

Body stores of emergent aquatic insects are associated with body size, sex, swarming behaviour and dispersal strategies

Rémi Gerber, Lilian Cabon, Christophe Piscart, Jean-Marc Roussel, Benjamin Bergerot

To cite this version:

Rémi Gerber, Lilian Cabon, Christophe Piscart, Jean-Marc Roussel, Benjamin Bergerot. Body stores of emergent aquatic insects are associated with body size, sex, swarming behaviour and dispersal strategies. Freshwater Biology, 2022, 67 (12), pp.2161-2175. $10.1111/fwb.14003$. hal-03829811

HAL Id: hal-03829811 <https://hal.inrae.fr/hal-03829811v1>

Submitted on 30 Oct 2022

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

[Distributed under a Creative Commons Attribution 4.0 International License](http://creativecommons.org/licenses/by/4.0/)

ORIGINAL ARTICLE

Body stores of emergent aquatic insects are associated with body size, sex, swarming behaviour and dispersal strategies

Rémi Gerber1,2 | **Lilian Cabon1,2** | **Christophe Piscart1** | **Jean-Marc Roussel2** | **Benjamin** Bergerot¹ [®]

1 CNRS, ECOBIO – UMR 6553, Univ Rennes, Rennes, France

2 DECOD (Ecosystem Dynamics and Sustainability), INRAE, Institut Agro, IFREMER, Rennes, France

Correspondence

Rémi Gerber, CNRS, ECOBIO – UMR 6553, Univ Rennes, Rennes F-35042, France. Email: remi.gerber@univ-rennes1.fr

Funding information

Earth Sciences and Astronomy Observatory (OSU) Rennes, OSUR; LTSER Zone Atelier Armorique

Abstract

- 1. Dispersal is a determining step in the life cycle of insects and a key factor of their energy budget. If the body stores of terrestrial insects are relatively welldocumented, those of merolimnic insects (i.e., with aquatic larvae and terrestrial winged adults) remain poorly investigated.
- 2. We assayed the body stores (protein, carbohydrate and lipid contents) of 18 families of merolimnic insects, encompassing a wide diversity of biological and ecological traits.
- 3. We highlighted allometry between body stores and mass. The proportion of triglyceride and free carbohydrate relatively decreases with an increase in absolute insect mass, whereas protein and glycogen contents are proportional.
- 4. We found a significant effect of insect order, Diptera having relatively more proteins than Trichoptera. For swarming taxa, males have relatively more triglycerides than their conspecific female. Passive dispersers are characterised by higher protein and glycogen contents
- 5. Each body store was associated with specific life-history traits, emphasising the importance of considering each body store independently to better understand the associated functions.
- 6. Dispersal strategies seem to be the main structuring factor underlying body store patterns of merolimnic insects, being significantly associated with the variation in adjusted triglycerides (30%), proteins (21%), glycogen (29%) and free carbohydrates (17%). However, our study estimated the role of flight only indirectly through the biological traits of insects. Future works should focus on identifying which substrates are used for flight, by comparing the proportion of body stores before and after flight, as well as protein identification to distinguish proteins associated with flight or reproduction. These further studies will help researchers in understanding the link between body stores of emerging merolimnic insects and the various facets of their ecology.

Gerber and Cabon to be considered as joint first author.

This is an open access article under the terms of the [Creative Commons Attribution](http://creativecommons.org/licenses/by/4.0/) License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

© 2022 The Authors. *Freshwater Biology* published by John Wiley & Sons Ltd.

Freshwater Biology WILEY

1 | **INTRODUCTION**

Freshwater ecosystems cover less than 1% of the Earth's surface but host a remarkable biodiversity with more than 140,000 known species, including approximately 88,000 insect species (Balian et al., 2007). Among these insects, 80% are merolimnic––that is their juvenile life stages are confined to freshwater environments while winged adults are terrestrial (Petersen et al., 2004). The flight of imagoes is crucial for the colonisation of new habitats (Kovats et al., 1996; Petersen et al., 2004), maintaining genetic diversity (Didham et al., 2012; Phillipsen et al., 2015) and defining community composition (Bacca et al., 2021; Tonkin et al., 2018). Flight also is important for fitnessrelated functions such as escaping predators, locating food and reproduction (Amat et al., 2012; Chen et al., 2011; David et al., 2015; Dudley, 2002; Tokeshi & Reinhardt, 1996). Variable flight distances have been reported depending on species, ranging from 160 m for some Ephemeroptera to 17 km for some Chironomidae (Muehlbauer et al., 2014). However, current knowledge on insect flight comes from studies that use heterogeneous methods and target one or a few species at once, leading to limited understanding of flight ability.

One way to approach the flight ability of merolimnic imagoes is to look at their energy content. Flight requires high energy consumption and is a key determinant of the energy budget of insects (Amat et al., 2012). The metabolic rate of insects when in flight can be 50–100-fold higher than their resting metabolic rate, and their flight muscles exhibit the highest metabolic rate of any known animal locomotor tissue (Beenakkers et al., 1984; Downer & Matthews, 1976; Dudley, 2002). Body size is a key constraint for insect metabolism and dispersal strategies. Firstly, the allometric relationship between resting metabolic rate and body mass is widely recognised (Harrison et al., 2014; Maino & Kearney, 2014), and light insects (<10 mg) are expected to consume proportionally less energy during flight than heavy ones (Niven & Scharlemann, 2005). Secondly, light species often are called "passive dispersers" as they are more subjected to wind dispersal (Peredo Arce et al., 2021), when bigger species can fly faster than the wind and choose their flight direction (Compton et al., 2002). Energy comes from the oxidation of one or several energy substrates (i.e., carbohydrates for short flights, to which lipids are added for long flights). Carbohydrates are stored freely in the cytoplasm and in the form of glycogen (Roma et al., 2010). Lipids are stored mostly in the form of triglycerides (Arrese & Soulages, 2010; Winkelmann & Koop, 2007). Some species also oxidise the amino acid proline (Amat et al., 2012; Beenakkers et al., 1984; Tigreros & Davidowitz, 2019). However, this knowledge of insect body stores is based largely on terrestrial species, and studies on emerging merolimnic imagoes are scarce.

Merolimnic insects encompass a large number of species divided into 12 orders (Dijkstra et al., 2014), and represent great variability in terms of phylogenetic relationships, morphology, size and lifehistory traits. Ephemeroptera, Plecoptera, Trichoptera (EPT) and

Diptera account for 85% of all species of merolimnic insects (Dijkstra et al., 2014). Within these orders, Diptera and Trichoptera are phylogenetically the closest and most recent orders, Ephemeroptera is the most ancestral one, and Plecoptera is intermediate (Ishiwata et al., 2011; Yeates et al., 2016). Merolimnic imagoes display variable feeding behaviours, and this may influence energy allocation and the amount of energy stored before their emergence from the water. Diptera and Plecoptera can feed as adults (Armitage, 1995; Brittain, 1990; Burtt et al., 1986; de Figueroa & Sánchez-Ortega, 1999, 2000; Rúa & de Figueroa, 2013), but Trichoptera rarely do so (Morse, 2009; Petersson & Hasselrot, 1994), and Ephemeroptera have no functional mouthparts, relying exclusively on the energy stored during larval development (Jacobus et al., 2019; Sartori & Brittain, 2015). Moreover, merolimnic insects have evolved different mating systems, notably Ephemeroptera and Chironomidae males form swarms (i.e., aggregative flights; Sullivan, 1981) to attract females (Brittain & Sartori, 2009; Harker, 1992; Tokeshi & Reinhardt, 1996). Some Trichoptera families also can swarm (e.g., Hydropsychidae, Leptoceridae) yet this behaviour has never been observed among Plecoptera (Holzenthal et al., 2015). As swarming requires great quantities of energy (Dudley, 2002), males of swarming species could have greater body stores than conspecific females.

In the present work, we studied body store patterns of 18 common families of merolimnic insects belonging to EPT and Diptera. We quantified their free carbohydrate, glycogen, lipid and protein contents, and we searched for variables that might explain variation among species offering a large range of morphology, body size and flying behaviour. The following questions were addressed: Is there allometry between body store and mass of merolimnic species imagoes? Do their body stores vary according to insect order, sex, and/ or between passive and active dispersal strategies? We hypothesised that mass strongly influences the quantity of body stores, with small taxa having relatively more stores than big ones. We also hypothesised that flight-related life-history traits (dispersal strategy and mating system) structure most of the pattern in body stores.

2 | **MATERIALS AND METHODS**

2.1 | **Study area and sampling**

Samples were taken in Brittany (western France), a region characterised by a patchy landscape of forests, pastures, crop cultures and farming zones under a temperate oceanic climate. Insects were caught using aquatic emergence traps (Cadmus et al., 2016). The traps consisted of a floating pyramidal tent $(1 \text{ m}^2 \text{ at its base})$ made of nylon mesh, with a collecting bottle placed at the top (Raitif et al., 2018). Depending on how the targeted insect groups emerged, the traps were placed either on the water (to catch insects emerging from the

 GERBER ET AL. *CRBER ET AL.* **ALL ASSESS** $\frac{1}{2}$ **3**

water surface such as Chironomidae; Pinder, 1986) or placed half on water and half on riverbanks (to capture insects crawling out of the water to moult such as Plecoptera; Hynes, 1976). The traps were lifted every 24 h to minimise energy expenditure after emergence. During the journey back to the laboratory, the insects were placed on ice before freezing.

Emerging merolimnic insects were collected during two sampling campaigns in 2018 (4–25 April; for Leuctridae only) and 2020 (12 March–10 July). Traps were regularly moved along the sampling area to catch species from various microhabitats and were checked nearly every day at the same time. They were carefully emptied of all insects when the sampling could not be done the day before. Emergence traps were settled in four sites close to our institutions: Chênelais (48.503093, −1.598211), Guyoult (48.470309, −1.628374), Petit Hermitage (48.488179, −1.571154) and Gueuche (48.614058, −1.008783). In addition, a citizen science-based sampling was organised to collect insects from four distant sites: Elorn (48.408397, −4.08789), Ruisseau de Condat (47.700761, −2.631534), Arz (47.713978, −2.403896) and Le Loc'h (47.761264, −3.505502). All emergence traps were settled in agricultural landscapes, except Le Petit Hermitage (forested area), and in small, 1–2 order (Strahler's stream order) headwater streams (except a pond at Le Loc'h).

2.2 | **Assessment of body stores**

2.2.1 | Sample preparation

Insects were sorted, sexed and identified at the family, genus or even species level when possible. For each taxonomic group (i.e., the lowest level of identification for each taxon), body store determination was performed on at least seven replicates and when the sampling allowed it, individuals were taken from different sites. The protocol was designed to maximise the number of species, and therefore intertaxa comparisons. The number of individuals *per* replicate depended on the average mass of the taxon. When insects were too light to be assayed individually (all of the Diptera, Baetidae and Leptophlebiidae), two to 10 individuals of the same sex were pooled in each replicate (Table 1). Each replicate was freeze-dried for at least 24 h and weighed (XP2U Mettler Toledo microbalance, \pm 0.01 mg) to provide the dry mass.

2.2.2 | Body store assays

The insects were placed in phosphate buffer (300 and 600 μl for replicates below and above 500 μg, respectively), homogenised in a bead-beating tube for 1 min 30 s at 25 Hz, and centrifuged for 5 min at 500 *g* and 4°C. Protein and glycogen assays were realised according to Andrade et al. (2020). The total protein content was assayed with the Bradford method (Bradford, 1976) using bovine serum albumin as a standard. The free carbohydrate and glycogen contents were assayed with the Anthrone method. Additionally

to Andrade et al. (2020), we assayed the free carbohydrates as suggested by Foray et al. (2012). Briefly, we collected 150 μl of the initial supernatant of each insect sample into a microplate well and evaporated the well contents for 120 min at room temperature until a volume of approximately 10 μl was reached. Then, 240 μl of anthrone reagent was added to each well and the plate was incubated for 15 min at room temperature and then heated 15 min at 90°C in a water bath, after which the absorbance of the samples was read at 625 nm using D-glucose as the standard. Triglycerides were assayed using the EnzyChrom™ Triglyceride Assay Kit (cat. no. ETGA-200), following the manufacturer's instructions.

We expected that body stores will not be proportional to insect mass (i.e., allometric relationships between body stores and mass). Therefore, data analyses were performed on the body store mass (mg) adjusted by allometric relationships computed for each type of body stores (proteins, free carbohydrates, glycogen, and triglycerides). For pooled replicates (i.e., small species), the mean individual body store mass was used.

3 | **DATA ANALYSES**

In our analyses, each body store was considered individually or pooled by fuels (glycogen, triglyceride and free carbohydrate) or total body store (fuels + proteins). As proteins are the main constituent of insect muscles (Vigoreaux, 2006) and flight muscles can constitute up to 65% of the body mass (Marden, 2000), we used protein content as a proxy for flight muscle. A larger quantity of flight muscle is associated with a better flight endurance (Marden, 2000).

Three explanatory factors of body stores were considered: (1) insect order; (2) mating system, including sex and swarming behaviour; and (3) dispersal strategy, including passive and active dispersal according to the DISPERSE database (Sarremejane et al., 2020). We ran tests to check for a possible effect of sampling sites on the quantity of body stores (see Appendix S1).

3.1 | **Allometric analyses**

Analysis of allometry followed the procedure proposed by Warton et al. (2006). To determine the relationship between body stores and body mass, we conducted standardised major axis (SMA) regression (model 2 regression) with the "smatr" package "sma" function (Warton et al., 2012). SMA is a least squares method (the regression line is estimated by minimising the sum of squares of residuals) but unlike linear regressions, the direction in which residuals are measured is not vertical. The SMA is the line that minimises the sum of squares of the shortest distances from the data points to the line, calculated on standardised data, then rescaled to the original axes (Warton et al., 2006). To use independent data, the analyses were carried out between the body store categories as response variables and the remaining body mass (i.e., the individual dry mass minus the measured body store mass) as explanatory

sampled, and the numbers in brackets indicate how many replicates per site. Che, Chênelais; Guy, Guyoult; Her, Petit Hermitage; Gue, Gueuche; Elo, Elorn; Con, Ruisseau de Condat; Arz, Arz; Loc, Le Loc'h; sampled, and the numbers in brackets indicate how many replicates per site. Che, Chênelais; Guy, Guyoult; Her, Petit Hermitage; Gue, Gueuche; Elo, Elorn; Con, Ruisseau de Condat; Arz, Arz; Loc, Le Loc'h; Mix, pooled replicates of individuals from different sites. The column "Dispersal strategies" is based on the DISPERSE database (see Sarremejane et al. (2020) for details). Rows correspond to the taxa Mix, pooled replicates of individuals from different sites. The column "Dispersal strategies" is based on the DISPERSE database (see Sarremejane et al. (2020) for details). Rows correspond to the taxa considered in analyses. "-" means identification was not performed. considered in analyses. "-" means identification was not performed.

^aNumber of individuals per replicate can be two or five depending on the replicate. e Number of individuals per replicate can be two or five depending on the replicate.

 GERBER ET AL. *BERBER ET AL.* **BERBER ET AL. BERGER ET A**

variable, both logarithmically transformed. The "sma" function allows us to calculate the regression lines between body stores and the remaining body mass for taxonomic groups. As the data are logarithmically transformed, the slope *b* of the regression line is equal to the allometric coefficient (see Peig & Green, 2009, for details). If the slope $b = 1$, the link between body stores and mass is isometric (i.e., increasing proportionally with the absolute increase in size). If *b > 1*, light individuals have relatively less body stores than heavy individuals (hyper-allometry). If *b < 1*, light individuals have relatively more body stores than heavy individuals (hypo-allometry).

In order to investigate the allometry intertaxa, a first series of regressions was performed on the mean by taxonomic groups to get the coefficient *b* for each body stores. The overall body store slope was compared to metabolic rate coefficients (2/3 and 3/4) by testing the correlation between residual and fitted values. Then, in a second series of regressions, we tested for differences in slopes between orders (using the mean of body stores by taxonomic groups) and between sexes (using the mean of body stores by taxonomic groups AND sex), using the Bartlett-corrected likelihood ratio statistic. When differences were detected, pair-wise comparison tests were run. When no differences were detected (i.e., there is a

common slope), between-order and between-sex differences were tested by calculating the differences between the regression lines for elevations (i.e., if significant, regression lines are separated by a gap indicating differences in body store quantity for a same body mass) and for shifts (i.e., if significant, regression lines are separated along a parallel axis and indicate a difference in the body store quantity due to a difference in body mass; see Warton et al., 2006 for details). Taxa with only one male or one female (Table 1; Nemouridae and Chloroperlidae among Plecoptera; Polycentropodidae among Trichoptera) were excluded from the sex analyses. We also explored the allometry intrataxa by conducted a third series of SMA regression within the taxonomic groups (Table 2).

Data on each body store category were allometrically adjusted as follows, considering the mean of each body store by taxonomic group (*n* = 22; Table 1), or by taxonomic group and sex, when investigating the mating system. The allometrically adjusted ratio used is $\bar{Y}_i = Y_i / X_i^b$ (Albrecht et al., 1993): for each individual i, \bar{Y}_i is the allometrically adjusted body store ratio (which will be used in the further analyses); *Yi*, the mass of the considered body store and *Xi* the body mass. The allometric coefficient *b* (i.e., slope) was obtained from the first series of SMA regressions.

TABLE 2 Coefficients of standardised major axis (SMA) regression between body stores and the remaining mass for each taxonomic group

				Overall body stores		Fuel		Proteins	
Order	Family	Genus	Species	Slope	\boldsymbol{p}	Slope	\boldsymbol{p}	Slope	\boldsymbol{p}
Diptera	Chironomidae	Conchapelopia		0.70	0.004	-0.5	0.964	1.34	0.02
	Chironomidae	Micropsectra		1.21	0.001	-0.33	0.369	2.01	0
	Chironomidae	Phaenopsectra		0.55	0.112	0.29	0.912	2.29	0.122
	Chironomidae	Polypedilum	٠	0.54	0.008	-0.55	0.228	1.62	0.037
	Chironomidae	Rheotanytarsus	٠	0.55	0.19	0.28	0.256	2.18	0.05
	Empididae		÷,	1.02	0.034	0.84	0.144	1.7	0.692
	Sphaeroceridae			0.8	0.381	0.53	0.794	2.49	0.213
Ephemeroptera	Baetidae	Baetis		NA	NA	-0.62	0.24	NA	NA
	Ephemerellidae	Serratella	ignita	NA	NA	NA	NA	-1.79	0.881
	Leptophlebiidae	Habrophlebia	$\overline{}$	0.48	0.171	0.39	0.842	1.92	0.461
Plecoptera	Chloroperlidae		\blacksquare	NA	NA	NA	NA	2.21	0.29
	Perlodidae	Isoperla	grammatica	2.1	0.695	-0.38	0.071	3.73	0.671
	Leuctridae	Leuctra		1.1	0.006	0.32	0.011	2.82	0.019
	Nemouridae	Nemoura		0.71	0.001	0.14	0.064	1.58	0.004
Trichoptera	Glossosomatidae	Agapetus		0.7	0.605	0.61	0.115	2.13	0.877
	Hydropsychidae	Hydropsyche		0.56	0.019	0.78	0.214	0.64	0.021
	Lepidostomatidae	Lepidostoma	hirtum	2.37	0.401	1.1	0.804	5.58	0.258
	Limnephilidae	Limnephilus		NA	NA	NA	NA	-0.23	0.843
	Polycentropodidae	Polycentropus		0.82	0.018	0.23	0.746	1.33	0.016
	Psychomyiidae	Lype		NA	NA	0.19	0.942	NA	NA
	Rhyacophilidae	Rhyacophila	dorsalis	1.5	0.017	1.3	0.014	1.83	0.017
	Sericostomatidae	Sericostoma		1.26	0.139	0.86	0.013	2.18	0.425

Note: The columns "slope" represent allometric coefficient of the regression. The columns "*p*" give the significance of the regressions. Numbers in green represent significant linear regressions (i.e., *p <* 0.05).

6 WILEY-Freshwater Biology CONSERVING BY SERBER ET AL.

3.2 | **Principal component analysis of body stores**

A principal component analysis (PCA) was performed on adjusted ratios, with free carbohydrate, glycogen and triglyceride and proteins as variables using the "FactomineR" package (Lê et al., 2008). Five missing data (NA) were treated by the "estim_npcPCA" function of the "missMDA" package (Josse & Husson, 2016). The npc value (i.e., the number of components to be retained for the PCA) given by this function was 0, meaning that NAs had to be substituted by the mean value of each variable. As the data were normalised, NAs were replaced by zeros. The coordinates of the first two principal components were used as response variables, and the factors of the PCA projections (order and dispersal) were used as explanatory variables. Active and passive dispersals were combined in three modalities: active dispersal (affinity to active dis $persal = 2$ or 3, eight taxa), passive dispersal (affinity to passive $dispersal = 2$ or 3, four taxa) and weak dispersal (affinity to active and passive dispersals $= 0$ or 1, eight taxa). Shapiro-Wilk tests and Bartlett's tests were conducted to evaluate the assumptions of normality and homoscedasticity. When heteroscedasticity or nonnormality were detected, Kruskal–Wallis tests were performed and when assumptions were met, ANOVA tests were used. When necessary, Dunn's *post hoc* tests were performed using the "FSA" package to test pairwise differences (Ogle et al., 2021). The alpha levels were adjusted using the Holm–Bonferroni correction to correct for multiple comparisons.

3.3 | **Effect of mating system on body store patterns**

We tested the differences of body stores between sexes for swarming taxa and for non-swarming taxa with Wilcoxon–Mann–Whitney tests for paired sample. The same tests were used to compare body stores between sexes for Diptera and Trichoptera as the number of replicates for each order is low (>10). Ephemeroptera and Plecoptera did not have enough replicates (more than five) to make reliable betweensex comparisons. We also compared variation in body stores between males and females in the orders Diptera and Trichoptera.

3.4 | **Contribution of insect order, sex and dispersal on body store patterns**

In order to determine the relative contribution of each explanatory factor (order, mating system and dispersal strategy) to body store patterns, partitions of variance were performed using the "varpart" function in the "vegan" package (Oksanen et al., 2015). We used three explanatory matrices––one for each factor. As the "varpart" function does not deal with NAs and as flight ability data were not available for Sphaericidae (Diptera), this taxon was removed from the variance partitioning. A total of six partitions of variance were

performed, one for each body store, one by combining fuels and one combining all body stores. The analyses were conducted on the mean by taxa and sex. Significance of each explanatory factor was checked using ANOVA tests.

All statistical analyses were performed with R software v.4.1.1 (R Core Team, 2021).

4 | **RESULTS**

We sampled 556 merolimnic imagoes, representing 22 taxa of 18 families and four orders (Diptera, Ephemeroptera, Plecoptera and Trichoptera; Table 1).

4.1 | **Allometry of body stores**

By considering all body stores together (sum of glycogen, triglycerides, free carbohydrates and proteins), a significant hypo-allometry was detected ($b = 0.70$; Figure 1a). Light individuals had relatively more body stores than heavy individuals. The allometry coefficients did not differ from the 3/4 nor $2/3$ metabolic rates ($p = 0.418$, *r* = 0.19, *df* = 15 and *p =* 0.481, r = 0.21, *df* = 15, *n =* 17, respectively).

Owing to the small number of Ephemeroptera (three taxa), their allometric coefficients could not be statistically compared with other orders (i.e., Diptera, Plecoptera and Trichoptera). For Plecoptera, only proteins were significantly correlated with the remaining mass of individuals. Consequently, the protein allometric coefficients of Plecoptera, Diptera and Trichoptera were compared, but the fuels (the sum of glycogen, triglycerides and free carbohydrates) and overall body store coefficients were compared between Diptera and Trichoptera only.

For both overall body stores and fuels, Diptera and Trichoptera shared the same allometric relationships (common slope *b =* 0.86; *p =* 0.608, likelihood ratio statistic = 0.99, *df* = 2, *n* = 13 and *b = 0*.63; *p =* 0.196, likelihood ratio statistic = 3.26, *df* = 2, *n =* 14, respectively). Diptera had more overall body stores for a given mass than Trichoptera (higher elevation: $p = 0.044$, Wald statistic = 4.05, $df = 1$; lower shift: $p = 0.001$, Wald statistic = 10.88, *df* = 1; Figure 1a). No difference in elevation was detected for fuels (*p = 0*.685, Wald statistic = 0.16, *df =* 1, *n =* 14), but Trichoptera had an absolute higher fuel content than Diptera, as a consequence of their higher mass, (higher shift, *p <* 0.001, Wald statistic = 12.45, *df =* 1, *n =* 14; Figure 1b).

For proteins, Diptera, Trichoptera and Plecoptera had the same allometric relationships (common slope *b =* 1.05, *p =* 0.411, likelihood ratio statistic = 1.78, $df = 2$, $n = 18$; Figure 1c). Differences in elevation were detected, together with a shift between Trichoptera and Diptera only. Diptera had more proteins for a given mass than Trichoptera (higher elevation: $p = 0.029$, pairwise test statistic = 2.59; lower shift: $p = 0.004$, pairwise test statistic = 10.99).

When investigating allometry intrataxa, nearly half of the taxonomic groups exhibited significant linear regressions for the overall

FIGURE 1 Standardised major axis (SMA) regression by order between the logarithms of body stores (mg) and the remaining body mass for the mean of each taxa. (a) SMA regression on all body stores, with the coloured lines for each order. Diptera and Trichoptera share a common slope ($b = 0.86$) and they differ in elevation (*p =* 0.044) and in shift (*p =* 0.001). (b) SMA regression on fuel content (i.e., sum of mass of glycogen, free carbohydrates and triglycerides). Diptera and Trichoptera share a common slope $(b = 0.63)$ and differ in shift ($p < 0.001$). (c) SMA regression on proteins. Diptera, Trichoptera and Plecoptera share a common slope ($b = 1.05$). Diptera and Trichoptera differ in elevation (*p =* 0.016) and in shift (*p =* 0.03). No significant correlations are observed for Ephemeroptera and for Plecoptera for regressions on the overall body stores and the fuels; therefore, the regression lines are not shown.

body stores and the proteins, whereas we found only three taxa with significant allometry for the fuel (*p* <0.05; Table 2). For the overall body stores, no allometric pattern was visible (i.e., the slope indicated either hypo-, iso- or hyper-allometry depending on the taxa). For the proteins, seven of eight taxa with significant repressions exhibited a hyper-allometry (i.e., light individuals have relatively less proteins than heavy individuals).

4.2 | **Site effect**

The site effect for overall body stores, fuels, proteins, triglycerides and free carbohydrates was tested as random term. For glycogen, the site effect was added directly as explanatory factor. Sites did not have a significant effect on body stores (Table 3). Although there is no overall effect, trends are observed on the dotplots (Figure 2),

Note: Mixed-effects models were used with adjusted body stores were used as response variables, taxa, sex and their interactions as explanatory variables, and site as random effect. For Glycogen, we used a generalised linear model with taxa, sex and their interactions, and site as explanatory variables.

FIGURE 2 Dotplot of the conditional modes of site effect as random term for each body stores. Conditional mode is the difference between the average predicted response and the response predicted for a particular individual. Blue dots, conditional values; black lines, conditional SDs.

especially for Triglycerides and Fuels. In particular, the "Le Loc'h" site differs from the "Guyoult" site. These differences could be explained by variations in taxonomic composition.

4.3 | **Relationship between body stores and insect order**

The first three principal components of the PCA (PC1, PC2 and PC3) explained 41.7%, 29.1% and 22.9% (respectively) of the total variance of body stores (Figure 3a). Free carbohydrate content contributed mainly to PC1 (34.3%). Adjusted triglyceride contributed mainly to PC2 (54.3%),

and adjusted glycogen to PC1 (23%), PC2 (29.8%) and PC3 (25.6%). Adjusted proteins contributed at 26.8% to PC1 and 62.4% to PC3. Diptera, Ephemeroptera, Trichoptera and Plecoptera separated along PC1, but only Diptera and Ephemeroptera differed significantly on PC1 (pairwise comparison Dunn test: $p = 0.026$, *Z*-statistic = 2.85; Figure 3b).

4.4 | **Relationship between body stores and dispersal**

comparison Dunn test: *p =* 0.005, *Z* = 3.12; Figure 3c).

TABLE 3 Significance of the site effect

for each body stores

Passive and weak dispersal significantly differed on PC1 (pairwise

FIGURE 3 Principal component analysis (PCA) of the merolimnic insect body stores. (a) Correlation circle showing correlations among body stores (Prot, proteins; Glyco, glycogen; free C., free carbohydrates; Trigly, triglycerides) and between body stores and PCA axes. Projections of insects on PCA axes according to (b) Insect orders (Diptera; Ephemeroptera; Plecoptera; Trichoptera) and (c) Dispersal affinity. Body stores are allometrically adjusted.

By testing the influence of dispersal strategies on each body store, we found that passive dispersers had significantly lower adjusted triglycerides than weak disperser (pairwise comparison Dunn test: $p = 0.050$, $Z = -2.40$). Active and weak dispersers had less adjusted glycogen than passive dispersers (pairwise comparison Dunn test: *p =* 0.013, *Z*= −2.85 and *p =* 0.053, *Z* = 2.22, respectively). Passive dispersers tended to have more proteins than active ones (pairwise comparison Dunn test: *p =* 0.053, *Z* = −2.22). However, when considering directly the adjusted overall body stores, fuels and free carbohydrates, no difference between dispersal strategies were detected.

4.5 | **Relationship between sex and body stores**

When considering the allometry between males and females, no significant differences in slope nor shift were detected for total body stores (Figure 4a). Overall, males had more fuel and triglycerides than females (difference in elevation: $b = 0.64$, $p < 0.001$, Wald statistic = 11.72, *df* = 1 [Figure 4b]; *b* = 0.59, *p* = 0.019, Wald statistic = 5.51, $df = 1$, $n = 38$ [Figure 4e], respectively). By testing swarming and non-swarming taxa separately, we found that males of a given taxon had more free carbohydrates and fuel than

their conspecific female, independently of their swarming behaviour (Wilcoxon–Mann–Whitney tests for paired sample, *p <* 0.05). However, for swarming taxa only, males of a given taxon had more triglycerides than their conspecific females (Wilcoxon–Mann– Whitney tests for paired samples, $p = 0.002$, V-statistic = 1).

Between-sex differences also were mediated by insect order. For Diptera and Trichoptera, females had less adjusted fuel, and free carbohydrates than males (Wilcoxon–Mann–Whitney tests for paired samples: *p* <0.05). However, female Diptera also had more adjusted proteins and less triglyceride than males (Wilcoxon–Mann– Whitney tests for paired sample, $p < 0.031$).

4.6 | **Contribution of insect order, mating system and dispersal strategy**

For the adjusted overall body stores, none of the explanatory factors were significant (ANOVA, *p* >0.05, *n* = 28). Thus, their relative contributions could not be interpreted. Mating system significantly explained 26% of the fuel variation (ANOVA: $p = 0.013$, $F = 4.29$, $df = 3$, $n = 28$; Figure 5a). Dispersal strategy is the only significant factor for explaining variation of adjusted triglycerides (30% of

FIGURE 4 Standardised major axis (SMA) regressions by sex between the logarithms of body stores (mg) and the remaining body mass for each individual. (a) Sum of all body stores; (b) Fuels––male and female differ in elevation (*p <* 0.001); (c) Free carbohydrates; (d) Glycogen; (e) Triglycerides–male and female differ in elevation (*p* = 0.019); (f) Proteins. No differences in slope nor shift

variance explained, ANOVA: *p* = 0.004, *F* = 6.84, *df* = 2, *n* = 28; Figure 5b) and the variation of adjusted glycogen (29% of variance explained, ANOVA: $p = 0.007$, $F = 6.59$, $df = 2$, $n = 28$; Figure 5e). It also explained 17% of the free carbohydrate variation (ANOVA: *p* = 0.033, *F* = 3.83 *df* = 2, *n* = 28; Figure 5d). Insect order explained 38% of the free carbohydrate variation (ANOVA: $p = 0.038$, $F = 6.61$ $df = 3, n = 28$.

Altogether, the three factors significantly explained the adjusted proteins (Figure 5c). Mating system explained 36% (ANOVA: *p =* 0.002, *F* = 6.11 *df* = 3, *n = 28*), dispersal strategies explained 30% $(ANOVA: p = 0.018, F = 4.55 df = 2, n = 28)$ and order explained 20% (ANOVA: *p =* 0.014, *F* = 4.95 *df* = 3, *n = 28*). Nevertheless, a large part of the variance remained unexplained (from 32% [Figure 5b] to 82% [Figure 5e]). Variation jointly explained by two or three factors (i.e., where the circles overlap) were not considered here and can be positive or negative, which explains why the sum of the variations for each factor do not always match with the total variance explained.

5 | **DISCUSSION**

In this study, we present a novel dataset on body stores of proteins, carbohydrates and triglycerides from >550 imagoes belonging to 22 taxa and four major orders of merolimnic insects, Diptera, Ephemeroptera, Plecoptera and Trichoptera. Firstly, we found that, according to their mass, small-bodied insect taxa stored a higher proportion of fuelling substrates than heavier insects, highlighting allometry between body stores and dry mass. This was particularly true for Trichoptera and Plecoptera, for which a hypo-allometry relationship was observed, indicating that light insects had relatively more energy stores than heavier insects. Previous studies stated that light insects consumed less energy during flight than heavy insects because they have a lower mass-specific flight metabolic rate (i.e., the rate of energy consumption in flight divided by the body mass; Niven & Scharlemann, 2005). Based on maximum power production relative to the power required to maintain steady flight (Hepburn et al., 1998), recent studies on bees showed that light individuals had more available power during flight than large ones (Duell, 2018; Grula et al., 2021; Helm et al., 2021), suggesting that body stores of light insects might be reduced without affecting their flight ability. Yet, we observed the opposite pattern, suggesting that allometric variation in overall body stores is not only linked to the energetic cost of flight for merolimnic taxa.

When considering each body store independently, light insects exhibited relatively more free carbohydrates and triglycerides than heavy insects. For glycogen and proteins, the relationship between body stores and size was close to isometric: glycogen and proteins increase proportionally with body size. Generally, fuel stored is greater for small insects and the quantity of muscle (i.e., proteins) remained proportional to the size. Interestingly, the resting metabolic rate (i.e.,

FIGURE 5 Analyses of variance partitioning from allometrically adjusted body stores showing the relative contribution of the effect of the mating system (in yellow), dispersal mode by flight (i.e., active or passive) (in blue) and order (in orange) in explaining body store strategies. Values are the proportion of variance explained by each factor, including the fractions shared with other factors (i.e., where the circles overlap). Values of the fractions shared with other factors are not shown. Residuals represent the unexplained variation. (a) Fuels, (b) Triglycerides, (c) Proteins, (d) Free carbohydrates and (e) Glycogen. Each number on the coloured section indicates the explained variance (%) by each combination of variables. Residuals show the unexplained variance. Values lower than 0% are considered as 0. Note: *When ANOVA tests are significant (*p <*0.05).

the rate of energy consumption at rest) has long been recognised to scale hypo-allometrically with mass. For insects, the resting metabolic rate scales to mass $0.66-0.9$, depending on the metabolic models (Harrison et al., 2014). Again, these results suggested that the relative increase of fuel in a light insect is not linked to the cost of flight, but rather to the relative increase of their resting metabolic rate.

We also looked at variation of allometry within each taxon. Even if the sampling did not enable us to highlight systematic allometric relationships among the body stores within our taxonomic groups, the results suggest that, within species, the protein contents increased relatively with size. As shown in different taxonomic groups (Green et al., 2001), the intraspecies allometry for proteins seemed to differ from the interspecies allometry.

When removing the allometry effect, we showed that Diptera had relatively more overall body stores, in particular more adjusted proteins, than Trichoptera. Insect order partly explains the body store patterns. It can be hypothesised that between-order variations result from the diversification of feeding habits from the ancestral group of Ephemeroptera, for which imagoes do not feed, to the more recently evolved Diptera that present a wide range of feeding habits (Courtney et al., 2017). Johnson et al. (2018) showed that diversification of feeding habits may strongly contribute to the diversification of insects, especially in Paraneoptera. The diversification of feeding resources may dramatically change the nutritional quality of food (Arrese & Soulages, 2010) and consequently change the body store patterns of merolimnic insects. Variations among

12 a b b CERBER ET AL. CERBER ET AL. CERBER ET AL. CERBER ET AL.

taxonomic groups also could be explained by environmental conditions. Two studies emphasised the potential role of environmental variation on triglycerides and glycogen contents for three species of Ephemeroptera (Koop et al., 2008; Winkelmann & Koop, 2007). In particular, water temperature, which is known to influence the size of the emerging adult, should be considered in further study of body store patterns (Lytle, 2002; Sweeney & Vannote, 1978).

We found differences in adjusted body stores between males and females. At comparable size, males contained more fuels, specifically more triglycerides, than females. Differences between males and females have been reported already in the literature. For example, Sartori et al. (1992) showed that males of *Siphlonurus aestivalis* (Ephemeroptera, Siphlonuridae) consumed lipids for their swarming flights whereas females did not use lipids for flight. Indeed, males of swarming species fly considerably longer than females, resulting in a greater energy consumption (Dudley, 2002; Lancaster & Downes, 2013). We found that males of swarming taxa contained more triglycerides than their conspecific females, contrary to non-swarming taxa. This suggests that the energy substrates used for the swarm would be mainly triglycerides. In particular, within Diptera males had more long-flight energy substrates (i.e., fuels, triglycerides) and energy availability (i.e., free carbohydrates circulating in the haemolymph directly available; Amat et al., 2012). Therefore, males should be able to fly longer since long flights first demand glycogen to initiate flight, then only lipids are subsequently required (Beenakkers et al., 1984; Chapman et al., 2013; Tigreros & Davidowitz, 2019). This ability to maintain flight longer could be used for swarming, as all the Diptera in our study can swarm. The high energy content found in males of swarming taxa and the high energy cost of maintaining flight suggest that body stores may be strongly driven by sexual selection. Thus, swarming behaviour could structure a part of the body stores, especially the triglycerides.

In our study, passive dispersers had relatively more proteins and glycogen (a fuel for flight) than weak dispersers and active dispersers. Mostly represented by small chironomids in our study, passive dispersers need to constantly beat their wings to maintain flight (i.e., they cannot glide to save energy power; Bhat et al., 2019). The highest proportion of glycogen (i.e., a common flight fuel for Diptera; Dudley, 2002) could help chironomids to extended flight and dispersal. According to Armitage (1995), the action of wind on swarming males may disperse populations away. In addition to reproductive success, we can hypothesise that the higher quantity of triglycerides in male Chironomidae could improve their dispersal ability. Nevertheless, chironomids colonise new habitats predominantly through the passive dispersal of fertilised females (Armitage, 1995). This result is corroborated by McLachlan (1985) who showed that females have wider wings than males, allowing extended flight. This is consistent with the high protein content found in chironomid females, associated with more muscle (Vigoreaux, 2006) and flight endurance (Marden, 2000).

We expected active dispersers to have more fuel than weak dispersers, but weak dispersers had relatively more triglycerides.

Potentially, triglycerides are allocated to other functions such as reproduction, but this has yet to be investigated. The typology of the dispersal strategies used here allowed us to analyse general trends of dispersal. However, these categories can be questioned on several aspects; for example, dispersal can vary within the same species depending on the population (Bonada & Dolédec, 2018). Even though these hypotheses need to be consolidated, our results highlight that physiological studies can help understand the conclusions of studies based solely on insect dispersal distances.

Furthermore, our study shows how body storage strategies could result in a trade-off between insect order, mating systems and dispersal. These factors did not explain the variation of the overall body stores. But each adjusted body store was partly explained by at least one factor (i.e., insect order, mating system and dispersal), thereby reinforcing the link between physiology and life-history traits. Interestingly, adjusted flight fuels, glycogen and triglycerides are well-explained by dispersal, which is in line with the importance of flight in the energy stores of insects (Amat et al., 2012). Moreover, we do not find the same patterns between overall flight fuels and each constituent (i.e., triglycerides, free carbohydrates and glycogen), suggesting the use of different types of body stores for reproduction depending on the species. Adjusted proteins are well-explained by combination of the mating system, the dispersal strategy and the insect order. It is congruent with the diversity of protein use, either fuel for flight (Bursell, 1963), muscle components (Marden, 2000) or fuel for reproduction (Chapman et al., 2013; Hahn et al., 2008). For example, we found that Diptera females had more adjusted proteins than males, highlighting an important role of proteins for reproduction (Chapman et al., 2013). Future work should be planned to identify the type of proteins and ascertain whether proteins serve as energy reserves for flight or reproduction or as constituents of muscle tissue.

Finally, we showed that a large part of variance of body stores remained unexplained (from 32% to 82% according to the type of body store), indicating that there are other influential variables that were not accounted for in this study. For instance, merolimnic insects differ in terms of voltinism (Bonada & Dolédec, 2018), life duration, period of emergence in the year (Kjær et al., 2021) and morphological traits, all of which can influence body stores. In addition to life-history traits, environmental factors might influence body store patterns within taxa (Shipley et al., 2012; Slansky Jr. & Haack, 1986), but also between taxa (Koop et al., 2008; Mathieu-Resuge et al., 2021). Very little information is available on the biology and ecology of merolimnic imagoes especially at the local scale where variability among populations can be high (Bonada & Dolédec, 2018).

6 | **CONCLUSIONS**

Our results on body storage strategies of emerging imagoes belonging to a large set of merolimnic insect families indicate that the body stores scaled allometrically with mass. This could be explained by the resting metabolic rate as it scales with insect body mass at the same magnitude. We also highlight a differentiation of body stores depending on order and sex, overall males had a greater amount of body stores than females. Finally, each body store is explained by a different life-history trait emphasising the need to choose which body store to measure depending on the question being addressed. Flight-related factors (i.e., mating system through swarm and dispersal) structured most of the body stores. To better investigate the body storage strategies of insects, future works should focus on identifying flight substrates and protein identification. Linking body stores of emerging merolimnic insects to the various facets of their ecology also is challenging.

AUTHOR CONTRIBUTIONS

Conceptualisation: BB, JM, CP. Developing methods: RG, BB, JM, CP. Conducting the research: LC, RG, CP. Data analysis: LC, RG. Data interpretation LC, RG, BB, JM, CP. Preparation figures & tables: LC, RG. Writing LC, RG, BB, JM, CP.

ACKNOWLEDGMENT

We particularly thank: all the people involved during the citizen science program (Lionel Picard of "GRoupe d'ETude des Invertébrés Armoricains" [http://www.gretia.org/\)](http://www.gretia.org/); Anthony le Gall of "Maison de la Rivière" [\(https://www.maison-de-la-riviere.com;](https://www.maison-de-la-riviere.com) Sizun); Jean-Pierre Pichard and Romain Bazire of "Réserve naturelle des étangs du Petit et du Grand Loc'h" (Guidel, [https://www.chasserenbretag](https://www.chasserenbretagne.fr/fdc56/) [ne.fr/fdc56/\)](https://www.chasserenbretagne.fr/fdc56/); Sébastien Gautier (with Matthieu and Pauline's help) of "Office français de la biodiversité" (<https://ofb.gouv.fr/>); as well as all those motivated but prevented by the COVID-19 epidemic. We thank anonymous reviewers and the handling editor for helpful comments and corrections on earlier drafts that significantly improved the final version of this article.

FUNDING INFORMATION

This research was funded the Earth Sciences and Astronomy Observatory of Rennes (OSUR, grant 2019) and the LTSER Zone Atelier Armorique (DIPATE project 2020).

DATA AVAILABILITY STATEMENT

Data are available from the authors upon reasonable request.

ORCID

Rémi Gerber <https://orcid.org/0000-0002-5520-9747> *Jean-Marc Rousse[l](https://orcid.org/0000-0002-6301-3234)* <https://orcid.org/0000-0002-6301-3234> *Benjamin Bergerot* <https://orcid.org/0000-0003-4313-4925>

REFERENCES

- Albrecht, G. H., Gelvin, B. R., & Hartman, S. E. (1993). Ratios as a size adjustment in morphometrics. *American Journal of Physical Anthropology*, *91*, 441–468. [https://doi.org/10.1002/ajpa.13309](https://doi.org/10.1002/ajpa.1330910404) [10404](https://doi.org/10.1002/ajpa.1330910404)
- Amat, I., Besnard, S., Foray, V., Pelosse, P., Bernstein, C., & Desouhant, E. (2012). Fuelling flight in a parasitic wasp: Which energetic substrate to use? *Ecological Entomology*, *37*, 480–489.

 GERBER ET AL. *IS* GERBER ET AL. **IS** $\begin{bmatrix} 13 & 13 \\ 14 & 15 & 16 \end{bmatrix}$

- Andrade, V. S., Wiegand, C., Pannard, A., Gagneten, A. M., Pédrot, M., Bouhnik-Le, C. M., & Piscart, C. (2020). How can interspecific interactions in freshwater benthic macroinvertebrates modify trace element availability from sediment? *Chemosphere*, *245*, 125594.
- Armitage, P. D. (1995). Behaviour and ecology of adults. In P. D. Armitage, P. S. Cranston, & L. C. V. Pinder (Eds.), *The Chironomidae: Biology and ecology of non-biting midges*, (pp. 194–224). Springer.
- Arrese, E. L., & Soulages, J. L. (2010). Insect fat body: Energy, metabolism, and regulation. *Annual Review of Entomology*, *55*, 207–225.
- Bacca, R. C., Pires, M. M., Moreira, L. F. B., Stenert, C., & Maltchik, L. (2021). The role of environmental and spatial factors in the assembly of aquatic insect communities in southern Brazilian temporary ponds. *Austral Ecology*, *46*, 228–238.
- Balian, E. V., Segers, H., Martens, K., & Lévêque, C. (2007). An introduction to the freshwater animal diversity assessment (FADA) project. *Hydrobiologia*, *595*, 3–8.
- Bay, E. C. (1993). Chironomid (Diptera: Chironomidae) larval occurrence and transport in a municipal water system. *Journal of the American Mosquito Control Association*, *9*, 275–284.
- Beenakkers, A. M. T., Van der Horst, D. J., & Van Marrewijk, W. J. A. (1984). Insect flight muscle metabolism. *Insect Biochemistry*, *14*, 243–260. [https://doi.org/10.1016/0020-1790\(84\)90057-X](https://doi.org/10.1016/0020-1790(84)90057-X)
- Bhat, S. S., Zhao, J., Sheridan, J., Hourigan, K., & Thompson, M. C. (2019). Aspect ratio studies on insect wings. *Physics of Fluids*, *31*, 121301. <https://doi.org/10.1063/1.5129191>
- Bonada, N., & Dolédec, S. (2018). Does the Tachet trait database report voltinism variability of aquatic insects between Mediterranean and Scandinavian regions? *Aquatic Sciences*, *80*, 1–11.
- Bradford, M. M. (1976). A rapid and sensitive method for the quantitation of microgram quantities of protein utilizing the principle of proteindye binding. *Analytical Biochemistry*, *72*, 248–254.
- Brittain, J. E. (1990). Life history strategies in Ephemeroptera and Plecoptera. In I. C. Campbell (Ed.), *Mayflies and stoneflies: Life histories and biology* (pp. 1–12). Springer.
- Brittain, J. E., & Sartori, M. (2009). Ephemeroptera:(mayflies). In V. H. Resh, & R. Cardé (Eds.), *Encyclopedia of insects* (pp. 328–334). Elsevier.
- Bursell, E. (1963). Aspects of the metabolism of amino acids in the tsetse fly, Glossina (Diptera). *Journal of Insect Physiology*, *9*, 439–452. [https://doi.org/10.1016/0022-1910\(63\)90054-4](https://doi.org/10.1016/0022-1910(63)90054-4)
- Burtt, E. T., Perry, R. J. O., & McLachlan, A. J. (1986). Feeding and sexual dimorphism in adult midges (Diptera: Chironomidae). *Ecography*, *9*, 27–32.
- Cadmus, P., Pomeranz, J. P., & Kraus, J. M. (2016). Low-cost floating emergence net and bottle trap: Comparison of two designs. *Journal of Freshwater Ecology*, *31*, 653–658.
- Chapman, R. F., Simpson, S. J., & Douglas, A. E. (2013). *The insects: Structure and function* (5th ed.). Cambridge University Press.
- Chen, H., Li, Z., Bu, S. H., & Tian, Z. Q. (2011). Flight of the Chinese white pine beetle (coleoptera: Scolytidae) in relation to sex, body weight and energy reserve. *Bulletin of Entomological Research*, *101*, 53–62.
- Compton, S. (2002). Sailing with the wind: Dispersal by small flying insects. In J. M. Bullock, R. Kenward, & R. Hails (Eds.), *Dispersal Ecology: 42nd Symposium of the British Ecological Society* (1st ed., pp. 113–133). Blackwell Pub.
- Courtney, G. W., & Cranston, P. S. (2015). Order Diptera. In J. H. Thorp, & D. C. Rogers (Eds.), *Thorp and Covich's Freshwater Invertebrates* (4th ed., pp. 1043–1058). Academic Press.
- Courtney, G. W., Pape, T., Skevington, J. H., & Sinclair, B. J. (2017). Biodiversity of Diptera. *Insect Biodiversity: Science and Society*, *1*, 229–278.
- David, G., Giffard, B., Halder, I., van Piou, D., & Jactel, H. (2015). Energy allocation during the maturation of adults in a long-lived insect: Implications for dispersal and reproduction. *Bulletin of Entomological Research*, *105*, 629–636.

14 [|] GERBER et al.

- de Figueroa, J. M. T., & Sánchez-Ortega, A. (1999). Imaginal feeding of certain Systellognathan stonefly species (Insecta: Plecoptera). *Annals of the Entomological Society of America*, *92*, 218–221.
- de Figueroa, J. M. T., & Sánchez-Ortega, A. (2000). Imaginal feeding of twelve Nemouroidean stonefly species (Insecta, Plecoptera). *Annals of the Entomological Society of America*, *93*, 251–253.
- Didham, R. K., Blakely, T. J., Ewers, R. M., Hitchings, T. R., Ward, J. B., & Winterbourn, M. J. (2012). Horizontal and vertical structuring in the dispersal of adult aquatic insects in a fragmented landscape. *Fundamental and Applied Limnology-Archiv furHydrobiologie*, *180*, $27 - 40.$
- Dijkstra, K.-D. B., Monaghan, M. T., & Pauls, S. U. (2014). Freshwater biodiversity and aquatic insect diversification. *Annual Review of Entomology*, *59*, 143–163. [https://doi.org/10.1146/annurev](https://doi.org/10.1146/annurev-ento-011613-161958)[ento-011613-161958](https://doi.org/10.1146/annurev-ento-011613-161958)
- Downer, R. G. H., & Matthews, J. R. (1976). Patterns of lipid distribution and utilisation in insects. *American Zoologist*, *16*, 733–745.
- Downes, J. A. (1969). The swarming and mating flight of Diptera. *Annual Review of Entomology*, *14*, 271–298. [https://doi.org/10.1146/annur](https://doi.org/10.1146/annurev.en.14.010169.001415) [ev.en.14.010169.001415](https://doi.org/10.1146/annurev.en.14.010169.001415)
- Dudley, R. (2002). *The biomechanics of insect flight: Form, Function, Evolution*. Princeton, New Jersey: Princeton University Press.
- Duell, M. E. (2018). *Matters of size: Behavioral, morphological, and physiological performance scaling among stingless bees (Meliponini)*. Arizona State University.
- Foray, V., Pelisson, P.-F., Bel-Venner, M.-C., Desouhant, E., Venner, S., Menu, F., Giron, D., & Rey, B. (2012). A handbook for uncovering the complete energetic budget in insects: The van Handel's method (1985) revisited. *Physiological Entomology*, *37*, 295–302. [https://doi.](https://doi.org/10.1111/j.1365-3032.2012.00831.x) [org/10.1111/j.1365-3032.2012.00831.x](https://doi.org/10.1111/j.1365-3032.2012.00831.x)
- Green, A. J., Figuerola, J., & King, R. (2001). Comparing interspecific and intraspecific allometry in the Anatidae. *Journal für Ornithologie*, *142*, 321–334. <https://doi.org/10.1046/j.1439-0361.2001.00075.x>
- Grula, C. C., Rinehart, J. P., Greenlee, K. J., & Bowsher, J. H. (2021). Body size allometry impacts flight-related morphology and metabolic rates in the solitary bee Megachile rotundata. *Journal of Insect Physiology*, *133*, 104275. <https://doi.org/10.1016/j.jinsphys.2021.104275>
- Gullefors, B., & Petersson, E. (1993). Sexual dimorphism in relation to swarming and pair formation patterns in leptocerid caddisflies (Trichoptera: Leptoceridae). *Journal of Insect Behavior*, *6*, 563–577. <https://doi.org/10.1007/BF01048123>
- Hahn, D. A., James, L. N., Milne, K. R., & Hatle, J. D. (2008). Life-history plasticity after attaining a dietary threshold for reproduction is associated with protein storage in flesh flies. *Functional Ecology*, *22*, 1081–1090. <https://doi.org/10.1111/j.1365-2435.2008.01451.x>
- Harker, J. E. (1992). Swarm behaviour and mate competition in mayflies (Ephemeroptera). *Journal of Zoology*, *228*, 571–587.
- Harrison, S. S. C., & Dobson, M. (2008). Evolutionary drivers and the ecological traits of adult aquatic insects. In J. Lancaster, & R. A. Briers (Eds.), *Aquatic Insects: Challenges to Populations* (pp. 250–267). CABI.
- Harrison, J. F., Klok, C., & Waters, J. S. (2014). Critical PO2 is sizeindependent in insects: Implications for the metabolic theory of ecology. *Current Opinion in Insect Science*, *4*, 54–59. [https://doi.](https://doi.org/10.1016/j.cois.2014.08.012) [org/10.1016/j.cois.2014.08.012](https://doi.org/10.1016/j.cois.2014.08.012)
- Helm, B. R., Baldwin, M. A., Rinehart, J. P., Yocum, G. D., Greenlee, K. J., & Bowsher, J. H. (2021). Body and wing allometries reveal flightfecundity tradeoff in response to larval provisioning in Osmia lignaria (Hymenoptera: Megachilidae). *Journal of Insect Science*, *21*, 11. <https://doi.org/10.1093/jisesa/ieab035>
- Hepburn, H. R., Radloff, S. E., Steele, G. R., & Brown, R. E. (1998). Dimensional aspects of flight in the honey bees of Africa. *Journal of Apicultural Research*, *37*, 147–154. [https://doi.org/10.1080/00218](https://doi.org/10.1080/00218839.1998.11100966) [839.1998.11100966](https://doi.org/10.1080/00218839.1998.11100966)
- Holzenthal, R. W., Thomson, R. E., & Ríos-Touma, B. (2015). Order Trichoptera. In J. H. Thorp, & D. C. Rogers (Eds.), *Thorp and Covich's freshwater invertebrates* (pp. 965–1002). Elsevier.
- Hynes, H. B. N. (1976). Biology of plecoptera. *Annual Review of Entomology*, *21*, 135–153.
- Ishiwata, K., Sasaki, G., Ogawa, J., Miyata, T., & Su, Z.-H. (2011). Phylogenetic relationships among insect orders based on three nuclear protein-coding gene sequences. *Molecular Phylogenetics and Evolution*, *58*, 169–180.
- Jackson, J. K. (1988). Diel emergence, swarming and longevity of selected adult aquatic insects from a Sonoran Desert stream. *American Midland Naturalist*, *119*, 344–352.<https://doi.org/10.2307/2425817>
- Jacobus, L. M., Macadam, C. R., & Sartori, M. (2019). Mayflies (Ephemeroptera) and their contributions to ecosystem services. *Insects*, *10*, 170.
- Johnson, K. P., Dietrich, C. H., Friedrich, F., Beutel, R. G., Wipfler, B., Peters, R. S., Allen, J. M., Petersen, M., Donath, A., Walden, K. K. O., Kozlov, A. M., Podsiadlowski, L., Mayer, C., Meusemann, K., Vasilikopoulos, A., Waterhouse, R. M., Cameron, S. L., Weirauch, C., Swanson, D. R., … Yoshizawa, K. (2018). Phylogenomics and the evolution of hemipteroid insects. *Proceedings of the National Academy of Sciences of the United States of America*, *115*, 12775–12780.
- Josse, J., & Husson, F. (2016). missMDA: A package for handling missing values in multivariate data analysis. *Journal of Statistical Software*, *70*, 1–31.
- Kjær, C., Sørensen, P. B., Wiberg-Larsen, P., Bak, J., Bruus, M., Strandberg, B., Larsen, S. E., Rasmussen, J. J., & Strandberg, M. (2021). Vulnerability of aquatic insect species to insecticides, depending on their flight period and adult life span. *Environmental Toxicology and Chemistry*, *40*, 1778–1787.
- Koop, J. H. E., Schaeffer, M., Ortmann, C., & Winkelmann, C. (2008). Towards environmental assessment of river ecosystems by analyzing energy reserves of aquatic invertebrates. *Limnologica*, *38*, 378– 387.<https://doi.org/10.1016/j.limno.2008.05.004>
- Kovats, Z. E., Ciborowski, J. J. H., & Corkum, L. D. (1996). Inland dispersal of adult aquatic insects. *Freshwater Biology*, *36*, 265–276.
- Lancaster, J., & Downes, B. J. (2013). *Aquatic entomology*. Oxford University Press.
- Lê, S., Josse, J., & Husson, F. (2008). FactoMineR: An R package for multivariate analysis. *Journal of Statistical Software*, *25*, 1–18.
- Lytle, D. A. (2002). Flash floods and aquatic insect life-history evolution: Evaluation of multiple models. *Ecology*, *83*, 370–385. [https://doi.](https://doi.org/10.1890/0012-9658(2002)083%5B0370:FFAAIL%5D2.0.CO;2) [org/10.1890/0012-9658\(2002\)083\[0370:FFAAIL\]2.0.CO;2](https://doi.org/10.1890/0012-9658(2002)083%5B0370:FFAAIL%5D2.0.CO;2)
- Maino, J. L., & Kearney, M. R. (2014). Ontogenetic and interspecific metabolic scaling in insects. *The American Naturalist*, *184*, 695–701. <https://doi.org/10.1086/678401>
- Malik, P., Hegedüs, R., Kriska, G., & Horváth, G. (2008). Imaging polarimetry of glass buildings: Why do vertical glass surfaces attract polarotactic insects? *Applied Optics*, *47*, 4361–4374. [https://doi.](https://doi.org/10.1364/ao.47.004361) [org/10.1364/ao.47.004361](https://doi.org/10.1364/ao.47.004361)
- Marden, J. H. (2000). Variability in the size, composition, and function of insect flight muscles. *Annual Review of Physiology*, *62*, 157–178. <https://doi.org/10.1146/annurev.physiol.62.1.157>
- Martynov, A. (2016). The life cycles of Mayflies (Insecta: Ephemeroptera) of the Eastern Ukraine. Second report. *Proceedings of the National Museum of Natural History*, *14*, 86–94. [https://doi.org/10.15407/](https://doi.org/10.15407/vnm.2016.14.086) [vnm.2016.14.086](https://doi.org/10.15407/vnm.2016.14.086)
- Mathieu-Resuge, M., Martin-Creuzburg, D., Twining, C., Parmar, T. P., Hager, H., & Kainz, M. (2021). Taxonomic composition and lake bathymetry influence fatty acid export via emergent insects. *Freshwater Biology*, *66*, 2199–2209.<https://doi.org/10.1111/fwb.13819>
- McLachlan, A. (1985). The relationship between habitat predictability and wing length in midges (Chironomidae). *Oikos*, *44*, 391. [https://](https://doi.org/10.2307/3565779) doi.org/10.2307/3565779
- Morse, J. C. (2009). Trichoptera (caddisflies). In V. H. Resh, & R. Cardé (Eds.), *Encyclopedia of insects* (pp. 1015–1020). Elsevier
- Muehlbauer, J. D., Collins, S. F., Doyle, M. W., & Tockner, K. (2014). How wide is a stream? Spatial extent of the potential "stream signature" in terrestrial food webs using meta-analysis. *Ecology*, *95*, 44–55.
- Müller-Peddinghaus, E. (2011). *Flight-morphology of Central European caddisflies (Insecta: Trichoptera) in relation to their ecological preferences*. PhD Thesis, Duisburg-Essen University.
- Nijboer, R. (2004). The ecological requirements of Agapetus fuscipes Curtis (Glossosomatidae), a characteristic species in unimpacted streams. *Limnologica*, *34*, 213–223. [https://doi.org/10.1016/](https://doi.org/10.1016/S0075-9511(04)80046-X) [S0075-9511\(04\)80046-X](https://doi.org/10.1016/S0075-9511(04)80046-X)
- Niven, J. E., & Scharlemann, J. P. W. (2005). Do insect metabolic rates at rest and during flight scale with body mass? *Biology Letters*, *1*, 346– 349.<https://doi.org/10.1098/rsbl.2005.0311>
- Ogle D. H., DollJ. C., Wheeler P. & Dinno A. (2021). FSA: Fisheries stock analysis. R package version 0.9.1. [https://github.com/](https://github.com/fish-R-Core-Team/FSA) [fish-R-Core-Team/FSA](https://github.com/fish-R-Core-Team/FSA)
- Oksanen J., Blanchet F.G., Kindt R., Legendre P., Minchin P.R., O'Hara R.B., Simpson, G. L., Solymos, P., Stevens, M. H. H., Wagner, H. H. (2015). *Vegan community ecology package: Ordination methods, diversity analysis and other functions for community and vegetation ecologists*. R package ver 2.3.5.
- Peig, J., & Green, A. J. (2009). New perspectives for estimating body condition from mass/length data: The scaled mass index as an alternative method. *Oikos*, *118*, 1883–1891. [https://doi.](https://doi.org/10.1111/j.1600-0706.2009.17643.x) [org/10.1111/j.1600-0706.2009.17643.x](https://doi.org/10.1111/j.1600-0706.2009.17643.x)
- Peredo Arce, A., Hörren, T., Schletterer, M., & Kail, J. (2021). How far can EPTs fly? A comparison of empirical flying distances of riverine invertebrates and existing dispersal metrics. *Ecological Indicators*, *125*, 107465. <https://doi.org/10.1016/j.ecolind.2021.107465>
- Petersen, I., Masters, Z., Hildrew, A. G., & Ormerod, S. J. (2004). Dispersal of adult aquatic insects in catchments of differing land use. *Journal of Applied Ecology*, *41*, 934–950.
- Petersson, E., & Hasselrot, A. T. (1994). Mating and nectar feeding in the psychomyiid caddis fly Tinodes waeneri. *Aquatic Insects*, *16*, 177–187.
- Phillipsen, I. C., Kirk, E. H., Bogan, M. T., Mims, M. C., Olden, J. D., & Lytle, D. A. (2015). Dispersal ability and habitat requirements determine landscape-level genetic patterns in desert aquatic insects. *Molecular Ecology*, *24*, 54–69.
- Pinder, L. C. V. (1986). Biology of freshwater Chironomidae. *Annual Review of Entomology*, *31*, 1–23.
- R Core Team. (2021). *R: A language and environment for statistical computing*. R Foundation for Statistical Computing.
- Raitif, J., Plantegenest, M., Agator, O., Piscart, C., & Roussel, J.-M. (2018). Seasonal and spatial variations of stream insect emergence in an intensive agricultural landscape. *Science of the Total Environment*, *644*, 594–601. <https://doi.org/10.1016/j.scitotenv.2018.07.021>
- Roma, G. C., Bueno, O. C., & Camargo-Mathias, M. I. (2010). Morphophysiological analysis of the insect fat body: A review. *Micron*, *41*, 395–401.
- Rúa, J., & de Figueroa, J. M. T. (2013). Adult feeding habits of three Perloidea species (Plecoptera: Perlidae and Chloroperlidae). *Aquatic Insects*, *35*, 99–104.
- Sarremejane, R., Cid, N., Stubbington, R., Datry, T., Alp, M., Cañedo-Argüelles, M., Cordero-Rivera, A., Csabai, Z., Gutiérrez-Cánovas, C., Heino, J., Forcellini, M., Millán, A., Paillex, A., Pařil, P., Polášek, M., Tierno de Figueroa, J. M., Usseglio-Polatera, P., Zamora-Muñoz, C., & Bonada, N. (2020). DISPERSE, a trait database to assess the dispersal potential of European aquatic macroinvertebrates. *Scientific Data*, *7*, 386. <https://doi.org/10.1038/s41597-020-00732-7>
- Sartori, M., & Brittain, J. E. (2015). Order Ephemeroptera. In J. H. Thorp, & D. C. Rogers (Eds.), *Thorp and Covich's freshwater invertebrates* (4th ed., pp. 873–891). Elsevier.
- Sartori, M., Keller, L., Thomas, A., & Passera, L. (1992). Flight energetics in relation to sexual differences in the mating-behavior of a mayfly, *Siphlonurus aestivalis*. *Oecologia*, *92*, 172–176.
- Shipley, M. M., Wellington, K., Rao, A., Ritchie, T., & Vogtsberger, R. C. (2012). Fatty acid composition of a burrowing mayfly, Hexagenia limbata (Ephemeroptera: Ephemeridae), from a north Central Texas

Lake. *Journal of the Kansas Entomological Society*, *85*, 245–258. <https://doi.org/10.2317/JKES120831.1>

- Slansky, F., Jr., & Haack, R. A. (1986). Age-specific flight behavior in relation to body weight and lipid content of Ips calligraphus reared in slash pine bolts with thick or thin inner bark (phloem). *Entomologia Experimentalis et Applicata*, *40*, 197–207. [https://doi.org/10.1111/](https://doi.org/10.1111/j.1570-7458.1986.tb00502.x) [j.1570-7458.1986.tb00502.x](https://doi.org/10.1111/j.1570-7458.1986.tb00502.x)
- Sullivan, R. T. (1981). Insect swarming and mating. *The Florida Entomologist*, *64*, 44–65.
- Sweeney, B. W., & Vannote, R. L. (1978). Size variation and the distribution of hemimetabolous aquatic insects: Two thermal equilibrium hypotheses. *Science*, *200*, 444–446. [https://doi.org/10.1126/scien](https://doi.org/10.1126/science.200.4340.444) [ce.200.4340.444](https://doi.org/10.1126/science.200.4340.444)
- Tigreros, N., & Davidowitz, G. (2019). Flight-fecundity tradeoffs in wingmonomorphic insects. In R. Jurenka (Ed.), *Advances in insect physiology* (Vol. *56*, pp. 1–41). Academic Press. [https://doi.org/10.1016/](https://doi.org/10.1016/bs.aiip.2019.02.001) [bs.aiip.2019.02.001](https://doi.org/10.1016/bs.aiip.2019.02.001)
- Tokeshi, M., & Reinhardt, K. (1996). Reproductive behaviour in Chironomus anthracinus (Diptera: Chironomidae), with a consideration of the evolution of swarming. *Journal of Zoology*, *240*, 103– 112. <https://doi.org/10.1111/j.1469-7998.1996.tb05488.x>
- Tonkin, J. D., Altermatt, F., Finn, D. S., Heino, J., Olden, J. D., Pauls, S. U., & Lytle, D. A. (2018). The role of dispersal in river network metacommunities: Patterns, processes, and pathways. *Freshwater Biology*, *63*, 141–163. <https://doi.org/10.1111/fwb.13037>
- Vallenduuk, H. J., & Moller Pillot, H. (2007). *Chironomidae Larvae of the Netherlands and Adjacent Lowlands: General Ecology and Tanypodinae*. KNNV Publishing.
- Vigoreaux, J. O. (2006). *Nature's versatile engine: Insect flight muscle inside and out*. Springer.
- Warton, D. I., Duursma, R. A., Falster, D. S., & Taskinen, S. (2012). Smatr 3– An R package for estimation and inference about allometric lines. *Methods in Ecology and Evolution*, *3*, 257–259. [https://doi.](https://doi.org/10.1111/j.2041-210X.2011.00153.x) [org/10.1111/j.2041-210X.2011.00153.x](https://doi.org/10.1111/j.2041-210X.2011.00153.x)
- Warton, D. I., Wright, I. J., Falster, D. S., & Westoby, M. (2006). Bivariate line-fitting methods for allometry. *Biological Reviews of the Cambridge Philosophical Society*, *81*, 259–291. [https://doi.](https://doi.org/10.1017/S1464793106007007) [org/10.1017/S1464793106007007](https://doi.org/10.1017/S1464793106007007)
- Winkelmann, C., & Koop, J. H. E. (2007). The management of metabolic energy storage during the life cycle of mayflies: A comparative field investigation of the collector-gatherer Ephemera danica and the scraper Rhithrogena semicolorata. *Journal of Comparative Physiology B-Biochemical Systemic and Environmental Physiology*, *177*, 119–128. <https://doi.org/10.1007/s00360-006-0114-7>
- Yeates, D. K., Meusemann, K., Trautwein, M., Wiegmann, B., & Zwick, A. (2016). Power, resolution and bias: Recent advances in insect phylogeny driven by the genomic revolution. *Current Opinion in Insect Science*, *13*, 16–23.

SUPPORTING INFORMATION

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

How to cite this article: Gerber, R., Cabon, L., Piscart, C., Roussel, J.-M., & Bergerot, B. (2022). Body stores of emergent aquatic insects are associated with body size, sex, swarming behaviour and dispersal strategies. *Freshwater Biology*, *00*, 1–15. <https://doi.org/10.1111/fwb.14003>