. Three characters were invariable having a single state, that is, egg color (always white or greyish white), the number of ventral caudal tubercles in the pupa (always 2), and the presence of an apical bristle on the mesotibia (always present).

**TWO THIRDS OF MORPHOLOGICAL CHANGES ARE HOMOPLASTIC**

Under both the ACCTRAN and DELTRAN models, consistency index (CI) for the whole data set averaged over all 20,086 trees sampled from the two runs of MrBayes was $0.33 (\pm 4.37 \times 10^{-6})$, with its complement, the data homoplasy index (HI), being $0.67 (\pm 4.37 \times 10^{-6};$ Fig. 4A). This means that for a character change in *Drosophila* there is nearly as much as twice a chance that the character will take a recurrent or a preexistent state than a new one. Interestingly, average HI value after 10,000,000 random permutations of taxa over the tree using PAUP* was 0.75, with the empirical value of 0.67 falling outside the random distribution (Fig. 4A). Although this result supports the predominance of morphological homoplasy, it still shows that the potential space for novelties in drosophilids morphology is considerably large.

**ONLY ONE THIRD OF DERIVED CHANGES ARE SHARED**

PAUP* was used to infer the ancestral state at the root of the phylogenetic tree for the 490 characters. Under the ACCTRAN
Homoplasy measurements for whole data, states and characters. (A) Histogram of the whole-data homoplasy index (HI) after 10,000,000 random permutations of taxa over the phylogenetic tree. The empirical value of 0.67 (averaged over 20,086 sampled Bayesian trees) is shown in red and falls outside the random distribution. (B) Proportions of morphological states categorized as derived (Der) and ancestral (Anc) and subcategorized as autapomorphic (A), synapomorphic (S), homoplastic (H) and invariable (I). (C-D) Violin plots of the retention index ($r_i$) with autapomorphic attributed an $r_i$ value of 1 for the 487 morphological characters. Plots are arranged according to (C) the developmental stage and (D) the adult organs. White dots inside the boxes indicate the distribution median. L1, L2 and L3 refer to 1st, 2nd and 3rd larval instars, respectively. Polynomial regression suggesting a “developmental hourglass model” is shown as red curve in (C).

DEVELOPMENTAL CHANGES IN HOMOPLASY SUPPORT A MORPHOLOGICAL ‘HOURGLASS’ MODEL

Homoplasy, as measured by the character retention index with characters with only autapomorphic states attributed an $r_i$ value of 1 (here $r_i$), significantly differed among developmental stages (Kruskall–Wallis chi-squared = 34.013, df = 5, $P = 2.37 \times 10^{-6}$; Table S3; Fig. 4C). The difference formed a concave pattern (polynomial regression: $r_i = 0.755 - 0.245D + 0.037D^2$, $P = 0.003$, with $D$ referring to the developmental stage as egg, larva 1, larva 2, larva 3, pupa, and adult with a unit increment). This quadratic regression had a lower Akaike Information Criterion (AIC) than a linear regression (352.45 vs. 363.20, respectively). The concave model, 220 of the 490 root states have been secondarily derived. Among the 989 derived states, 429 were derived once. Of these, 137 states (~32%) are shared between at least two taxa, that is, synapomorphic states (Fig. 4B). Under the DELTRAN model, 122 root states were secondarily derived, and 113 of 405 apomorphic states were synapomorphic (~28%). That means that only one third of states occupying novel positions on the morphospace would remain unchanged for longer evolutionary times. This result cautions against homoplasy quantification from only phylogenetically informative characters, that is, after excluding characters with only autapomorphic derived states, as it is customary in cladistic analyses (Bryant 1995).
pattern, which supports the “developmental hourglass model” (see Discussion below), did not differ when autapomorphic characters were excluded ($r = 0.588 - 0.201 D + 0.030 D^2, P = 0.006$). No differences were found between the ACCTRAN and DELTRAN models.

**MALE AND FEMALE TERMINALIA HAVE THE LEAST HOMOPLASTIC CHARACTERS**

Homoplasy also differed between adult organs (Kruskal–Wallis chi-squared = 41.935, $df = 4, P = 1.72 \times 10^{-6}$), with male and female terminalia characters being the least homoplastic compared to the three somatic organs, that is, the head, the thorax, and the abdomen (Wilcoxon test: $W = 10986$, $P = 7.39 \times 10^{-9}$; Table S3; Fig. 4D). This difference did not persist when autapomorphic characters were excluded (Wilcoxon test: $W = 7119$, $P = 0.18$). Although on average, male terminalia had higher $r_w$ than female organs (0.73 vs. 0.61, respectively), the difference was not significant (Wilcoxon test: $W = 1107.5$, $P = 0.158$). Exclusion of autapomorphic characters did not lead to significant difference either (Wilcoxon test: $W = 349$, $P = 0.32$). This lack of strong divergence between the sexes in terminalia traits supports the hypothesis of genital coevolution for which several lines of evidence have recently been demonstrated in *Drosophila* (Kamimura 2007; Yassin and Orgogozo 2013) and other animals (Yassin 2016). No differences were found between the ACCTRAN and DELTRAN models.

**PARALLELISM AND CONVERGENCE ACCOUNT FOR 13% OF PAIRWISE SPECIFIC SIMILARITIES**

Proportions of instances belonging to each of the six possible cases of character changes significantly differed (Table S3, Fig. 5A and B). For similarity between pairs of species, the most frequent case was that of phylogenetic inertia (ACCTRAN = 53.77%, DELTRAN = 54.20%), that is, the two species have a state that derived from the state present in their last common ancestor. Under the ACCTRAN model, this figure was followed by both convergence (4.22%) and parallelism (3.61%), which together indicates that homoplasy does not exceed 13% of cases where two species have similar character states. Under DELTRAN, parallelism (4.22%) exceeded convergence (1.99%), with sum homoplasy counting for 10.3% of species comparisons with similar states. For differences between species, the most frequent case was that of Disparity 2 (ACCTRAN = 34.86%, DELTRAN = 34.72%), wherein different states arose from different ancestral states. This was followed by Disparity 1 wherein the ancestral states were similar but of different origins (ACCTRAN = 3.54%, DELTRAN = 4.56%). No (0.00%) and 94 cases (0.30%) of divergence were found in pairwise comparisons where different states diverge from the same different ancestral state under the ACCTRAN and DELTRAN models, respectively.

**REVERSALS OCCUR IN 7% OF DESCENDANT–ANCESTRAL COMPARISONS**

Two hundred sixty-one (53%) and 363 (74%) characters showed no single occurrence of reversals under the ACCTRAN and DELTRAN reconstructions, respectively. On average, reversals were found in 6.65% and 2.63% of all comparisons between terminal taxa and their ancestors according to the two models. The frequency of reversals correlated with both parallelism and convergence, although its effect on convergence was higher under the ACCTRAN model ($R^2 = 0.62$ and 0.29 for convergence and parallelism, respectively). Similar correlations were also found under the DELTRAN model ($R^2 = 0.65$ and 0.11 for convergence and parallelism, respectively). These results suggest that high incidences of convergence and parallelism could be mostly due to recurrent reversals (Table S3, Fig. 5C and D).

**Discussion**

**WHAT IS THE EXTENT OF MORPHOLOGICAL HOMOPLASY?**

Our analyses on 490 morphological traits indicate that nearly two thirds of morphological changes are homoplastic. Our ensemble consistency index (CI) estimate of 0.33 approaches that obtained in other cladistics analyses with >300 morphological characters, for example, in Diptera (CI = 0.40, Lambkin et al. 2013), Birds (0.24, Livezey and Zusi 2007), Birds and related Dinosaurs (0.27, Godefroit et al. 2013), Squamates (0.16, Conrad 2008; 0.21, Gauthier et al. 2012), and Placental Mammals (0.20, O’Leary et al. 2013). Our study differs in its narrower taxonomic scale (i.e., all species belonging to the same family) and its quantification of morphological homoplasy on an independently inferred molecular tree. Zou and Zhang (2016) have recently compared O'Leary et al.’s (2013) ~3,500 morphological traits with a molecular phylogeny and estimated a morphological ensemble CI value of ~0.25. Testing whether the potential of morphological innovations is larger in Dipterans (or Insects) than in Vertebrates would require further developments and refinements of the so-called “giant taxon-character matrices” (Simões et al. 2017; Laing et al. 2018).

**HOW MANY MORPHOLOGICAL CHARACTERS ARE THERE?**

In order to draw general conclusions about the extent of homoplasy, one should be aware about how representative is the studied sample of characters. Information on the total number of morphological traits is lacking, not only because their study often requires profound learning and expertise but also because of the lack of a standard conceptualization approach. To the best of our knowledge, our study represents the largest morphological dataset to be analyzed for the prevalence of homoplasy in an insect. Various *Drosophila* evolutionary developmental studies...
WHAT ARE THE GENOMIC UNDERPINNINGS OF MORPHOLOGICAL HOMOPLASY?

Our morphological analyses uncovered two interesting genomic evolutionary trends, namely the developmental “hourglass” model and the rapid evolution of sex-specific traits. The “hourglass model” postulates that intermediate stages of development are evolutionarily less variable than early and later stages (Irie and Kuratani 2014). In the context of our analysis, low variability
correlates with high homoplasy, that is, there are higher chances to find similar states among taxa at larval stages than at the egg, pupal or adult stages. The hourglass model was initially proposed based on morphological data in Vertebrates, but its support in insects mainly came from genomic and transcriptomic analyses in Drosophila (Cruickshank and Wade 2008; Kalinka et al. 2010; Yassin et al. 2010) and Diptera (Jiménez-Guri et al. 2013; Schep and Adryan 2013). Our results support the prevalence of the model at the morphological level in an insect but it is important to note that we did not study embryos, since the egg analyzed here is the female gamete with most features such as size, shape, and chorionic specializations (e.g., dorsal appendages, ridges, and sculpture) being determined during oogenesis. Moreover, characters were unevenly distribution on the phylogeny (Fig. 1). Investigating these missing data in future comparative studies should improve our tests of different developmental patterns.

Song and Bucheli (2010) found no significant difference in the phylogenetic signal between genital and non-genital traits in a meta-analysis of 41 studies in insects wherein autapomorphic characters were excluded. Similarly, we did not find a difference in the amount of homoplasy between adult genital and non-genital characters when autapomorphies were also excluded. Nonetheless, including autapomorphies revealed a sharp difference. Rapidly evolving characters should be richer in autapomorphic states, because such states have either recently appeared or rapidly changed in other lineages. Our finding hence is consistent with previous studies using fewer morphological characters in Drosophila (Civetta and Singh 1998) and with recent genomewide analyses revealing faster evolution of traits and genes with sex-limited expression (Haerty et al. 2007; Parsch and Ellegren 2013), most likely due to divergent sexual selection that could ultimately lead to fewer homoplasy.

Ever since the early days of Drosophila genetics, a major question has arisen on the relationship between the genetic mutations discovered in D. melanogaster and the morphological differences between wild species (Sturtevant 1921). Patterson and Stone (1952) argued that resemblance between morphological differences and D. melanogaster mutants might involve homologous or quite different loci, which seems to be the case as unraveled by recent evo-devo studies (e.g., Frankel et al. 2012, 2016b; Yassin et al. 2016a; Yassin et al. 2016b). On the online database Flybase, the effect of genetic mutations on different anatomical structures is curated, and for each gene in D. melanogaster links to its orthologous genes (when present) in other Drosophila species are also provided. However, data on the evolutionary differences in these structures are still scattered in the taxonomic literature for the nearly 4,500 drosophilid species. The matrix provided here represents a first attempt to curate these disperse data and future research should complement missing data, adding more taxa and enriching Flybase resources with structures evolution. This endeavor would significantly promote the identification of the genetic underpinnings of morphological changes leading to better character state delimitation. Given the prevalence of the “deep homology” of developmental pathways in animals, such knowledge would ultimately lead to a better understanding of the molecular basis of morphological homoplasy in Drosophila and beyond.

**AUTHOR CONTRIBUTIONS**

S. A. S. conceptualized and coded the morphological characters and performed the phylogenetic analyses. A. Y. conceived the project, designed the study, supervised the analyses and wrote the paper with input from first author.

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**DATA ARCHIVING**

Data associated with this article are all provided as supporting files.

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**Supporting Information**

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Supporting Information

**Table S1.** Phylogenetic classification, data origin and GINBANK accession numbers for taxa sampled in the present study.

**Table S2.** RGB coding and NBClust analysis for the 60 colors described in the taxonomic literature.

**Table S3.** Character evolution statistics (ci=consistency index, ri=retention index, rc=rescaled consistency index, hi=homoplasy index) and proportions of different evolutionary mechanisms for the 490 morphological characters analyzed in the present study.

Supporting Information