Drosophila glue protects from predation

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Animals can be permanently attached to a substrate in terrestrial environments at certain stages of their development. Pupa adhesion has evolved multiple times in insects and is thought to maintain the animal in a place where it is not detectable by predators. Here, we investigate whether pupa adhesion in Drosophila can also protect the animal by preventing potential predators from detaching the pupa. We measured the adhesion of Drosophila species sampled from the same area and found that pupa adhesion varies among species, which can be explained by different glue production strategies. Then, we compared attached and manually detached pupae in both field and laboratory assays to investigate the role of pupa adhesion to prevent predation. First, we found that attached pupae remain onsite 30% more than detached pupae in the field after 3 days, probably because they are less predated. Second, we observed that attached pupae are less efficiently predated by ants in the laboratory: they are not carried back to the ant nest and more ants are needed to consume them onsite. Our results show that pupa adhesion can prevent the animal from being taken away by predators and is crucial for Drosophila fly survival.

1. Background

Many animals are immobile and permanently attached to a substrate. Most of them are found in aquatic environments [1] and within living hosts for parasites. In terrestrial environments, animals can also be attached to a substrate during certain stages of development where no feeding from outside nutrients is required. Eggs from multiple invertebrate species are fixed on leaves, wood or on the host tissue in some parasitic species [2–4]. This is also the case in pupae, a non-feeding life stage of holometabolous insects between the larval and adult stages [5].

Modes of attachment and the ability to stick to the substrate appear to change rapidly during animal evolution. In stick and leaf insects, gluing eggs to a substrate has evolved independently seven times [6]. Wasp parasites have evolved different pupation strategies consisting in hanging their cocoon to a leaf (Meteorus pulchricomis) or attaching their cocoon on a leaf (Microplitis sp.) [7]. In flies, pupae of certain species such as Drosophila melanogaster and Phormia regina are glued to a substrate, whereas others such as Musca domestica, Calliphora erythrocephala or Sarcophaga falculata are not [8].

Permanent attachment of eggs and pupae has been associated with several functions. First, attachment can allow the organisms to remain in a favourable environment. Females of many butterfly species choose to lay and attach their eggs directly on the host plant on which their larvae will start feeding [3,9]. Butterflies laying their eggs during winter have evolved different strategies to avoid their eggs being blown away if the dead host leaf falls far away from the plant. They lay their eggs on herbal or wooden substrates near the host plant or they glue their eggs less strongly to the host leaf so that the eggs would detach from the dead leaf and fall close to the host plant in the case of strong wind...
we compared the ability of attached and detached pupae of species to remain on site in a natural environment. Finally, the ability of attached and detached pupae of one of these logical community to explore whether different species have of pupae from different substrate might have another, yet unexplored, effect against out of the soil [22], we hypothesized that pupa adhesion to a As ants were previously observed to dig up and carry pupae and crickets were identified as potential natural predators. L.05/010/dryad.x3fbb77h [30]. Combining isofemale lines of the same species when necessary, we managed to obtain mass culture for most species, including Drosophila hydei, Drosophila simulans and D. suzuki. These three stocks were raised in the laboratory for three to four months at 25°C before being used in the experiments described below.

2. Material and methods

(a) Fly culture

Flies were cultured at 25°C in plastic vials on standard medium (4 l: 83.5 g yeast, 335.0 g cornmeal, 40.0 g agar, 233.5 g saccharose, 67.0 ml Moldex, 6.0 ml propionic acid). For D. suzuki, this medium was supplemented with 200 g of r-glucose.

(b) Fly collection

Drosophila flies were collected at the Bois de Vincennes in Paris, France. On 3 July 2020, five traps made from 0.5 l plastic bottles were settled in the ornithological reserve (48°50'05.5" N; 2°26'11.4" E). Small holes were made in the sides of the bottles that allowed drosophilid flies to enter but prevented entry by larger insects. Traps were baited with pieces of banana and hung to tree branches or within the understorey vegetation. They were distantly placed, three of them in the forest, one at the margin of a meadow and another close to a pond (see [30]). Flies in the bottles were collected 2 and 5 days later. On 16 July 2020, drosophilid flies were also collected with a sweeping net over compost at the forest services facilities. Collected flies were transferred with an aspirator into plastic vials containing a piece of humid tissue paper for the time of transportation. In the laboratory, flies were checked under a binocular stereomicroscope and isolated by species in culture vials with standard cornmeal. When species could not be precisely determined, females were isolated in small culture vials with instant Drosophila medium (Formula 4–24, Carolina Biological Supply Company, Burlington, NC, USA) and species were then identified based on key morphological characters in the male progeny. Results of the fly collections are available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.x3fbb77h [30]. Combining isofemale lines of the same species when necessary, we managed to obtain mass culture for most species, including Drosophila hydei, Drosophila simulans and D. suzuki. These three stocks were raised in the laboratory for three to four months at 25°C before being used in the experiments described below.

(c) Adhesion assays

Third instar wandering larvae were transferred on glass slides (Menzel Superfrost microscope glass slide, ThermoScientific™ no. AGAB00080) with soft forceps and kept in a box with wet cotton. Fifteen to 21 h after transfer, pupae naturally attached to the glass slides with their ventral part adhering to the glass slide were used for the adhesion tests. The pull-off force necessary to detach the pupa from the glass slide was measured using a universal test machine (LS15/H/230 V Lloyd Instruments) with a 5 N force sensor (YL-LC-005-A1 Lloyd Instruments), in a set-up similar to the one published earlier [31]. Double-sided adhesive tape (tesa, extra strong, no. 05681-00018) was attached to a cylindrical metal part in contact with the force sensor. The force was set to 0 before each run. The force sensor was moved down with a constant speed of 1 mm min⁻¹ until it pressed the pupa with a force of 0.07 N (0.25 N for D. hydei) and stiller at a force of 0.03 N (0.21 N for D. hydei) for 10 s and finally moved up with a constant speed of 0.2 mm s⁻¹ until the pupa was detached. Force-by-time curves were recorded using NEXYGENPlus soft- ware (Lloyd Instruments). We used the maximal force reached during the experiment, corresponding to the force at which the pupa was detached, as the adhesion force of the individual. Pupae whose pupal case broke during the assay (D. suzuki: 1 of 27, D. simulans: 2 of 37; D. hydei: 7 of 50) or pupae which were not detached (D. suzuki: 0 of 27, D. simulans: 5 of 37, D. hydei: 12 of 50) were excluded from the analysis. After pupa detachment, images of glue prints remaining on glass substrates were taken with a Keyence VHX-2000 Z20 ×20 or ×100. Contours of prints areas were digitized manually by the same person using IMAGEJ (1.50d, java 1.8.0_212, 64-bit) [32]. Pictures were anonymized for manual contour acquisition so that the digitizer did not know the genotype. We measured the surface of the print corresponding to the pupa–substrate interface as defined previously [31]. Three prints for D. suzuki and one print for D. simulans were not detectable on the slides and were not used in the analysis.

To assess pupal size, we used pupae that were raised in the same conditions as for the adhesion tests but that were not used for the tests. We imaged attached pupae from the dorsal view and measured the area of the pupal case as described to measure the glue area.

(d) Predation assay in the field

In the laboratory, D. simulans third instar wandering larvae were collected with soft forceps and let to pupate in Petri dishes
(55 mm diameter) in a closed plastic box containing wet paper for 17–24 h at room temperature. Fifteen larvae were placed in each dish. All Petri dishes were then brought to the field. For the condition ‘attached pupae’, a few pupae were removed in order to have exactly 10 pupae glued in the lid of the Petri dish. For the condition ‘detached pupae’, pupae were all detached from the dish with featherweight entomology forceps and exactly 10 detached pupae were kept in the lid. To distinguish between the two conditions (‘non collées’ versus ‘collées’ in French), the letters ‘NC’ or ‘C’ were written on pieces of paper that were taped on the external surface of the lids to facilitate visualization and counting of the pupae. The lids of the Petri dishes were put in the centre of buckets previously installed in the ornithological reserve of Bois de Vincennes (see [30]). These 40 × 35 cm buckets contain local soil, leaf litter and vegetation; they are pierced at the bottom for water drainage and are semi-buried (10 cm deep) [33]. In total, 28 buckets were used and contained both one dish with ‘attached pupae’ and one dish with ‘detached pupae’ (56 Petri dishes in total). We randomly alternated the west/east orientation of the two conditions inside the buckets. Pupae were counted at 0 h (8 September 2020, day 1 morning, at 11.00), 6 h30 (day 1 afternoon), 22 h30 (day 2 morning), 31 h (day 2 afternoon), 47 h (day 3 morning) and 54 h (day 3 afternoon) after the start of the experiment without being touched. Animals present in the dishes at counting times were photographed and later identified based on the pictures (see [30]). Insects present in the dishes at the end of recording (54 h) were collected and kept in 90% ethanol for identification.

(e) Ant predation assay in the laboratory

Seven colonies of Temnothorax nylanderi were collected on 17 September 2020 in Bois de Vincennes (48°50’20.0” N 2°26’57.2” E), brought to the laboratory and allowed to move into artificial nests consisting of two microscope glass slides separated by a 1 mm auto-adhesive plastic foam harbouring three chambers, covered with a black plastic film to maintain darkness. Each artificial nest was placed in a foraging area consisting of a plastic box (11.5 × 11.5 × 5.5 cm) as described in [33]. Water was always available in a tube plugged with cotton. Colonies were fed frozen Drosophila and diluted honey and then they were starved for 10 days until the beginning of the experiment, on 13 October 2020. Prior and during the experiment, colonies were kept at room temperature on the bench and under indirect sunlight. Each day at around 9:30, and for 6 days (between 13 October and 21 October 2020), one glass slide presenting two pupae was put into the foraging area of each colony, at about 10 cm from the entrances of the artificial nest. On each day, half of the colonies were given one slide with two D. simulans pupae and the other half one slide with two D. suzukii pupae, and we alternated species every day. We used the same two Drosophila lines as for our adhesion assays described above.

The glass slides were prepared as follows. Six third instar wandering larvae were transferred on glass slides (Menzel Superfrost microscope glass slide, ThermoScientific™ no. AGAB000080) placed in a Petri dish kept in a box containing wet cotton and left to pupate for 14–19 h. On each slide, only one pupa was kept attached, the other ones were detached slightly with soft forceps and one detached pupa was left on the slide. Pupae were about 1 cm apart from each other on the slide. Left/right positions of the two pupae were randomly assigned per colony and changed every day. The initial locations of the pupae were identified by a mark under the slide.

Ant foraging areas were checked by eye every 5 min for 3 h after the glass slide with the two pupae was added. The number of ants in contact with the pupae was counted. If the pupa was brought to the nest, we noted the time when the pupa was present inside the nest for the first time. If the pupa was not brought to the nest, we noted the time when the pupa was fully consumed by the ants (no Drosophila body remnants visible by eye).

(f) Quantification and statistical analysis

We used R v. 3.6.1 (https://www.r-project.org) to conduct our statistical analyses. Adhesion forces were not normally distributed and statistical differences in forces between species were tested by Kruskal–Wallis tests followed by multiple pairwise Wilcoxon tests. To test whether adhesion forces were correlated to pupa–substrate contact areas for each species, we performed standardized major axis regressions using the sma function from the rsmatr-package in R [34]. Slopes were compared among species. For predation assays in Bois de Vincennes and in the laboratory, means were compared using the Wilcoxon tests because data were not normally distributed. In order to avoid a potential bucket effect on pupa disappearance in the predation assay in the field, we paired both ‘attached pupae’ and ‘detached pupae’ conditions in each bucket and we subsequently used paired tests. As the number of detached pupae decreases towards the end of the experiment, predators might be more likely to prey upon attached pupae at later time points. Therefore, we did not use a Cox proportional hazards model for survival analysis as the ratio of the hazard functions for individuals within the same bucket may change over time. We did not correct p-values for multiple testing, as suggested by Nakagawa [35].

3. Results

(a) Glue adhesion strength varies between species from the same location

We collected drosophilid flies in the forest near Paris in June 2020 and noted the presence of nine Drosophila species (see [30]). We established fly stocks of the most common species. Three of them, Drosophila simulans, D. hydei and D. suzukii, were assayed for pupal adhesion. We found that D. simulans detached at a median strength of 234.2 mN (figure 1a), similar to what has been found previously for its sister species D. melanogaster [31]. We measured a lower adhesion for D. suzukii pupae with a median strength of 78.7 mN and higher adhesion for D. hydei with a median strength of 482.6 mN. Adhesion strength was significantly different between the three species (Kruskal–Wallis $\chi^2 = 0.00$.) but not different between D. simulans and D. suzukii ($p=0.02$). There was no difference in slope among the three species ($p=0.04$) and the common slope was about 491, meaning that adhesion force increases by 491 mN for 1 mm$^2$ increase of the pupa–substrate contact area for each species. After dividing the adhesion force by the surface of contact, we found a difference in adhesion between D. simulans and D. hydei (figure 1c; Kruskal–Wallis $\chi^2 = 9.61$, $p=0.008$, followed by all pairwise comparison Wilcoxon test, $p=0.0008$) but not between D. simulans and D. suzukii ($p=0.1$) and D. suzukii and D. hydei ($p=0.6$).
Figure 1. Pupa adhesion varies between species originating from the same location. (a) Adhesion strength of three *Drosophila* lines collected in Vincennes. Force indicates the force required to detach a pupa naturally attached to a glass slide. Each dot corresponds to a single pupa and *n* indicates the total number of pupae tested for each species. Ends of the boxes define the first and third quartiles. The black horizontal line represents the median. The vertical line on the top of the box extends to the largest value no further than 1.5 × IQR from the upper hinge of the box. The vertical line on the bottom of the box extends to the smallest value at most 1.5 ×IQR of the hinge (IQR, inter-quartile range is the distance between the first and the third quartiles.). Data beyond the end of these lines are ‘outlying’ points. Asterisk indicates significant differences between *D. suzukii* and *D. hydei*, *D. suzukii* and *D. simulans* and *D. simulans* and *D. hydei* (*p < 0.05*). (b) Relationship between pupa adhesion force and pupa–substrate contact area. Each dot corresponds to a single pupa. *Drosophila* suzukii pupae are represented as circles, *D. simulans* as triangles and *D. hydei* as stars. (c) Adhesion strength corrected by the pupa–substrate contact area. Boxplots and asterisk as in (a). n.s. indicates not significant (*p > 0.05*).

To test whether the production of the glue was related to the size of the pupa in each species, we imaged attached pupae raised in the same conditions as the pupae used for the adhesion tests and measured the area of the pupal case. We found that the three species had different pupal sizes (Kruskal–Wallis *χ*^2^ = 73.876, *p < 2 × 10^-16*, followed by all pairwise comparison Wilcoxon test, *p < 0.001*) with *D. hydei* pupae presenting the biggest size (4.63 mm^2^) and *D. simulans* (2.78 mm^2^) and *D. suzukii* (2.12 mm^2^). The ratio of glue print area over pupal case area was 0.21 for *D. simulans* and *D. hydei*, suggesting that *D. suzukii* pupae produce less glue relative to their size than *D. simulans* and *D. hydei*.

(b) Attached pupae are taken away less frequently than detached pupae in a semi-natural environment
To test whether glue attachment may protect pupae from predation in a semi-natural environment, we chose to use *D. simulans*, as we could obtain a large number of pupae from our fly strain. We compared the disappearance of pupae naturally attached to the plastic lid of Petri dishes with pupae mechanically detached from the lid. We placed two dishes containing, respectively, 10 attached and 10 detached pupae in 28 open buckets in Bois de Vincennes (two Petri dishes in each bucket) for 54 h and monitored the number of remaining pupae twice a day (figure 2a,b). At the end of the experiment, 10 pupae (median = 10) remained in the dish with attached pupae (all still attached) compared to 6–7 pupae (median = 6.5) in the dish with detached pupae. We found that attached pupae stayed significantly more in the Petri dishes than detached ones, with differences becoming significant from day 2 morning until the end of the experiment (figure 2f); paired Wilcoxon rank tests with continuity correction: day 1 afternoon: *V* = 9, *p* = 0.2; day 2 morning: *V* = 89.5, *p* = 0.02; day 2 afternoon: *V* = 92, *p* = 0.01; day 3 morning: *V* = 91, *p* = 0.002; day 3 PM: *V* = 110, *p* = 0.005). After 3 days, attached pupae remained onsite 30% more than detached pupae. During the countings, a few species were observed in the Petri dishes: *T. nylanderi* ants, red spider mites, cockroaches and woodlice (see [30]). Red spider mites were seen to fix themselves to both attached and detached pupae (figure 2c). *Tenebrio* nylanderi was the only species that was clearly seen consuming both attached and detached pupae in the dishes (figure 2d). Cockroaches were also observed, but only on the first day (figure 2c). Woodlice were also found in the dishes but never in contact with the pupae.

(c) Attached pupae are predated less efficiently by ants
To further understand how predators may act when they encounter an attached or a detached pupa, we decided to monitor in the laboratory pupae predation by the ant *T. nylanderi*, which was the most commonly found predator of *D. simulans* pupae in our field assay. Seven ant colonies were collected in Bois de Vincennes. After 10-day starvation, each ant colony was given on each day, one glass slide with two pupae, an attached and a detached one. We used either two pupae of *D. simulans* (naturally strongly attached) or two pupae of *D. suzukii* (naturally loosely attached) and we alternated colonies each day. We examined the ant–*Drosophila* interactions every 5 min for 3 h after adding the glass slide with pupae.

We found that in all replicates, all pupae were consumed by the ants. For both fly species, we observed a difference between detached and attached pupae: detached pupae were mostly taken to the nest and eaten there while most of the attached pupae were eaten on site (figure 3a; number of
pupae taken to the nest in *D. simulans*: detached 21 of 21, attached 3 of 21, $\chi^2 = 28.10$, $p < 10^{-6}$; in *D. suzukii*: detached 19 of 21, attached 7 of 21, $\chi^2 = 12.22$, d.f. = 1, $p = 0.0005$). In nine cases (6 for *D. suzukii* and 3 for *D. simulans*), both the attached and the detached pupa were taken to the nest; in all those cases, the detached pupa was always taken to the nest first (about 37 min earlier for *D. suzukii* and 50 min earlier for *D. simulans* in median; figure 3a).

Ants spent 15 and 45 min (median) outside the nest in contact with *D. simulans* and *D. suzukii* detached pupae, respectively, while they spent, respectively, six and two times longer in contact with attached pupae (figure 3b; respectively, 100 and 110 min in *D. simulans* and *D. suzukii*, paired Wilcoxon rank tests with continuity correction *D. simulans*: $V = 4.5$, $p = 0.0001$; *D. suzukii*: $V = 18$, $p = 0.0007$). The maximum number of ants observed in direct contact with the fly pupa over the duration of the experiment was significantly higher for attached pupae than for detached pupae (figure 3a,c; median of 9 for *D. simulans* and 8 for *D. suzukii* for attached pupae compared to the median of 2 for detached pupae in both species, paired Wilcoxon rank tests with continuity correction *D. simulans*: $V = 2.5$, $p = 0.0002$; *D. suzukii*: $V = 10.5$, $p = 0.0007$). The difference was still significant after correcting for the amount of time, by comparing the maximum number of ants in contact with the pupa until the first pupa is brought to the nest or fully consumed (paired Wilcoxon rank tests with continuity correction, *D. simulans*: 3 versus 2, $V = 24$, $p = 0.02$; *D. suzukii* 6 versus 2, $V = 7$, $p = 0.004$).

The time for the first ant to touch a pupa was slightly significantly different between attached and detached pupae only in *D. suzukii* (paired Wilcoxon rank tests with continuity correction, *D. suzukii*: 15 versus 20 min, $V = 31$, $p = 0.03$; *D. simulans*: 15 versus 20 min, $V = 62.5$, $p = 0.3$) but not different between *D. simulans* and *D. suzukii* (Wilcoxon rank tests with continuity correction, attached pupae: $W = 195.5$, $p = 0.5$; detached pupae: $W = 200$, $p = 0.6$). No...
differences were found between *D. simulans* and *D. suzukii* regarding the time until the detached pupae is brought to the nest from the start of the experiment (45 versus 65 min, \( W = 160.5, p = 0.296 \)), the time that ants spent in contact with attached pupa (\( W = 194, p = 0.5 \)) and detached pupa (\( W = 161, p = 0.1 \)), the maximum number of ants over the duration of the experiment on detached pupae (\( W = 196.5, p = 0.5 \)) or on attached pupae (\( W = 190.5, p = 0.5 \)), the maximum number of ants until the first pupa disappeared on detached pupae (\( W = 231.5, p = 0.8 \)) and the maximum number of ants on attached pupae (\( W = 265, p = 0.3 \)).

4. Discussion

(a) In the same environment, pupa adhesion strength varies among species

Using our previously published pull-force measurement assay [31], we provide here, to our knowledge, the first evidence that pupa adhesion varies between *Drosophila* species. Our result is in agreement with the rapid evolution of glue genes [16]. Our analysis unravels at least two mechanisms leading to changes in the quantity of glue produced and resulting in changes in...
adhesion among species: (i) a change in body size (probably linked with a change in salivary gland size), as observed between D. hydei and D. simulans, and (ii) a change in the amount of glue production independently of body size, as observed between D. suzukii and D. simulans. We note that our experiment might not reflect natural conditions as we have not tested adhesion on natural substrates and in natural habitats.

The variation we observed in adhesion force among individuals within a given species is much higher than measurement error (our universal test machine has an accuracy of ±0.5%) and could be owing to individual variation in size, shape, weight, glue production or position of the pupa relative to the substrate.

At the end of the larval stage, larvae stop feeding and start to search for a site to pupate. Pupation site choice during the larval stage is important for pupal survival. Pupation site preference has been thoroughly investigated in the laboratory [36–38] and more rarely in nature [39]. This choice depends on abiotic factors such as temperature [37], darkness [40] or the nature of the substrates [41]. In particular, D. simulans prefers to pupate on rough and humid surfaces while D. hydei prefers dry and smooth surfaces. Drosophila simulans was often reported as pupating in fruits in the laboratory [36,38] and from field sampling [39], but other natural sites have not been investigated. In D. suzukii, recent studies in the field have found that pupae are present in the soil rather than in fruits [20,22]. Additionally, pupation site preference depends on biotic factors and particularly on the presence of conspecifics and alien species. In D. simulans and D. hydei and in other Drosophila species, pupae are aggregated with conspecifics [39,42]. In D. simulans and Drosophila buzzatii, larvae change their site choice in the presence of heterospecific larval cues [19]. The differences we observed in adhesion strength among species may reflect differences in their ecology. For example, an adhesive substance might not be required for animals pupating on sticky substrates (rotten fruits and mushrooms, sap, etc.). Furthermore, species exhibiting distinct adhesion strengths in our laboratory conditions may nevertheless stick with similar forces in their respective natural habitats.

(b) Fixation of the pupa prevents predation

Comparing the disappearance of attached and detached D. simulans pupae in the field, we found that being attached allows pupae to stay on site more efficiently. Because pupae were contained within the lid of Petri dishes, we infer that they were not blown away by light wind, but we cannot be sure that missing pupae were predated. In our design, predators have the choice between attached and detached pupa within a bucket. This design reduces the possible effect of buckets when comparing attached and detached pupa disappearance.

The ant T. nylanderi was the main predator that we observed in the field consuming pupae. This observation is in agreement with previous studies which found that ants prey on fruit fly pupae [20,22,29]. During our laboratory assay, T. nylanderi ants predated attached and detached pupae with distinct behaviours: they brought most of the detached pupae to the nest while they ate the attached ones directly on site. The latter strategy requires ants to spend more time outside the nest and to recruit more foragers. The presence of parasites, predators and competitors in the wild would certainly make this strategy costly. Additionally, T. nylanderi is a solitary foraging species that recruits nest-mates one by one [43]. Under natural conditions, it would take a relatively long time to gather many foragers around the pupae. In our field assay, no more than two ants were observed together in a lid (see [30]). We found that ants act similarly on D. suzukii and D. simulans pupae, suggesting that there is no difference in strategy to predate loosely attached pupae such as D. suzukii or more strongly attached ones such as D. simulans, and that both species are equally attractive as prey.

To our knowledge, this study is the first to show that fly pupa adhesion can protect from predation. Our experiments are simple and can be easily applicable to other species, and not only for pupae but also for eggs, to check if this phenomenon is general in flies and insects.

(c) Alternative functions of Drosophila glue and alternative strategies to protect pupae from predation

Here, we demonstrated that pupal adhesion protects from predation by preventing predators like ants from taking the pupa away. Pupa adhesion may have alternative roles, such as maintaining the individual in a favourable environment [3,9,13] so that the pupa would be hidden from predators, protected from microorganism contaminants or/and have optimal conditions for pupal development. If not attached, the pupa could be moved away by abiotic factors such as wind or rain, or biotic factors such as competitors. Pupa attachment could also help the adult to emerge from the pupal case, as suggested in butterflies [11] or facilitate pupal aggregation and thus dilution of predation risk as in freshwater caddisflies [12]. Pupal congregation has been observed in Drosophila species [39,42], but its contribution to protection from predators has not been tested.

Pupal adhesion is only one of several strategies for pupae to escape predators. A common strategy is cryptic coloration to hide from visual predators [44,45]. The brownish colour of Drosophila pupae could contribute to hiding the animal when pupating in the leaf litter or in the soil. In some cases, pupae mimic non-living things such as leaves or sticks like the common maplelet butterfly chrysalis [45]. To avoid non-visual predators, pupa has also evolved chemical defences either to chemically hide from predators [46] or to make the pupa toxic [47]. Pupae have also evolved different types of physical defence (spin, hard pupal case, urticating hairs, etc.). In Drosophila, the pupa is covered by a relatively thick (about 20 µm, [48]) and hard cuticle which has been hypothesized to protect the animal from predator attacks.

For the first time, to our knowledge, we report that pupal adhesion varies among Drosophila species and that pupa attachment can protect from predation. Our results unravel a previously unknown important trait for Drosophila survival in the wild, the ability of pupae to firmly adhere to a substrate. Further studies of Drosophila glue combining genetic and phenotypic approaches should provide insight on the molecular basis for diverse biadhesive properties, adapted to various habitats and climates.

Data accessibility. Supplementary material, tables, raw data and scripts are available from the Dryad Digital Repository: https://doi.org/10.5061/dryad.x3fbg7hg [30].

Authors’ contributions. F.B., M.M. and V.C.-O. designed the experiments, V.C.-O. supervised the project, S.R.P. collected, identified and raised wild-caught flies, F.B. performed all the other experiments, F.B., M.M.
and V.C.-O. analysed the results, F.B. wrote the original draft and all authors edited and contributed to the final version of this paper.

**Competing interests.** We declare we have no competing interests.

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