INTRODUCTION

Integration between phenotype expression—be it in morphology, physiology or behaviour—and overall body size represents a hallmark of living forms (Schmidt-Nielsen, 1984). The formation and evolution of such covariation, or allometry sensu lato, has received long-standing attention in evolutionary biology, not least because body size is closely linked to fitness, rapidly responds to selection (Blanckenhorn, 2000; Fairbairn, Blanckenhorn, & Székely, 2007; Peters, 1986), and frequently elicits correlated evolutionary responses in integrated traits (Gould, 1966; Lande, 1979; Pelabon et al., 2014). However, even though covariation between traits may itself represent an adaptation shaped by natural selection (Cheverud, 1982, 1984), it can constrain or bias evolutionary trajectories of complex phenotypes (Gould, 1966; Klingenberg, 2014; Schluter, 1996). Therefore, in order to understand and predict evolutionary change, we not only need to understand how allometric variation arises and is maintained, but also how static scaling relationships hamper or bias the evolution of
differentiation. Yet, despite more than a century of research on scaling relationships, their ultimate and proximate drivers as well as their evolutionary consequences remain an enigmatic problem (Casasa, Schwab, & Moczek, 2017; O’Brien, Katsuki, & Emlen, 2017; Pelabon et al., 2014; Shingleton & Frankino, 2018; Shingleton, Frankino, Flatt, Nijhout, & Emlen, 2007).

While many researchers emphasize the evolutionary constancy of static scaling relationships via various mechanisms (Bolstad et al., 2015; Gould, 1966, Kleiber, 1947; Vojé, Hansen, Egset, Bolstad, & Pelabon, 2014), others highlight the potential of particularly rapid divergence (Casasa et al., 2017; Emlen, 1996; Frankino, Zwaan, Stern, & Brakefield, 2005; Puniamoorthy, Blankenhorn, & Schäfer, 2012; Rohner & Blankenhorn, 2018; Wilkinson, 1993). At least some discrepancies about the speed of allometric evolution can be ascribed to methodological and conceptual differences in how scaling relationships are studied. Traditionally, allometry-related concepts were based on the (allometric) coefficient in an exponential equation (e.g., Gould, 1966; Huxley, 1932; here referred to as ‘narrow-sense’ allometry); however, many contemporary researchers use the term allometry to describe various forms of covariation between size and organismal shape (Crabtree, Macagno, Moczek, Rohner, & Hu, 2020; Larson et al., 2018), or, in fact, virtually any phenotype of interest (sexual dimorphism: Fairbairn, 1997; life history: Marbà, Duarte, & Agusti, 2007; behaviour: Dial, Greene, & Irschick, 2008). As some researchers apply a more inclusive concept of allometry than others, this necessarily causes disagreement over what extent allometries differ and hence on how fast they evolve. Irrespective of the concept of allometry, however, there is little doubt that static allometries can affect the course of evolution. Yet it remains unclear how often and to what extent this is the case, whether such effects vary among traits or forms of selection, and to what degree they depend on the conceptual/mathematical approach that is used to study allometry in the broadest sense.

The forewing of Drosophila melanogaster has been widely used to study the evolution of static allometry and developmental plasticity more generally with a variety of approaches (Bolstad et al., 2015; Debat, Begin, Legout, & David, 2003; Gidaszewski, Baylac, & Klingenberg, 2009; Gilchrist & Partridge, 2001; Houle, Jones, Fortune, & Szteganacz, 2019; Robertson & Reeve, 1952; Weber, 1990). Despite the ecological and morphological diversity observed among drosophilids, static narrow-sense allometries of the length of one of the major longitudinal wing veins are very similar across species (Bolstad et al., 2015). Because the drosophilids rapidly diversified about 40 million years ago, this suggests a strongly conserved scaling relationship. While such stasis has traditionally been attributed to selection and/or constraints, (multivariate) stabilizing selection on (multivariate) pleiotropic effects has been shown to be the most likely explanation for the relative constancy of allometric scaling of this particular wing vein in drosophilids (Bolstad et al., 2015; Houle et al., 2019). Given that conserved genes involved in wing development have pleiotropic effects on potentially fitness-related traits (McKay & Lieb, 2013; Ruíz-Losada, Blom-Dahl, Córdoba, & Estella, 2018) and the typical polygenic nature of allometric relationships, similar mechanisms are expected to act in other species or clades. Unfortunately, however, our understanding of wing allometry (and wing shape evolution in general) is primarily based on a few species of Drosophila with most research focusing exclusively on D. melanogaster. It is thus unclear to what extent the findings in drosophilids apply to other groups and if so, how far they can be extrapolated to other dipterans or insects in general.

Drosophilids belong to a division of Diptera called the Schizophora. This group originated around 65 million years ago and underwent rapid diversification. Today, with more than 50,000 species, the Schizophora accounts for more than a third of all fly species (Wiegmann et al., 2011; Wiegmann & Yeates, 2017). The clade is vastly ecologically diverse, is found in all terrestrial habitats and shows a striking diversity in wing morphology that relates to both natural and sexual selection (Dudley, 2002a). Therefore, by investigating static wing allometry across members of this diverse clade should help understand the ecological, developmental or genetic causes and consequences of the evolution of allometry, and whether the evolutionary stasis found in drosophilids also occurs in other groups of flies.

Using geometric morphometric methods, I here investigate the evolution of static and evolutionary allometry in the wings of higher Diptera. I first investigate overall variation in wing morphology among five species belonging to four different families and evaluate the relative contribution of species, sex and size differences. I then investigate the evolution of static allometries in more detail and investigate differences between species and sexes. Lastly, I assess whether static allometries within species are associated with evolutionary divergence across Schizophora by evaluating evolutionary allometry among 33 different families (belonging to 11 of 13 superfamilies following McAlpine (1989); Table S1). I discuss how the patterns across Schizophora relate to the evolutionary stasis found in drosophilids and evaluate the potential roles of selection and pleiotropy in driving static and evolutionary scaling relationships in Diptera.

2 | MATERIALS AND METHODS

2.1 | Dipteron wing morphology

Compared to other insects (e.g., odonates or stoneflies), the forewings of (higher) Diptera have a particularly derived and reduced wing venation pattern (Grimaldi & Engel, 2005; Shimmi, Matsuda, & Hatakeyama, 2014). Despite this reduction, fly wings represent complex functional structures lending dipterans their outstanding flight capacity that facilitates their vast degree of ecological diversification (Dudley, 2002a). Although wing shape varies strongly across all Diptera, wing venation is relatively invariant among Schizophora, allowing to trace the evolution of homologous characters (Ennos, 1989; Henri, 1958, 1981; Redtenbacher, 1886). Wing vein nomenclature used here follows Oosterbroek (2006) (also see McAlpine, 1981).

To quantify static allometries in wing morphology, five species belonging to four distantly related dipteran families (Scathophagidae, Muscidae, Sepsidae and Drosophilidae) were reared under controlled conditions. Other species or clades. Unfortunately, however, our understanding of wing allometry (and wing shape evolution in general) is primarily based on a few species of Drosophila with most research focusing exclusively on D. melanogaster. It is thus unclear to what extent the findings in drosophilids apply to other groups and if so, how far they can be extrapolated to other dipterans or insects in general. Drosophilids belong to a division of Diptera called the Schizophora. This group originated around 65 million years ago and underwent rapid diversification. Today, with more than 50,000 species, the Schizophora accounts for more than a third of all fly species (Wiegmann et al., 2011; Wiegmann & Yeates, 2017). The clade is vastly ecologically diverse, is found in all terrestrial habitats and shows a striking diversity in wing morphology that relates to both natural and sexual selection (Dudley, 2002a). Therefore, by investigating static wing allometry across members of this diverse clade should help understand the ecological, developmental or genetic causes and consequences of the evolution of allometry, and whether the evolutionary stasis found in drosophilids also occurs in other groups of flies. Using geometric morphometric methods, I here investigate the evolution of static and evolutionary allometry in the wings of higher Diptera. I first investigate overall variation in wing morphology among five species belonging to four different families and evaluate the relative contribution of species, sex and size differences. I then investigate the evolution of static allometries in more detail and investigate differences between species and sexes. Lastly, I assess whether static allometries within species are associated with evolutionary divergence across Schizophora by evaluating evolutionary allometry among 33 different families (belonging to 11 of 13 superfamilies following McAlpine (1989); Table S1). I discuss how the patterns across Schizophora relate to the evolutionary stasis found in drosophilids and evaluate the potential roles of selection and pleiotropy in driving static and evolutionary scaling relationships in Diptera.

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laboratory conditions. Although they all can be reared on decaying organic matter at the larval stage (as is common in flies), they vary strongly in the ecology of their adult forms. The yellow dung fly *Scathophaga stercoraria* (Linnaeus, 1758) is a large, predatory, cold-adapted species of Scathophagidae common across the Holarctic. The common housefly, *Musca domestica* Linnaeus, 1758, represents an extreme generalist common across the globe. The afro-tropical *Sepsis lateralis* Wiedemann, 1830 and the holarctic *Saltella sphondylii* (Schrank, 1803) both belong to Sepsidae, but strongly differ in their overall appearance, adult ecology and wing morphology (Pont & Meier, 2002). *Drosophila prolongata* (Singh and Gupta, 1977) is a cold-loving fruit fly found in South-East Asia. Laboratory cultures of all species were provided with a plastic dish filled with a food source that acted as an oviposition plate. These plates were removed after 24–48 hr. Eggs were retrieved and haphazardly distributed among plastic containers that held varying amounts of food (see Rohner, Teder, Esperk, Lüpold, & Blanckenhorn, 2018 for additional information). This resulted in varying degrees of larval competition for food and hence caused strong variation in adult body size. *Drosophila prolongata* was reared on standard *Drosophila* medium, whereas the other four species were reared in homogenized, previously frozen cow dung. The size variation within species and sex was extensive and approximates the maximal size range that can be achieved with food manipulation, facilitating accurate estimation of allometric slopes despite large differences in sample size per species and sex (*D. prolongata*: 166 females, 119 males; *M. domestica*: 56 females, 45 males; *S. sphondylii*: 37 males, 39 females; *Sc. stercoraria*: 213 females, 210 males; *Se. lateralis*: 52 females, 43 males). Upon adult eclosion and complete hardening, animals were killed, and their wings were removed and embedded in Euparal on glass slides. Slides were dried and later photographed using a LeicaDFC490 camera mounted on a Leica MZ12 microscope.

To investigate wing shape and its covariation with size, I integrated several methods of geometric morphometrics. Only eight landmarks that could unambiguously be identified in all specimens were used. These were placed at the intersections between the longitudinal veins R1, R2+3 and R4+5 and the costa, the most proximal edge of the wing cell R2+3, as well as the anterior and posterior major axes of shape variation (using the geomorph function *calcProcrustes*). To assess whether size contributes to the major axis of shape variation, Pearson correlation coefficients were used to investigate the relationship between PC1 and log-centroid size.

### 2.2 Variation in wing shape

First, a principal component analysis (PCA) was used to visualize the major axes of shape variation (using the *plotTangentSpace*()-function as implemented in geomorph). To assess whether size contributes to the major axis of shape variation, Pearson correlation coefficients were used to investigate the relationship between PC1 and log-centroid size.

#### 2.3 Static allometries

A Procrustes ANOVA was used to test for species and sex differences in allometry across the whole data set using the function *procD.lm()* as implemented in geomorph. Because this model indicated that allometries differed between species and sexes (significant log-centroid size × sex × species interaction; see Results), sex-specific multivariate regressions of shape on size were calculated for each species separately. The vectors of coefficients of these regressions represent the multivariate ‘broad-sense’ form of static allometry for each sex and species, respectively. These allometric vectors (A) were then used to compute allometric spaces using ordination of allometric vectors to illustrate variation in allometric scaling (Gerber, Eble, & Neige, 2008; Gerber & Hopkins, 2011; Strelin, Benitez-Vieyra, Fornoni, Klingenberg, & Cocucci, 2018). To this end, I used the R-function *pcorpcor()* based on the covariance matrix of all static allometric vectors. Such ordination resulted in an allometric space where each point represents an allometric vector (rather than an individuals’ shape as in an ordinary morphospace), where distances between points relate to the similarity in this particular allometric space.

For a more detailed assessment of pairwise sex and species differences in allometric scaling, I computed pairwise vector correlations following:

\[
\mathbf{r}_{\mathbf{A}_i \mathbf{A}_j} = \frac{|\mathbf{A}_i \cdot \mathbf{A}_j|}{||\mathbf{A}_i|| \times ||\mathbf{A}_j||}.
\]

That is, the dot product of the allometric vectors \(\mathbf{A}_i\) and \(\mathbf{A}_j\) was standardized by their norm (Claude, 2008; Pitchers, Pool, & Dworkin, 2013; Schäfer et al., 2018). Because correlations between sexes were much higher than interspecific correlations, pooled within-sex vectors were compared when calculating correlations between species. Confidence intervals surrounding these correlations were estimated using a nonparametric bootstrapping procedure (10,000 iterations). The norm of the allometric vector was used to quantify the strength of the allometric effect in unit Procrustes distance per log-millimetre.

The shape score method proposed by Drake and Klingenberg (2008) was used to visualize the allometric relationship across all species. That is, the shape data (y) were projected onto a vector in the direction of a common allometric slope (\(\beta\), as derived from a multivariate regression with all data combined) as \(s = y^\prime(\beta|x^\prime|)^{-0.5}\). The regression score \(s\) can then be used to visualize the strength and shape of the overall relationship between shape and size (Drake & Klingenberg, 2008).
To quantify the scaling relationship among schizophoran species, I selected 33 pinned specimens belonging to 33 different families of both major subsections of the Schizophora (all specimens stem from the private collection of G. Bächli; families and representative species listed in Table S1). This taxon sampling covers 11 of the 13 superfamilies sensu McAlpine (1989). These species are ecologically highly diverse and differ markedly in their size (ranging from the usually 1–3 mm long leafminer flies (Agromyzidae) to the much larger flesh flies (Sarcophagidae) of more than 20 mm body length). Wing morphology, that is venation, colouration, thickness and size, also varied markedly (Figure 1).

Specimens were re-hydrated in a plastic container fitted with wet cotton overnight. Wings were carefully removed, embedded in Euparal, photographed and digitized as described above. After Procrustes transformation, wing shape was subjected to a phylogenetic regression on log-centroid size to estimate evolutionary allometry taking into account the phylogenetic relationship among the dipteran families by Wiegmann et al. (2011) (*procD.pgls* function implemented in *geomorph*). This vector was then compared to the static allometric vectors found within species. Sex could not be taken into account for this analysis. As a phylogenetically weighed estimate of the average static allometry, and to prevent multiple testing, I first computed pooled within-sex static allometries and then used a maximum-likelihood approach implemented in the R-package phytools (*fastAnc*; Revell, 2012) to estimate the shared direction of static allometry among the five species. The correlation between this ‘ancestral’ static allometry and evolutionary allometry was computed as above, but its significance was tested using the approach proposed by Klingenberg and Marugan-Lobon (2013) using the closed-form expression for the area of a hypersphere cap by Li (2011). In brief, the sum of all vectors that have an angle of $\alpha$ or less relative to a fixed

**FIGURE 1** Wing morphology of selected species used in the comparative data set. Two-dimensional landmarks were placed at the intersections between the longitudinal veins $R_1$, $R_{2+3}$, and $R_{4+5}$, and the costa, the most proximal edge of the wing cell $r_{2+3}$, as well as the anterior and posterior ends of both cross veins R-M and DM-Cu.

2.4 Evolutionary allometry across Schizophora

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vector can be represented as the cap of the hypersphere. Dividing the area of this cap by the total surface of the hypersphere (representing a random sample of a uniform distribution) equals the probability that a vector drawn at random from a uniform distribution has an angle $\leq \alpha$. This ratio then represents the $p$-value (see Klingenberg and Marugan-Lobon (2013) for a detailed description of the method).

3 | RESULTS

3.1 | Variation in wing shape

The major axis of variation (PC1) mostly related to the placement of the anterior cross vein (R-M) and an elongation of the proximal part of the wing relative to the distal part. PC2 mostly affected the posterior cross vein and landmarks at the tip of the wing (Figure 2a). PC1 and PC2 explained 59.4% and 21.8% of the total shape variation, respectively, and discriminate between species (Figure 2a), as is expected because they differ markedly in their wing shape (Figure 3). Intraspecific variation in shape was pronounced, but sexual dimorphism does not seem to constitute a major contributor. In contrast, PC1 was strongly related to log-centroid size ($r = .87$, $p < .001$; Figure 1b). As this correlation was also present within species (Sc. stercoraria: $r = .67$, $p < .001$; D. prolongata: $r = .73$, $p < .001$; Sa. sphondylil: $r = .83$, $p < .001$; Se. lateralis: $r = .86$, $p < .001$; M. domestica: $r = .45$, $p < .001$; Figure 1b), this suggests that the association between size and PC1 is not merely driven by interspecific differentiation but that a similar covariation holds within species. Repeating this principal component analysis based on the pooled within-species variance–covariance matrix leads to the same conclusion as the resulting first common principal component (explaining ~35% of the total variance) correlated strongly with log-centroid size ($r = .85$).

3.2 | Static allometries

A Procrustes ANOVA indicated that static allometries differed between species and sexes (significant log-centroid size $\times$ sex $\times$ species interaction in Table 1). Corroborating the overall patterns of the principal component analysis on wing shape, species differences ($\eta^2 = 0.50$) and log-centroid size ($\eta^2 = 0.46$) accounted for most of the variance in shape, whereas sex and the interaction terms had a much smaller effect size (all $\eta^2 \leq 0.03$; Table 1). To investigate differences in allometries in more detail, sex-specific multivariate regressions were fitted for each species separately. All species showed covariation between shape and size when analysed using multivariate regression, but the strength (i.e., the vector norm) of this effect differed between species and somewhat between sexes (Table 2). Musca domestica showed relatively weak deformations (average between sexes: 0.06), followed by Sc. stercoraria (0.10) and D. prolongata (0.12), whereas the two sepsids showed much stronger responses (Se. lateralis: 0.18; Sa. sphondylil: 0.17). Sex differences were particularly pronounced in Sa. sphondylil and Se. lateralis where females reacted much stronger to an increase in log-centroid size (Table 2).

Ordination of the sex-specific allometric vectors yielded an allometric space where the first two principal components explained 60.7% and 25.7% of the total variance, respectively. This is a large portion of the total variance, indicating that variation in wing shape allometries can be summarized in relatively few dimensions. PC1 and PC2 clearly distinguish between species (see Figure 2c), but in contrast to the morphospace (Figure 2a), the allometric space indicates comparably small interspecific variation relative to intraspecific variation in the form of sex differences in allometric slopes.

Pairwise vector correlations between static allometric vectors were moderate to strong between Scathophaga, Musca, Saltella and Sepsis (average correlation $r = .47$; Table 3). Only the pairwise comparison between D. prolongata and Scathophaga as well as Saltella was weak and not significantly different from zero (Table 3). When visualizing the common allometric variation using regression scores (Figure 2d), however, all species followed a similar allometric trajectory. The shape deformation associated with this regression score was very similar to the variation captured by PC1 (Figure 2a vs. d), reiterating the notion that a common allometric variation may explain a large portion of the total variance in wing shape.

3.3 | Evolutionary allometry across Schizophora

Wing shape varied strongly among families (Figure 4). The multivariate extension of Blomberg’s Kappa ($K_{mult} = 0.49$, $p = .024$; Figure 4) was smaller than expected under Brownian motion ($K_{brown} = 1.00$), indicating little phylogenetic inertia (Adams, 2014). There was, however, evidence for evolutionary allometry in wing shape ($r = .38$, $p = .005$) which mostly entailed changes in the aspect ratio and the positioning of the two cross veins (Figure 4). The vector of this evolutionary shape change correlated significantly with the ancestral (i.e. phylogenetically weighed average) static relationship ($r = .49$, $p = .023$).

4 | DISCUSSION

Studying multivariate static allometries in five distantly related species of flies led to three main findings: Firstly, although species strongly diverged in wing shape (Figures 2a and 3), size-related changes represent a major contributor to overall variation, both within and among species (Figure 2b). Secondly, static allometries differ between species and sexes, yet allometric changes are correlated across species, suggesting a shared developmental programme underlying size-dependent shape variation (Figure 2c and Table 3). Lastly, static allometries also relate to the evolutionary divergence across the Schizophora. Together, these findings point towards a general mechanism that canalizes or maintains the covariation between shape and size despite rapid ecological and morphological diversification during the Cenozoic. I in turn discuss the putative roles of selection and developmental constraints in the evolution of wing allometry in the Schizophora.

Flight is the primary means of dispersal in (most) dipterans and is therefore critically involved in mate finding, courtship,
thermoregulation, foraging and various other behaviours (Dillon, Wang, Garrity, & Huey, 2009; Dudley, 2002a). Because the physical forces involved in flight directly relate to size, and natural as well as sexual selection act on performance and manoeuvrability, selection is expected to act on size-dependent compensatory adjustments of wing morphology (Dudley, 2002b; Unwin & Corbet, 1984) and flight behaviour (Lehmann, 1998, 2002; Sane, 2003). Stabilizing selection on a shared developmental programme that accommodates size-related

FIGURE 2  (a) The first two principal components cumulatively explain ~ 80% of the total variation in shape and differentiate between the five species investigated here. Shape change associated with an increase in PC1 and PC2 is indicated in sketches. (b) Plotting PC1 against log-centroid size reveals that the first principal component is strongly related to variation in size, both within and among species. (c) Ordination of sex-specific multivariate allometries in an allometric space shows clear differences in allometric relationships among species; this, however, seems small compared to intraspecific variation in the form of sex differences. (d) Drake & Klingenberg's regression score, a metric representing a projection of the individual wing shape data on the allometric vector across all samples. The shape deformations associated with an increase in the shape score correspond well to the shape changes captured by PC1 (a)
physical constraints therefore appears as a likely general mechanism that canalizes allometry in a wide range of species. Such an adaptive scenario would not only explain the similarities in static allometries between species, but also the association between static allometries and evolutionary divergence. Because the allometric changes mostly involve the positioning of the cross veins and the overall aspect ratio, which most probably relate to the stiffness of the wing blade and lift production, respectively, such an adaptive scenario seems likely.

However, direct evidence for a functional relevance of wing shape and its allometry are scarce. While repeatable plastic and genetic responses, as well as functional implications have been demonstrated for wing size (Azevedo, James, McCabe, & Partridge, 1998; Gilchrist, Azevedo, Partridge, & O’Higgins, 2000; Rohner, Pitnick, et al., 2018; Rohner, Roy, Schäfer, Blanckenhorn, & Berger, 2019; Starmer & Wolf, 1989; Frazier, Harrison, Kirkton, & Roberts, 2008), the functional relevance of the relative positioning of individual wing veins remains...
Sex-specific allometries and the length (=norm) of the sex-specific allometric vector (|A|)

|                          | Sexual shape dimorphism | Sex-specific allometries |
|--------------------------|-------------------------|--------------------------|
|                          | Procrustes distance × 10^3 | Correlation between sexes | |A| males × 10^2 | |A| females × 10^2 |
| Drosophila prolongata     | 17.9 [15.3, 20.0]         | 0.93 [0.89, 0.98]        | 11.6 [10.1, 13.1] | 12.8 [11.4, 14.1] |
| Musca domestica           | 21.4 [17.2, 25.2]         | 0.89 [0.84, 0.97]        | 5.5 [3.4, 6.9]    | 6.9 [4.3, 9.2]    |
| Saltella sphondylii       | 21.4 [15.5, 25.0]         | 0.94 [0.90, 0.99]        | 13.8 [10.6, 17.0] | 20.3 [16.5, 23.5] |
| Scathophaga stercoraria   | 12.0 [9.6, 13.9]          | 0.92 [0.86, 0.98]        | 9.8 [7.9, 11.7]   | 9.3 [7.2, 11.4]   |
| Sepsis lateralis          | 11.1 [7.0, 14.2]          | 0.97 [0.94, 0.99]        | 16.8 [13.8, 19.7] | 19.6 [17.1, 22.1] |

Note: Complementary 95% bootstrap intervals are given.

Alternative mechanisms that could lead to shared patterns of static allometry are genetic and/or developmental constraints. Wings might be particularly prone to pleiotropy because of rather few major signalling pathways that govern the patterning of imaginal discs giving rise to the adult wings, halteres and legs (e.g., hedgehog, decapentaplegic, EGFR; Ruiz-Losada et al., 2018). These discs further share a temporal pattern of expression of the same regulatory elements (McKay & Lieb, 2013). Pleiotropy may thus be more common than previously expected and influence organismal evolution more than currently acknowledged (Sabarís, Laiker, Preger-Ben Noon, & Frankel, 2019). However, venation patterns do not seem to be particularly constrained to evolve per se given abundant additive genetic variation for selection to act upon (Mezey & Houle, 2005; Schäfer et al., 2018). Multivariate static allometries also do not seem to be hampered by absolute constraints. The two closely related black scavenger flies Sepsis fulgens and Se. punctum, for instance, show high vector correlations between multivariate allometric shape changes, yet this correlation is clearly distinct from unity (r = .87 [0.76, 0.92] 95% CI). Wing shape changes associated with rearing temperature differed even more between the two sepsids (vector correlation based on 15 landmarks amount to 0.33 or 0.60 depending on the temperature range used; Rohner et al., 2019), demonstrating that also other forms of plasticity evolve in comparably short periods of time. Evidence for rapid evolution of plasticity also comes from the yellow dung fly, for which populations on different continents show different patterns of thermal plasticity and static allometry (Schäfer et al., 2018). While surely biasing the phenotypic variation that is exposed to selection in general, developmental and/or genetic constraints do not seem to put an absolute limit on the evolution of wing allometry and thus are unlikely to account for the significant similarities that persisted despite rapid ecological diversification throughout the Cenozoic.

In drosophilids, the slow evolution of allometric scaling has been previously attributed to stabilizing selection on pleiotropic effects, causing evolutionary stasis (Bolstad et al., 2015; Houle et al., 2019). The multivariate approach followed here at least superficially recapitulates the patterns found for narrow-sense allometries in the Drosophilidae (Bolstad et al., 2015; Houle et al., 2019) and might indicate that similar processes act on a much larger phylogenetic scale. However, although the results found here indicate a shared axis of plasticity among species overall, it also shows that interspecific variation in allometry differs between landmarks. The length of vein R_{4+5} for instance shows similar covariation with size in all five species (e.g., R_{4+5}; Figure 3), whereas other measurements show much more interspecific variation (R_{1}; Figure 3). This suggests that not all landmarks are under the same evolutionary forces and implies that the choice of measurements used for bivariate allometries must heavily influence the evolutionary inference. It is further worth pointing out that D. prolongata had the lowest vector correlations compared with the others (Table 3). Whether this is a species-restricted effect or whether this applies to drosophilids in general remains unclear at this point.

Sexual shape dimorphism and sex-specific allometric scaling have been documented in several species and seem to be the rule, rather than the exception at least among flies (Gidaszewski

TABLE 2 Summary table providing information about the strength of sexual shape dimorphism (in unit Procrustes distance), the correlation between sex-specific static allometries and the length (=norm) of the sex-specific allometric vector (|A|)

|                          | Sexual shape dimorphism | Sex-specific allometries |
|--------------------------|-------------------------|--------------------------|
|                          | Procrustes distance × 10^3 | Correlation between sexes | |A| males × 10^2 | |A| females × 10^2 |
| Drosophila prolongata     | 17.9 [15.3, 20.0]         | 0.93 [0.89, 0.98]        | 11.6 [10.1, 13.1] | 12.8 [11.4, 14.1] |
| Musca domestica           | 21.4 [17.2, 25.2]         | 0.89 [0.84, 0.97]        | 5.5 [3.4, 6.9]    | 6.9 [4.3, 9.2]    |
| Saltella sphondylii       | 21.4 [15.5, 25.0]         | 0.94 [0.90, 0.99]        | 13.8 [10.6, 17.0] | 20.3 [16.5, 23.5] |
| Scathophaga stercoraria   | 12.0 [9.6, 13.9]          | 0.92 [0.86, 0.98]        | 9.8 [7.9, 11.7]   | 9.3 [7.2, 11.4]   |
| Sepsis lateralis          | 11.1 [7.0, 14.2]          | 0.97 [0.94, 0.99]        | 16.8 [13.8, 19.7] | 19.6 [17.1, 22.1] |
Sexual shape dimorphism was variable and almost twice as strong in *Sa. sphondylii* compared to *Se. lateralis*, suggesting that sexual shape dimorphism can differ rather strongly within families. However, the overall effect of sex seems small compared with species differentiation and allometry (Figure 2). Similar patterns were found for the direction of sex-specific allometric slopes (high between-sex correlations in Table 2). However, in contrast to sex differences in the direction of allometry, the strength of the respective allometric change varied (leading to relatively large sex differences in allometric spaces, as the latter also incorporates direction and strength of the allometric effect). Whether the strength and direction of allometry relate to sex-specific differences in function remains unclear but warrants further scrutiny.

5 | CONCLUSIONS

How often and by which means the mechanisms that underlie static size-dependent trait expression can bias evolutionary change remains an open question. Comparing static allometric slopes in a phylogenetic framework suggests a shared developmental programme despite rapid ecological diversification among schizophoran flies. Presupposing functional implications of wing shape, strong stabilizing selection on a common allometric relationship is the most likely explanation. However, pleiotropy may represent a previously underestimated source of constraint. Whether selection or pleiotropy is more important in driving similarities in allometric relationships is, at this point, difficult to assess due to the lack of functional data. The broad comparative framework applied here may, however, prove useful for exposing variation in allometric scaling that can then be used to assess the functional relevance of wing shape and its allometry.

ACKNOWLEDGMENTS

I thank two anonymous reviewers and the editor for very helpful and constructive comments on earlier versions of this manuscript. I am indebted to Dr. Gerhard Bächli for providing me access to his private collection. I also thank Armin P. Moczek and Wolf U. Blanckenhorn for comments on an earlier version of this manuscript. Finally, I would like to acknowledge financial support from the Swiss National Science Foundation in the form of an Early Postdoc.Mobility fellowship (P2ZHP3_184003).

AUTHOR CONTRIBUTIONS

PTR designed the study, analysed all data and wrote the manuscript.

DATA AVAILABILITY STATEMENT

Data will be available in Dryad upon acceptance. https://doi.org/10.5061/dryad.xd2547dcd.

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REFERENCES

Adams, D. C. (2014). A generalized K statistic for estimating phylogenetic signal from shape and other high-dimensional multivariate data. *Systematic Biology*, 63, 685–697.

Adams, D. C., & Otárola-Castillo, E. (2013). geomorph: An R package for the collection and analysis of geometric morphometric shape data. *Methods in Ecology and Evolution*, 4, 393–399.

Azevedo, R. B. R., James, A. C., McCabe, J., & Partridge, L. (1998). Latitudinal variation of wing:thorax size ratio and wing-aspect ratio in *Drosophila melanogaster*. *Evolution*, 52, 1353–1362.
Blankenhorn, W. U. (2000). The evolution of body size: What keeps organisms small? The Quarterly Review of Biology, 75, 385–407. https://doi.org/10.1086/393620

Bolstad, G. H., Cassara, J. A., Marquez, E., Hansen, T. F., van der Linde, K., Houle, D., & Pelabon, C. (2015). Complex constraints on allometry revealed by artificial selection on the wing of Drosophila melanogaster. Proceedings of the National Academy of Sciences of the United States of America, 112, 13284–13289.

Casasa, S., Schwab, D. B., & Moczek, A. P. (2017). Developmental regulation and evolution of scaling: Novel insights through the study of Onthophagus beetles. Current Opinions in Insect Science, 19, 52–60.

Cheverud, J. M. (1982). Phenotypic, genetic, and environmental morphological integration in the cranium. Evolution, 36, 499–516.

Cheverud, J. (1984). Quantitative genetics and developmental constraints on evolution by selection. Journal of Theoretical Biology, 110, 155–171.

Claude, J. (2008). Morphometrics with R. New York, NY: Springer.

Crabtree, J. R., Macagno, A. L. M., Moczek, A. P., Rohner, P. T., & Hu, Y. (2020). Notch signaling patterns head horn shape in the bull-headed dung beetle Onthophagus taurus. Development Genes and Evolution. https://doi.org/10.1007/s00427-020-00645-w [Epub ahead of print].

Debat, V., Begin, M., Legout, H., & David, J. R. (2003). Allometric and nonallometric components of Drosophila wing shape respond differently to developmental temperature. Evolution, 57, 2773–2784.

Dial, K. P., Greene, E., & Irischk, D. J. (2008). Allometry of behavior. Trends in Ecology and Evolution, 23, 394–401.

Dillon, M. E., Wang, G., Garrity, P. A., & Huey, R. B. (2009). Review: Thermal preference in Drosophila. Journal of Thermal Biology, 34, 109–119.

Drake, A. G., & Klingenberg, C. P. (2008). The pace of morphological change: Historical transformation of skull shape in St Bernard dogs. Proceedings of the Royal Society B-Biological Sciences, 275, 71–76.

Dudley, R. (2002a). The biomechanics of insect flight: Form, function, evolution. Princeton, NJ: Princeton University Press.

Dudley, R. (2002b). Mechanisms and implications of animal flight maneuverability. Integrative and Comparative Biology, 42, 135–140.

Emlen, D. J. (1996). Artificial selection on horn length-body size allometry in the horned beetle Onthophagus acuminatus (Coleoptera: Scarabaeidae). Evolution, 50, 1219–1230.

Ennos, A. R. (1989). Comparative functional morphology of the wings of Diptera. Zoological Journal of the Linnean Society, 96, 27–47.

Fairbairn, D. J. (1997). Allometry for sexual size dimorphism: Pattern and process in the coevolution of body size in males and females. Annual Review of Ecology and Systematics, 28, 659–687.

Fairbairn, D. J., Blankenhorn, W. U., & Székely, T. (2007). Sex, size and gender roles: Evolutionary studies of sexual size dimorphism. Oxford, UK: Oxford University Press.

Frankino, W. A., Zwaan, B. J., Stern, D. L., & Brakefield, P. M. (2005). Natural selection and developmental constraints in the evolution of allometries. Science, 307, 718–720.

Frazier, M. R., Harrison, J. F., Kirkton, S. D., & Roberts, S. P. (2008). Cold rearing improves cold-flight performance in Drosophila via changes in wing morphology. Journal of Experimental Biology, 211, 2116–2122.

Gerber, S., Eble, G. J., & Neige, P. (2008). Allometric space and allometric disparity: A developmental perspective in the macroevolutionary analysis of morphological disparity. Evolution, 62, 1450–1457.

Gerber, S., & Hopkins, M. J. (2011). Mosaic heterochrony and evolutionary modularity: The trilobite genus Zacanthopsis as a case study. Evolution, 65, 3241–3252.

Gidaszewski, N. A., Baylac, M., & Klingenberg, C. P. (2009). Evolution of sexual dimorphism of wing shape in the Drosophila melanogaster subgroup. BMC Evolutionary Biology, 9, 110.

Gilchrist, A. S., Azevedo, R. B., Partridge, L., & O’Higgins, P. (2000). Adaptation and constraint in the evolution of Drosophila melanogaster wing shape. Evolution and Development, 2, 114–124.

Gilchrist, A. S., & Partridge, L. (2001). The contrasting genetic architecture of wing size and shape in Drosophila melanogaster. Heredity (Edinb), 86, 144–152.

Gould, S. J. (1996). Allometry and size in ontogeny and phylogeny. Biological Reviews of the Cambridge Philosophical Society, 41, 587–640.

Grimaldi, D., & Engel, M. S. (2005). Evolution of the insects. Cambridge, UK: Cambridge University Press.

Hennig, W. (1958). Die Familien der Diptera Schizophora und ihre phylogenetischen Verwandtschaftsbeziehungen. Beiträge Zur Entomologie, 8, 505–688.

Hennig, W. (1981). Insect phylogeny. Chichester, UK: John Wiley & Sons.

Houle, D., Jones, L. T., Fortune, R., & Szetpanacz, J. L. (2019). Why does allometry evolve so slowly? Integrative and Comparative Biology, 59, 1429–1440. https://doi.org/10.1093/icb/icz099

Huxley, J. S. (1932). Problems of relative growth. London, UK: Methuen & Company Limited.

Kleiber, M. (1947). Body size and metabolic rate. Physiological Reviews, 27, 511–541.

Klingenberg, C. P. (2014). Studying morphological integration and modularity at multiple levels: Concepts and analysis. Philosophical Transactions of the Royal Society of London B: Biological Sciences, 369, 20130249.

Klingenberg, C. P., & Marugan-Lobon, J. (2013). Evolutionary covariation in geometric morphometric data: Analyzing integration, modularity, and allometry in a phylogenetic context. Systematic Biology, 62, 591–610.

Lande, R. (1979). Quantitative genetic analysis of multivariate evolution, applied to brain: Body size allometry. Evolution, 33, 402–416.

Larson, J. R., Manyama, M. F., Cole, J. B., Gonzalez, P. N., Percival, C. J., Liberton, D. K., & Hallgrimsson, B. (2018). Body size and allometric variation in facial shape in children. American Journal of Physical Anthropology, 165, 327–342.

Lehmann, F. O. (1998). The control of wing kinematics and flight forces in fruit flies (Drosophila spp.). Journal of Experimental Biology, 201, 385–401.

Lehmann, F. O. (1999). Ambient temperature affects free-flight performance in the fruit fly Drosophila melanogaster. Journal of Comparative Physiology B, 169, 165–171.

Lehmann, F. O. (2002). The constraints of body size on aerodynamics and energetics in flying fruit flies: An integrative view. Zoology, 105, 287–295.

Li, S. (2011). Concise formulas for the area and volume of a hyperspherical cap. Asian Journal of Mathematics and Statistics, 5, 66–70.

Marbá, N., Duarte, C. M., & Agusti, S. (2007). Allometric scaling of plant life history. Proceedings of the National Academy of Sciences of the United States of America, 104, 15777–15780.

McAlpine, J. F. (1981). Morphology and terminology—Adults. In: J. F. McAlpine & D. D. M. Wood (Eds.), Manual of nearctic diptera, Vol. 1. Ottawa, Research Branch Agriculture Canada, Monograph 27. pp. 9–63.

McAlpine, J. F. (1989). Phylogeny and classification of the Muscomorpha. In: J. F. McAlpine & D. D. M. Wood (Eds.), Manual of nearctic diptera. Ottawa, Research Branch Agriculture Canada, Monograph 32.

Mckay, D. J., & Lieb, J. D. (2013). A common set of DNA regulatory elements shapes Drosophila appendages. Developmental Cell, 27, 306–318.

Mezey, J. G., & Houle, D. (2005). The dimensionality of genetic variation for wing shape in Drosophila melanogaster. Evolution, 59, 1027–1038.

O’Brien, D. M., Katsuki, M., & Emlen, D. J. (2017). Selection on an extreme weapon in the frog-legged leaf beetle (Sagra femorata). Evolution, 71, 2584–2598.
Oosterbroek, P. (2006). The European families of the diptera: Identification – biology – ecology. The Netherlands: KNNV Publishing.

Pelabon, C., Firmat, C., Bolstad, G. H., Voje, K. L., Houle, D., Cassara, J., ..., Hansen, T. F. (2014). Evolution of morphological allometry. Annals of the New York Academy of Sciences, 1320, 58–75.

Peters, R. H. (1986). The ecological implications of body size. Cambridge, UK: Cambridge University Press.

Pitchers, W., Pool, J. E., & Dworkin, I. (2013). Alitudinal clinal variation in wing size and shape in African Drosophila melanogaster: One cline or many? Evolution, 67, 438–452.

Pont, A. C., & Meier, R. (2002). The Sepsidae (Diptera) of Europe. Fauna Entomologica Scandinavica, 37, 1-221.

Puniamoorthy, N., Blankenhorn, W. U., & Schäfer, M. A. (2012). Differential investment in pre- vs. post-copulatory sexual selection reinforces a cross-continenal reversal of sexual size dimorphism in Sepsis punctum (Diptera: Sepsidae). Journal of Evolutionary Biology, 25, 2253–2263.

Redtenbacher, J. (1886). Vergleichende studien über das flügelgeänder der insecten. Annalen des k. k. Naturhistorischen Hofmuseums, 1, 11–231.

Revell, L. J. (2012). phytools: An R package for phylogenetic comparative biology (and other things). Methods in Ecology and Evolution, 3, 217–223.

Robertson, F. W., & Reeve, E. (1952). Studies in quantitative inheritance.1. The effects of selection of wing and thorax length in Drosophila melanogaster. Journal of Genetics, 50, 414–448.

Rohlf, F. J. (2009). TpsDig. Stony Brook, NY: Department of Ecology and Evolution, State University of New York.

Rohner, P. T., & Blankenhorn, W. U. (2018). A comparative study of the role of sex-specific condition dependence in the evolution of sexually dimorphic traits. The American Naturalist, 192, E202–E215.

Rohner, P. T., Pitnick, S., Blankenhorn, W. U., Snook, R. R., Bächli, G., & Lümpold, S. (2018). Interrelations of global macroecological patterns in wing and thorax size, sexual size dimorphism, and range size of the Drosophilidae. Ecography, 41, 1707–1717.

Rohner, P. T., Roy, J., Schäfer, M. A., Blankenhorn, W. U., & Berger, D. (2019). Does thermal plasticity align with local adaptation? An interspecific comparison of wing morphology in sepsid flies. Journal of Evolutionary Biology, 32, 463–475.

Rohner, P. T., Teder, T., Esperk, T., Lümpold, S., & Blankenhorn, W. U. (2018). The evolution of male-biased sexual size dimorphism is associated with increased body size plasticity in males. Functional Ecology, 32, 581–591.

Ruiz-Losada, M., Blom-Dahl, D., Córdoba, S., & Estella, C. (2018). Specification and patterning of Drosophila appendages. Journal of Developmental Biology, 6, 17.

Sabaris, G., Laiker, I., Prager-Ben Noon, E., & Frankel, N. (2019). Actors with multiple roles: Pleiotropic enhancers and the paradigm of enhancer modularity. Trends in Genetics, 35, 423–433.

Sane, S. P. (2003). The aerodynamics of insect flight. Journal of Experimental Biology, 206, 4191–4208.

Schäfer, M. A., Berger, D., Rohner, P., Kjaersgaard, A., Bauerfeind, S., Guillaume, F., ... Blankenhorn, W. U. (2018). Geographic clines in wing morphology relate to biogeographic history in New World but not Old World populations of dung flies. Evolution, 72, 1629–1644.

Schluter, D. (1996). Adaptive radiation along genetic lines of least resistance. Evolution, 50, 1766–1774.

Schmidt-Nielsen, K. (1984). Scaling: Why is animal size so important? Cambridge, UK: Cambridge University Press.

Shinn, O., Matsuda, S., & Hatakeyama, M. (2014). Insights into the molecular mechanisms underlying diversified wing venation among insects. Proceedings of the Royal Society B-Biological Sciences, 281, 20140264.

Singleton, A. W., & Frankino, W. A. (2018). The (ongoing) problem of relative growth. Current Opinion in Insect Science, 25, 9–19.

Singleton, A. W., Frankino, W. A., Flatt, T., Nijhout, H. F., & Emlen, D. J. (2007). Size and shape: The developmental regulation of static allometry in insects. BioEssays, 29, 536–548.

Starmer, W. T., & Wolf, L. L. (1989). Causes of variation in wing loading among Drosophila species. Biological Journal of the Linnean Society, 37, 247–261.

Strelin, M. M., Benitez-Vieyra, S., Fornoni, J., Klingenberg, C. P., & Cocucci, A. (2018). The evolution of floral ontogenetic allometry in the Andean genus Coiophora (Loasaceae, subfam. Loasoideae). Evolution and Development, 20, 29–39.

Unwin, D. M., & Corbet, S. A. (1984). Wingbeat frequency, temperature and body size in bees and flies. Physiological Entomology, 9, 115–121.

Voje, K. L., Hansen, T. F., Egset, C. K., Bolstad, G. H., & Pelabon, C. (2014). Allometric constraints and the evolution of allometry. Evolution, 68, 866–885.

Weber, K. E. (1990). Selection on wing allometry in Drosophila melanogaster. Genetics, 126, 975–989.

Wiegmann, B. M., Trautwein, M. D., Winkler, I. S., Barr, N. B., Kim, J. W., Lambkin, C., ... Yeates, D. K. (2011). Episodic radiations in the fly tree of life. Proceedings of the National Academy of Sciences of the United States of America, 108, 5690–5695.

Wiegmann, B. M., & Yeates, D. K. (2017). Phylogeny of Diptera. In A. H. Kirk-Spriggs, & B. J. Sinclair (Eds.), Manual of afrotropical Diptera. Volume 1. Introductory chapters and keys to Diptera families. Suricata 4 (pp. 253–265). Pretoria, South African: South African National Biodiversity Institute.

Wilkinson, G. S. (1993). Artificial sexual selection alters allometry in the stalk-eyed fly Cyrtodiopsis dalmanni (Diptera, Diopsiidae). Genetical Research, 62, 213–222.

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

How to cite this article: Rohner PT. Evolution of multivariate wing allometry in schizophoran flies (Diptera: Schizophora). J Evol Biol. 2020;00:1–11. https://doi.org/10.1111/jeb.13613