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ORIGINAL ARTICLE

Size-dependent flight capacity and propensity in a range-expanding invasive insect



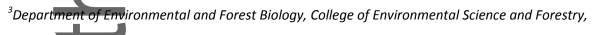
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Abstract

For capital-breeding insects, all resources available for adult metabolic needs are accumulated during larval feeding. Therefore, body size at adult eclosion represents the total energetic capacity of the individual. For female capital breeders, body size is strongly correlated with lifetime fecundity, while in males, body size correlates with fitness are less understood. In capitalbreeding species with wingless, flightless, or dispersal-limited females, flight potential for male Lepidoptera has important implications for mate-finding and may be correlated with body size. At low population densities, failure to mate has been identified as an important Allee effect and can drive the success or failure of invasive species at range edges and in species of conservation concern.

The European gypsy moth (EGM), *Lymantria dispar* (L.), a capital breeder, was introduced to North America in 1869 and now ranges across much of eastern North America. In EGM, females are flightless and mate-finding is entirely performed by males. We quantified male EGM flight capacity and propensity to fly relative to morphological and physiological characteristics using fixed-arm flight mills. A range of male body sizes were produced by varying the protein content of standard artificial diets while holding other dietary components constant. Wing length, a proxy for body size, as well as relative thorax mass and forewing aspect were all important predictors of total flight distance and maximum speed. These results have important implications for mate-finding and invasion dynamics in gypsy moth and may apply broadly to other capital-breeding insects.

Keywords flight mill; Lymantria dispar; life-history; invasion dynamics; selective pressure

Introduction

Variation in resources and their allocation affects the quantity and fitness of offspring that individuals produce in a given environment (Boggs & Freeman, 2005; Boggs, 2009). For holometabolous insects including Lepidoptera, energy and nutrients acquired in the larval stage are later redistributed to fuel the formation of adult body structures during the pupal stage and meet adult metabolic and reproductive needs (Boggs, 2009). While income breeding species feed as adults and use accumulated resources to supplement reproduction, capital breeders depend entirely on resources accrued as larvae for all adult functions (Tammaru & Haukioja, 1996; Davis *et al.*, 2016). Thus, in capital breeders, body size is indicative of the total energy and material reserve available for adult dispersal, mate finding, and reproduction.

Estimating fitness for female capital-breeding insects is done relatively easily by counting the number of eggs a female produces (Hough & Pimentel, 1978; Tammaru *et al.*, 1996a; Tisdale & Sappington 2001, Parry *et al.*, 2001). On the other hand, determining correlates of male fitness by quantifying the number of eggs fertilized following a controlled mating event or by measuring male flight capacity is more difficult and is done less frequently (e.g., Boggs & Gilbert, 1979; Svärd & Wiklund, 1939; Tammaru *et al.*, 1996b; Tobin *et al.*, 2014). Male flight may be particularly important for capital-breeding species where males do most or all of the mate-finding (Roff, 1991; Berec *et al.*, 2018; Javoiš *et al.*, 2019). Mate location is critical for overcoming mate-finding Allee effects in small isolated **populations** or those at low density near range edges (Contarini *et al.*, 2009; Yamanaka & Liebhold, 2009; Rhainds, 2010). Therefore, factors that influence male flight capacity and mate-finding ability in capital-breeding species (e.g. resource acquisition and body size) may impact persistence of small populations and establishment in range-expanding populations.

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The European strain of gypsy moth was introduced to North America in 1869 and now ranges from northern Minnesota to the Outer Banks of North Carolina (Grayson & Johnson, 2018). Adults are sexually dimorphic and, in North America, have flight-capable males and winged but flightless females (Doane & McManus, 1981). Thus, energy expended by adult females is allocated primarily to the production and emission of pheromone and the manufacturing and provisioning of eggs, whereas males allocate their resources to flight. While the strong linear relationship between female pupat mass and fecundity is well established (e.g., Hough & Pimentel, 1978, Faske *et al.*, 2019), no studies to our knowledge have assessed the relationship between body size and flight capacity (e.g. Davis *et al.*, 2012; Evenden *et al.*, 2014; Jones *et al.*, 2016), and flight-related morphology such as wing load and forewing aspect ratio affect flight capability in other Lepidoptera (Le Roy *et al.* 2019), male gypsy moth flight capacity may also be affected by body size and flight-related morphology.

Fixed-arm flight mills provide a valuable avenue for investigating effects of physiological, morphological, and environmental variation on flight capacity and dispersal under controlled conditions (Minter *et al.*, 2018; Naranjo, 2019). Although tethered flight mills have been used to determine the effect of morphological and physiological characteristics on females of the Asian gypsy moth, a strain with strong female flight ability (Shi *et al.*, 2015, Yang *et al.*, 2017), to our knowledge none have done so for any strain of male gypsy moth. Our study assessed the relationship between resource acquisition and allocation and male flight capacity for European gypsy moth, an economically and ecologically important invasive capital breeder. Specifically, we used flight mills to assess the direction and influence of several morphological and physiological characteristics on male gypsy moth flight performance. We predicted that (i) due to greater stored energetic resource availability with increasing male body size, increasing body size would correspond

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to greater flight distance, speed, and flight propensity, and (ii) variation in wing and thoracic morphology would affect male flight distance and speed.

Materials and methods

Insect rearing

In natural environments, the mass of male gypsy moth pupae and adults varies greatly as a function of dier quality, population density, and temperature experienced as a larva (Carter *et al.*, 1991; Lindroth *et al.*, 1997; Lazarevic *et al.*, 2004, Jahant-Miller *et al.*, 2020). To investigate the effects of male size on flight capacity required methodology to produce a range of sizes while minimizing the effects of other confounding factors on flight metrics. To produce a range of male sizes for exterimentation, we used an agar-based artificial gypsy moth diet (Leonard & Doane, 1966) and manipulated protein content while holding all other nutrients constant. Males were reared on four diet formulations (high-protein, medium-high protein, medium-low protein, low protein; Table S1). To compensate for reduced protein content, the equivalent amount by weight of food grade cellulose powder was added to keep the weight ratio of other ingredients equal across all diet mixes. Fresh diet was poured in 0.47 l plastic-lined paper soup containers and provided to larvae *ad libitum*. Larvae were reared in growth chamber at 25°C and a photoperiod of 15:9 (L:D). Upon pupation, individuals were weighed to the nearest 0.0001 g and returned to the growth chamber in individual 59.15 ml plastic containers until adult emergence.

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Flight mill design

Flight mills were assembled using the design described by Jones *et al.* (2010; Appendix 1). During each assay, a moth was attached distally to a rotating arm (18.34 cm diameter) and a magnetic sensor detected a small magnet below the flight mill arm at each full rotation as the moth powered the mill (Fig. 1A). The magnetic sensor was connected to a microcontroller (Fig. S1) that read up to eight sensors and converted rotation and time information to a USB signal for logging by a terminal program (TeraTerm) on a personal computer. Opposite-facing magnets reduced rotational friction by levitating the flight mill arm above the axle.

Flight assay

Pupae were checked at least twice daily for adult emergence. Moths that emerged and were not flown right away were held individually in small plastic containres at 4°C until ready to be flown. All flight assays were initiated within 24 hours of emergence. Prior to harness placement, moths were cold anesthetized at approximately –18°C for up to two minutes, then weighed to the nearest 0.0001 g. Inverted y-shaped wire harnesses (Fig. 1B) were glued dorsally to the descaled first abdominal segment using cyanoacrylate gel super glue. Moths were reweighed with the harness and attached to the distal end of the flight mill arm using thin plastic tubing. A piece of soft polymer clay equal to the weight of the moth and harness was attached to the opposite end as a counterbalance.

Flight assays began at the start of the photophase (light period) and ran for 24 hours with photoperiod of 15:9 (L : D). Photophase and scotophase temperatures were 25°C and 15°C, respectively, which are similar to ambient temperatures in gypsy moths encounter in their invasive range. During flight assays, total distance and duration of each individual flight was recorded, along with morphological (pupal mass, moth mass, wing length, width, and area, thorax width, height and

shape, relative thorax mass, relative abdominal mass, and forewing aspect) and physiological (mass lost during flight, relative mass lost during flight) moth characteristics. Moths that had stopped flying were encouraged to fly every 15 minutes using gentle tactile stimulation (i.e., paintbrush touches to the legs, posterior edge of the wings, or antennae) for the first eight hours of each flight assay. Moths that became detached from the flight mill during the flight assay or that did not fly at all due to injury during pre-flight procedures were excluded from analyses. Following the 24-hour flight assay, the moths were removed from the flight mills, frozen, and reweighed to quantify mass lost during flight. The right forewing of each male was removed and scanned, and the digital image of each wing was analyzed using ImageJ software to determine wing length, wing width, and wing area. Thorax mass and abdominal mass were measured by carefully separating each tagma using a scalpel and weighing them individually on a digital balance to the nearest 0.0001 g. Wing load was calculated as initial moth mass divided by two times the wing area (g/cm²). Forewing aspect ratio was calculated as four times the squared length of the forewing divided by forewing area (mm/cm²). Relative thorax mass was calculated as the mass of the thorax divided by the pre-flight moth mass, and relative abdominal mass was calculated as the mass of the abdomen divided by the pre-flight moth mass. Thorax width and height were measured to the nearest 0.01 mm using a digital caliper. Thorax shape was calculated as the width of the thorax divided by the height. Mass lost during flight was measured as the initial mass of the moth minus the mass of the moth measured immediately after the flight assay, and ratio of mass lost was calculated as mass lost during flight divided by initial moth mass

Flight propensity was measured as the number of flights consisting of more than 100 full rotations initiated upon tactile stimulation divided by the total number of touches for each individual moth. One-hundred rotations was equal to approximately 58 m of linear flight.

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Variable selection

Total distance flown and maximum speed were used as dependent variables. Based on biological significance, wing length, which correlates strongly with pupal mass (Tobin *et al.*, 2016; Jahant-Miller *et al.*, 2020), was chosen to represent body size, and proportionate mass lost during flight (RatioMD), forewing aspect ratio (FWa), and proportionate thorax mass (RTM) were also included in the model and each represented a different physiological or morphological component. Variables selected for the full models were not strongly correlated (P < 0.4). Flight mill identity was included in both models as a random effect to capture any variation in measured flight capacity metrics caused by any physical differences in flight mill design. Since insects do not need to produce lift during tethered flight, wing load was not included in flight capacity analyses.

Statistical analysis

All data analyses were performed in R (version 3.6.1, R Core Development Team, 2019). Significance level for all analyses was $\alpha = 0.05$. Effects of morphological and physiological characteristics on metrics of male flight capacity were analyzed using mixed effects linear regression analysis. Fixed effects and pairwise interaction effects were tested using stepwise model selection (package *ImerTest*; Kuznetsova *et al.*, 2017). For each dependent variable, full and reduced models were compared using AIC and ANOVA hypothesis testing. Model normality and homoscedasticity were checked using residual plots. Total distance flown was square root transformed to improve normality. Effect of wing length on flight propensity was compared using one-way ANOVA.

A total of 96 male gypsy moths were flown. Mean pupal mass of flown moths was 0.41 g \pm 0.013 (mean \pm s.e.) and ranged from 0.19–0.73 g, values that encompass much of the variation seen in males from natural populations (Jahant-Miller *et al.* 2020). Mean wing length was 2.05 cm \pm 0.021 and ranged from 1.64–2.53 cm (Table 1). High protein diet produced the greatest mean pupal mass (0.4362 \pm 0.014 g), followed by medium-high (0.4002 \pm 0.048 g), medium-low (0.3409 \pm 0.027 g), and low (0.1823 \pm 0.014 g). High, medium-high, and medium-low protein diets supported relatively high survival rates, while low protein diet did not. As a result, no moths reared on low-protein diet were used in flight assays. In total, high-protein, medium-high, and medium-low diets produced 70, 11, and 15 moths flown in flight assays, respectively.

Flight capacity

Wing length was the strongest predictor for both flight capacity metrics (total distance and maximum speed) in the mixed regression models (Table 2, 3). Total distance flown ($F_{1, 99} = 98.47$, P < 0.0001; Fig. 2A) and maximum speed ($F_{1, 93} = 80.17$, P < 0.0001; Fig. 3A) were significantly impacted by wing length. targe moths in the upper quartile for wing length (n = 27, $X_{WL} = 2.33$ cm [2.22–2.53 cm]) flew on average 31.6 km [11.6–68.2 km], or 282% more than small moths in the lower quartile (n = 24, $X_{WL} = 1.80$ cm [1.64–1.89 cm]) which flew on average 11.2 km [0.12–44.9 km]. Forewing aspect ratio had a significant negative effect on total distance ($F_{1, 91} = 30.32$, P < 0.0001; Fig. 2D) and maximum speed ($F_{1, 93} = 17.70$, P < 0.0001; Fig. 3B). Reduced forewing aspect was associated with greater flight speed and greater total distance flown. Relative thorax mass had a significant positive effect on total distance ($F_{1, 91} = 28.17$, P < 0.001; Fig. 2B).

Flight propensity

All but five moths recorded at least one response flight consisting of more than 100 full rotations; wing lengths of those that did not were 17.0, 17.22, 17.62, 19.13, and 20.34 mm. Wing

length had a significant effect on proportion of sustained flights initiated ($F_{4,91}$ = 9.43, P < 0.001; Fig. 3C); males with greater wing length were more likely to initiate sustained flight than males with smaller wings.

Discussion

Insect body size is often correlated with greater flight distance and speed (e.g., Dingle *et al.*, 1980; Shirai 1991; Bruzzone *et al.*, 2009; Evenden *et al.*, 2014; Fahrner *et al.*, 2014; Jones *et al.*, 2016). This study is the first to assess the impact of body size and flight morphometrics on measures of flight canacity for male gypsy moth. Wing length, a proxy for body size, was the most important predictor for both metrics of flight capacity; large moths were more likely than small moths to initiate sustained flight, and once initiated, larger moths tended toward longer, faster sustained flights than smaller moths. The greatest distance flown in the present study was more than 68 km and was accomplished by one of the largest moths (24.6 mm wing length). However, many midsize moths (19 24 mm wing length) in our study also flew total distances greater than 30 km. Moths with wing lengths between 19 mm and 21 mm are common in both endemic and high-density populations (Tobin *et al.*, 2016; Jahant-Miller *et al.*, 2020). Due to altered physical properties of tethered flight relative to free flight, such as a forced flight path, lack of natural flight stimuli, and the fact that tethered insects do not need to produce lift, extrapolation of tethered flight distances to flight potential in natural settings should be done with caution (Minter *et al.*, 2018).

Other significant predictors for metrics of flight capacity included forewing aspect ratio for speed and distance, and relative thorax mass for distance, and relative mass lost during flight for distance. Aspect ratio describes the shape of a wing where higher values indicate a longer, narrower wing and lower values indicate a shorter, broader wing. For aircraft and some flying animals

including insects, aspect ratio is negatively correlated with flight speed and positively correlated with flight efficiency during gliding flight since longer, narrower wings produce less drag (Savile, 1957; Dudley & Srygley, 1994; Davis *et al.*, 2012). Energy storage for insect flight and flight musculature are thoracic components, and flight metrics such as distance and speed are often correlated with increased relative thorax mass (Dudley & Srygley, 1994; Marden, 2000; David *et al.*, 2015). Relative mass lost during flight had a negative relationship with total distance flown, indicating that larger moths may use a smaller proportion of their stored resources than smaller moths and may have the capacity to cortinue flying longer than our flight assay allowed.

Most female gypsy moth mate only once and appear non-selective, thus, the first male to reach the female has a reproductive advantage (Doane & McManus, 1981). This suggests that being the fastest filer with the greatest endurance would be ideal; however, large body size can generally only be attained at the cost of longer larval development. All else being equal (i.e. foliar quality, temperature, etc.), genetically larger individuals take more time to develop than smaller conspecifics (Hough & Pimentel, 1978; Teder *et al.*, 2014, but see Thompson *et al.*, (2020) for latitudinal differences among populations). Adaptation towards greater flight capacity may ultimately put larger males at a mating disadvantage unless environmental conditions require mate-finding flights that cover greater distances, such as in low density populations. Conversely, due to protandrous emergence in gypsy moth, larger males could be at an advantage if delayed emergence relative to smaller males improves synchrony with female emergence (Robinet *et al.*, 2007), especially in low density populations. While body size is the greatest factor impacting flight capacity, changes to wing or thorax morphology can also alter gypsy moth flight ability and could do so without affecting development time.

Density-dependent shifts in mating or mate-finding morphology occur in other arthropods. In the amphipod *Gammarus roeselii*, individuals exhibit differential allocation to mating morphology in response to population density (Lipkowski *et al.*, 2019). In high density *G. roeselii* populations where mating competition is relatively intense, individuals express larger antennae to increase mate selectivity and larger grasping legs to reduce the likelihood of having a mate stolen. In a wingdimorphic planthopper *Prokelisia dolus* Wilson, flight-capable males in low-density populations with fully developed wings had greater mate-finding ability compared to wingless morphs, but at the cost of reduced fecundity (Langellotto & Denno, 2001).

Morphological and physiological adaptation that facilitates dispersal has been documented for several plant and animal phyla at invasion fronts, as well as areas in which species frequently undergo extinction and colonization events (Phillips *et al.*, 2006; Darling *et al.*, 2007; Forsman *et al.*, 2010; Hill *et al.*, 2011; Schäfer *et al.*, 2018). In cane toads, *Rhinella marina* (Linneaus), for example, dispersal ability increases with leg length, and more recently established populations (e.g., those at the invasion front) in Australia have longer legs than populations behind the invasion front (Phillips *et al.*, 2006). Similar intraspecific variation in dispersal related morphology and physiology has been reported for the butterfly *Melitaea cinxia* (Linneaus) and for invasive starlings, *Sturnus vulgaris* Linneaus, in Australia (Haag *et al.*, 2005).

In low-density populations, emigration can exacerbate inverse density-dependent Allee effects such as mate-finding success and cooperative feeding behaviors, and ultimately lead to range retraction (Taylor & Hastings, 2005; Tobin *et al.*, 2009). However, in gypsy moth, male dispersal from established populations in the range interior to the range margin could improve mating success by alleviating Allee effects caused by mate-finding failure typical of very low density populations (Sharov *et al.*, 1995; Liebhold & Bascompte, 2003; Contarini *et al.*, 2009; Tobin *et al.*, 2009).

Increased dispersal ability of male gypsy moths from just behind the invasion front to nascent satellite populations ahead of it has the potential to increase establishment success and the rate of range expansion.

Size-dependent effects of flight ability on mate-finding may be relevant to other capitalbreeding Lepidoptera, especially populations experiencing Allee effects, and in which females are non-disperang prior to mating. A body-size effect on mate finding success has been shown in some other capital-breeding Lepidoptera. For example, male autumnal moth *Epirrita autumnata* Borkhausen mating in low density field populations were larger than mean population size, indicating that there may be a size-dependent advantage, although it is unclear whether it is from mate-finding or mating competition (Tammaru *et al.*, 1996b). Evenden *et al.* (2015a) identified a mate-finding alter effect in a capital-breeding spring defoliator, *Malacosoma disstria* Hubner; caged females were mated more frequently in high-density populations than low. Although large males originating from endemic populations, there was no apparent mate-finding advantage for larger body size (Evenden *et al.*, 2015a, b). Mated males of the saturniid *Rothschildia lebeau* Guerin-Meneville tended to be larger on average than non-mated males in a field study that used caged females to attract endemic males (Agosta, 2010).

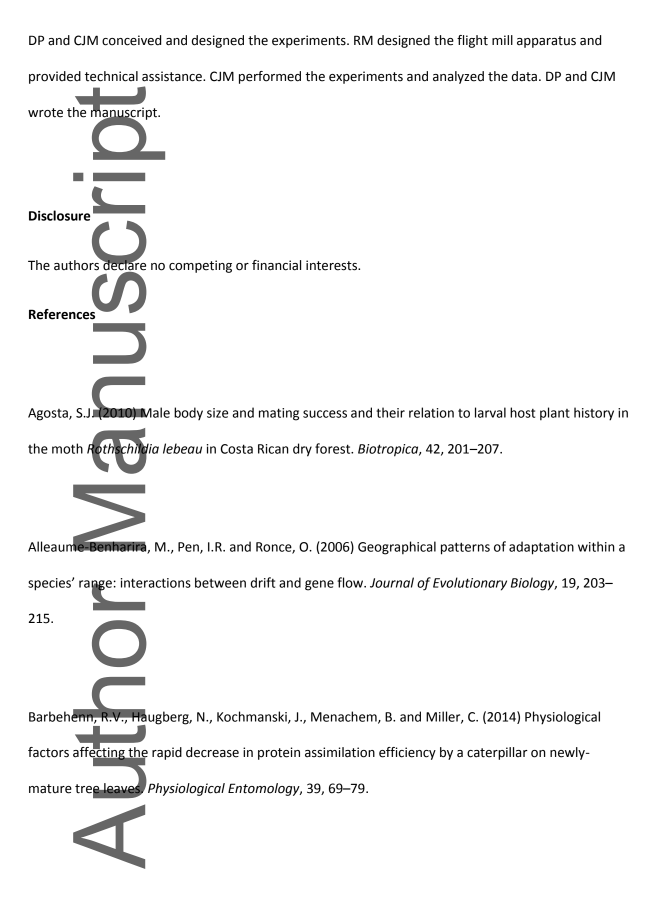
Adaptive shifts in wing morphology facilitating dispersal ability at the invasion front are unknown in gypsy moth. Counterselection favoring small male body size may be driven by short growing seasons and cool climates encountered by range-expanding populations in northern regions. In addition, a significant component of gypsy moth range expansion is driven by humanvectored movement of life stages (Sharov & Liebhold, 1998; Bigsby *et al.* 2011; Tobin and Blackburn, 2014). Movement of propagules from established areas to range front populations may swamp any

adaptive change. Despite known barriers to adaptation, recent studies indicate that rapid adaptation is occurring in gypsy moth populations in response to environmental conditions at range margins (e.g., Thompson *et al.*, 2017, 2020, Faske *et al.*, 2019). Future studies should take advantage of common garden studies, which disentangle genetic and environmental effects on body size, to detect potential shifts in gypsy moth body size or flight morphology in this and other rangeexpanding invasive insects.

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Barton, K. (2020) Package 'MuMIn'. Model selection and model averaging base on information criteria. R package version 1.43.17. R Foundation for Statistical Computing, Vienna, Austria.

Berec, L., Kramer, A.M., Bernhauerová, V. and Drake, J. M. (2018) Density-dependent selection on mate search and evolution of Allee effects. *Journal of Animal Ecology*, 87, 24–35.

Bigsby, K.M., Fobin, P.C. and Sills, E.O. (2011) Anthropogenic drivers of gypsy moth spread. *Biological Invasions*, 13, 2077–2090.



Boggs, C.L. (2009) Understanding insect life histories and senescence through a resource allocation lens. *Functional Ecology*, 23, 27–37.

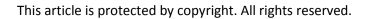


Boggs, C. L. and Freeman, K. D. (2005) Larval food limitation in butterflies: effects on adult resource allocation and fitness. *Oecologia*, 144, 353–361.



Bontrager, M. and Angert, A.L. (2019) Gene flow improves fitness at a range edge under climate

change. Evolution Letters, 3, 55–68.



Bruzzone, O.A., Villacide, J.M., Bernstein, C. and Corley, J.C. (2009) Flight variability in the woodwasp *Sirex noctilio* (Hymenoptera: Siricidae): An analysis of flight data using wavelets. *Journal of Experimental Biology*, 212, 731–737.



Burnham, K.P. and Anderson, D.R. (2004) Multimodel inference: Understanding AIC and BIC in model selection. *Sociological Methods & Research*, 33, 261–304.



Carter, M.R., Ravlin, F.W. and McManus, M.L. (1991) Changes in gypsy moth (Lepidoptera:

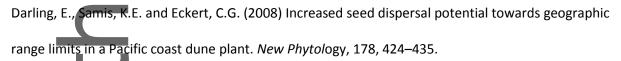
Lymantriidae) fecundity and male wing length resulting from defoliation. Environmental Entomology,

20, 1042–1047.



Contarini, M., Onufrieva, K.S., Thorpe, K.W., Raffa, K.F. and Tobin, P.C. (2009) Mate-finding failure as an important cause of Allee effects along the leading edge of an invading insect population.

Entomologia Experimentalis et Applicata, 133, 307–314.





David, G., Giffard, B., Van Halder, I., Piou, D. and 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.



Davis, A.K., Chi, J., Bradley, C. and Altizer, S. (2012) The redder the better: wing color predicts flight performance in monarch butterflies. *PLoS ONE*, 7, e41323.

Davis, R. B., Javoiš, I., Kaasik, A., Õunap, E. and Tammaru, T. (2016) An ordination of life histories using morphological proxies: Capital vs. income breeding in insects. *Ecology*, 97, 2112–2124.

Dingle, H., Blakley, N.R. and Miller, E.R. (1980) Variation in body size and flight performance in milkweed bugs (*Oncopeltus*). *Evolution*, 34, 371–385.

Dudley, R. and Srygley, R. (1994) Flight physiology of neotropical butterflies: Allometry of airspeeds during natural free flight. *Journal of Experimental Biology*, 191, 125–139.



Elliott, C.G. and Evenden, M.L. (2009) Factors influencing flight potential of *Choristoneura conflictana*. *Physiological Entomology*, 34, 71–78.



Evenden, M.L., Whitehouse, C.M. and Sykes, J. (2014) Factors influencing flight capacity of the mountain pine beetle (Coleoptera: Curculionidae: Scolytinae). *Environmental Entomology*, 43, 187–



196.

Evenden, M. L., Whitehouse, C. M. and Jones, B. C. (2015a) Resource allocation to flight in an outbreaking forest defoliator *Malacosoma disstria*. *Environmental Entomology*, 44, 835–845.

Evenden, M.L., Mori, B.A., Sjostrom, D. and Roland, J. (2015b) Forest tent caterpillar, *Malacosoma disstria* (Lepidoptera: Lasiocampidae), mate-finding behavior is greatest at intermediate population densities: implications for interpretation of moth capture in pheromone-baited traps. *Frontiers in Ecology and Evolution*, 3, 78.



Fahrner, S.J., Lelito, J.P., Blaedow, K., Heimpel, G.E. and Aukema, B.H. (2014) Factors affecting the flight capacity of *Tetrastichus planipennisi* (Hymenoptera: Eulophidae), a classical biological control agent of *Agrilus planipennis* (Coleoptera: Buprestidae). *Environmental Entomology*, 43, 1603–1612.

Faske, T.M., Thompson, L.M., Banahene, N., Levorse, A., Herrera, M.Q., Sherman, K., *et al.* (2019) Can gypsy moth stand the heat? A reciprocal transplant experiment with an invasive forest pest across its southern range margin. *Biological Invasions*, 21, 1365–1378.

Forbush, E. and Fernald, C. (1896) The Gypsy Moth. State Board of Agriculture.

Forsman, A., Merilä, J. and Ebenhard, T. (2010) Phenotypic evolution of dispersal-enhancing traits in insular voles. *Proceedings of the Royal Society B: Biological Sciences*, 278, 225–232.

Gibbs, M., Lace, L.A., Jones, M.J. and Moore, A.J. (2004) Intraspecific competition in the speckled wood butterfly *Pararge aegeria*: Effect of rearing density and gender on larval life history. *Journal of Insect Science*, 4, 16.



Grayson, K.L. and Johnson, D.M. (2018) Novel insights on population and range edge dynamics using an unparalleled spatiotemporal record of species invasion. *Journal of Animal Ecology*, 87, 581–593.



Haag, C.R., Saastamoinen, M., Marden, J.H. and Hanski, I. (2005) A candidate locus for variation in dispersal rate in a butterfly metapopulation *Proceedings of the Royal Society B: Biological Sciences*, 272, 2449–2456.

Hough, J.A. and Pimentel, D. (1978) Influence of host foliage on development, survival, and fecundity of the gypsy moth. *Environmental Entomology*, 7, 97–102.



Hunter, M.D. (2002) Landscape structure, habitat fragmentation, and the ecology of insects.

Agricultural and Forest Entomology, 4, 159–166.



Jahant-Miller, C., Tobin, P.C. and Parry, D., 2020. Spatial and temporal changes in male gypsy moth wing morphology reflect host tree phenology and habitat quality. *Agricultural and Forest*

Entomology, 22, 390–400.

Javoiš, J., Davis, R.B. and Tammaru, T. (2019) A comparative morphometric study of sensory capacity in geometrid moths. *Journal of Evolutionary Biology*, 32, 380–389.

Jones, H.B., Lim, K.S., Bell, J.R., Hill, J.K. and Chapman, J.W. (2016) Quantifying interspecific variation in dispersal ability of noctuid moths using an advanced tethered flight technique. *Ecology and Evolution*, 6, 181–190.

Kingsolver, I.G., Woods, H.A., Buckley, L.B., Potter, K.A., MacLean, H.J. and Higgins, J.K. (2011) Complex life cycles and the responses of insects to climate change. *Integrative and Comparative Biology*, 51, 719–732. Kirkpatrick, M. and Barton, N.H. (1997) Evolution of a species' range. *American Naturalist*, 150, 1–23.

Kuussaari, M., Saccheri, I., Camara, M. and Hanski, I. (1998) Allee effect and population dynamics in the Glanville fritillary butterfly. *Oikos*, 82, 384–392.

Langellotto, S.A. and Denno, R.F. (2001) Benefits of dispersal in patchy environments: Mate location

Kuznetsova, A., Brockhoff, P.B. and Christensen, R.H.B. (2017) ImerTest package: tests in linear

mixed effects models. Journal of Statistical Software, 82, DOI: 10.18637/JSS.V082.I13.

Langellotto, G.A. and Denno, R.F. (2001) Benefits of dispersal in patchy environments: Mate location by males of a wing-dimorphic insect. *Ecology*, 82, 1870–1878.



Larsen, W.A. and McCleary, S.J. (1972) The use of partial residual plots in regression analysis. *Technometrics*, 14, 781–790.



Lawrence, R.K., Mattson, W.J. and Haack, R.A. (1997) White spruce and the spruce budworm: Defining the phenological window of susceptibility. *Canadian Entomology*, 129, 291–318.

Lazarevic, J., Peric-Mataruga, V., Vlahovic, M., Mrdakovic, M. and Cvetanovic, D. (2004) Effects of rearing density on larval growth and activity of digestive enzymes in *Lymantria dispar* L. (Lepidoptera: Lymantriidae). *Folia*. *Biologica (Krakow)*, 52, 105–112.

Le Roy, C., Debat, V. and Llaurens, V. (2019) Adaptive evolution of butterfly wing shape: From morphology to behaviour. *Biological Reviews*, 94, 1261–1281.

Leonard, D.E. and Doane, C.C. (1966) An artificial diet for the gypsy moth, Porthetria dispar

(Lepidoptera: Lymantriidae). Annals of the Entomological Society of America, 59, 462–464.

Liebhold, A.M. (1995) Suitability of North American tree species to the gypsy moth: a summary of field and laboratory tests (Vol. 211). US Department of Agriculture, Forest Service, Northeastern

Liebhold, A. and Bascompte, J. (2003) The Allee effect, stochastic dynamics and the eradication of alien species. *Ecology Letters*, 6, 133–140.

Lindroth, R.L., Klein, K.A., Hemming, J.D. and Feuker, A.M. (1997) Variation in temperature and dietary nitrogen affect performance of the gypsy moth (*Lymantria dispar* L.). *Physiological Entomology*, 22, 55–64.



Forest Experiment Station.

Marden, J.H. (2000) Variability in the size, composition, and function of insect flight muscles. *Annual Review of Physiology*, 62, 157–178.

Minter, M., Pearson, A., Lim, K.S., Wilson, K., Chapman, J.W. and Jones, C.M. (2018) The tethered flight technique as a tool for studying life-history strategies associated with migration in insects. *Ecological Entomology*, 43, 397–411.



Naranjo, S.E. (2019) Assessing insect flight behavior in the laboratory: A primer on flight mill methodology and what can be learned. *Annals of the Entomological Society of America*, 112, 182–

199.

Parker, W.E. and Gatehouse, A.G. (1985) The effect of larval rearing conditions on flight performance in females of the African armyworm, *Spodoptera exempta* (Walker) (Lepidoptera: Noctuidae).

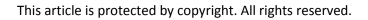
Bulletin of Entomological Research, 75, 35–48.



Parry, D., Spence, J.R. and Volney, W.J.A. (1998) Budbreak phenology and natural enemies mediate survival of first-instar forest tent caterpillar (Lepidoptera: Lasiocampidae). *Environmental Entomology*, 27, 1368–1374.

ips, B.L., Brown, G

Phillips, B.L., Brown, G.P., Webb, J.K. and Shine, R. (2006) Invasion and the evolution of speed in toads. *Nature*, 439, 803.



Pujol, B. and Pannell, J.R. (2008) Reduced responses to selection after species range expansion.

Science, 321, 96.

211–226.

Rhainds, M. (2010) Female mating failures in insects. Entomologia Experimentalis et Applicata, 136,

Robinet, C., Liebhold, A., and Gray, D. (2007) Variation in developmental time affects mating success and Allee effects. *Oikos*, 116, 1227–1237.

Schäfer, M.A., Berger, D., Rohner, P.T., Kjaersgaard, A., Bauerfeind, S.S., Guillaume, F., *et al.* (2018) Geographic clines in wing morphology relate to colonization history in New World but not Old World populations of yellow dung flies. *Evolution*, 72, 1629–1644.

Sexton, J.P., Strauss, S.Y. and Rice, K.J. (2011) Gene flow increases fitness at the warm edge of a species' range. *Proceedings of the National Academy of Sciences USA*, 108, 11704–11709.

H

Sharov, A.A. and Liebhold, A.M. (1998) Model of slowing the spread of gypsy moth (Lepidoptera: Lymantriidae) with a barrier zone. *Ecological Applications*, 8, 1170–1179.

Sharov, A.A., Liebhold, A.M. and Ravlin, F.W. (1995) Prediction of gypsy moth (Lepidoptera: Lymantriidae) mating success from pheromone trap counts. *Environmental Entomology*, 24, 1239– 1244.

Shi, J., Chen, F. and Keena, M.A. (2015) Differences in wing morphometrics of *Lymantria dispar* (Lepidoptera: Erebidae) between populations that vary in female flight capability. *Annals of the Entomological Society of America*, 108, 528–535.

Shirai, Y. (1991) Seasonal changes and effects of temperature on flight ability of the diamondback moth, *Plutella xylostella* (L.) (Lepidoptera: Yponomeutidae) *Applied Entomology and Zoology*, 26,

107–115.



Tammaru, T. and Haukioja, E. (1996) Capital breeders and income breeders among Lepidoptera: Consequences to population dynamics. *Oikos*, 73, 561–564.

Tammaru, T., Kaitaniemi, P. and Ruohomäki, K. (1996a) Realized fecundity in *Epirrita autumnata* (Lepidoptera: Geometridae): Relation to body size and consequences to population dynamics. *Oikos*,



Tammaru, T., Ruohomäki, K. and Saikkonen, K. (1996b) Components of male fitness in relation to body size in *Epirrita autumnata* (Lepidoptera, Geometridae). *Ecological Entomology*, 21, 185–192.

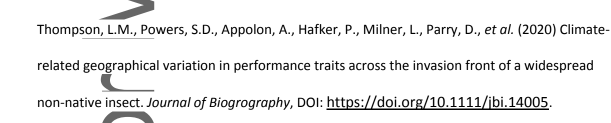
Tardif, R. and Secrest, J.P. (1970) Devices for cleaning and counting eggs of the gypsy moth. *Journal* of *Economic Entomology*, 63, 678–679.

Teder, T., Vellau, H. and Tammaru, T. (2014) Age and size at maturity: A quantitative review of diet-

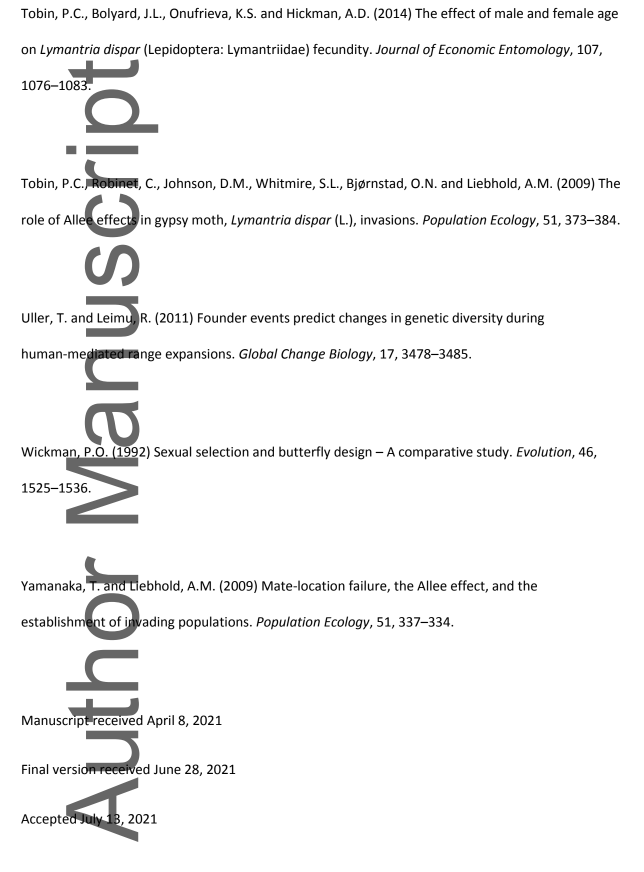
d Hastings, A. (2005) Allee effects in biological invasions. *Ecology Letters*, 8, 895–908.

induced reaction norms in insects. *Evolution*, 68, 3217–3228.

Taylor, C.M.



Tobin, P.C. and Blackburn, L.M. (2014) Long-distance dispersal of the gypsy moth (Lepidoptera: Lymantriidae) facilitated its initial invasion of Wisconsin. *Environmental Entomology*, 37, 87–93.



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Supporting Information

Appendix 1 Detailed flight mill design.

Fig. S1 Microcontroller schematic.

Table S1 Formulations for gypsy moth lab diet with varying quantities of protein. High proteindiet is equivalent to the standard gypsy moth lab diet first described in Doane and McManus(1981). Protein content was modified by removing protein components (wheat germ and casein)and adding equal volumes of cellulose to maintain consistent proportions of constituentingredients by volume. Ingredient quantities that deviated from the standard recipe are in bold.Manufacturer information for each diet ingredient is included in the last column.

Author

bt				-
Flight capacity metric	Mean ± s.d.	Median	Minimum	Maximum
Total distance (km)	19.96 <u>+</u> 16.74	15.98	1.24	68.22
Maximum speed (m/s)	1.21 ± 0.37	1.21	0.32	1.93
Wing length (mm)	2.06 ±0.21	2.03	1.64	2.53
Q				
Aut				

 Table 1 Summary statistics for dependent (total distance, maximum speed attained) variables and wing length.

Table 2 Intercept and slope coefficients for total distance (m) mixed linear regression. Total distance was square root

 transformed to improve residual normality.

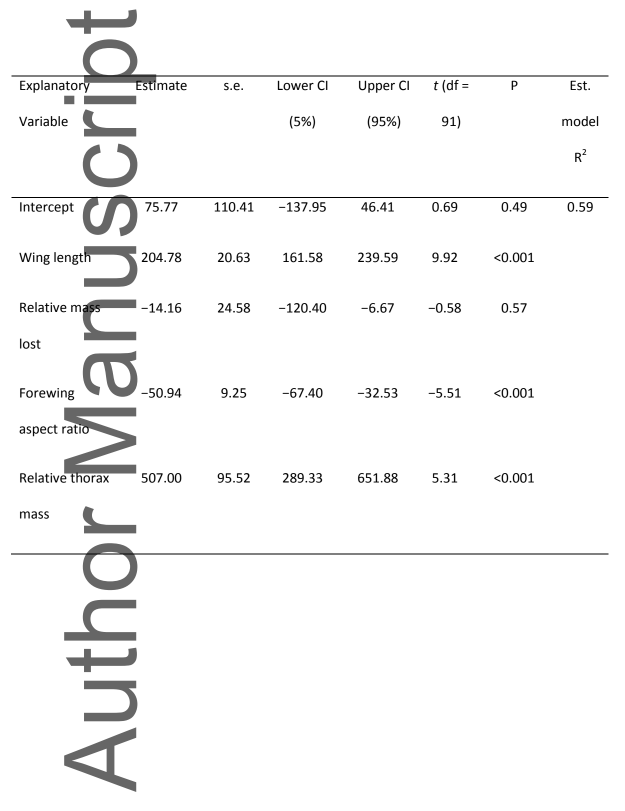


 Table 3 Intercept and slope coefficients for maximum speed (m/s) mixed linear regression.

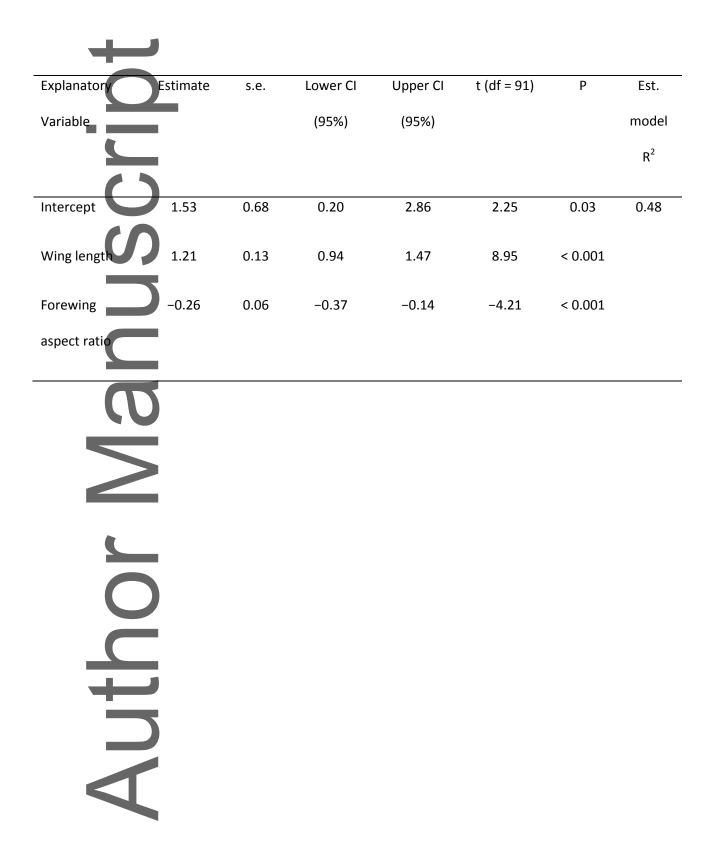


Figure Legends

Fig. 1 (A) Schematic of fixed-arm flight mill with tethered moth. Rotating arm is held above mill using opposite-facing magnets to reduce rotational friction. A magnetic sensor detected magnetic impulse at each full rotation and computer program linked to flight mill recorded each rotation with a timestamp. (B) Male gypsy moth with inverted-y shaped harness glued to descaled first abdominal segment. Image was taken following 24 h flight assay.

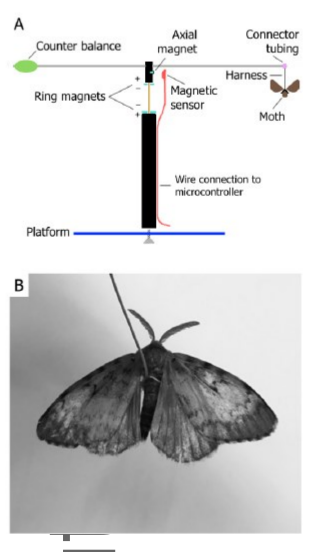


Fig. 2 Scatter plots for the effect of wing length, forewing aspect ratio, relative thorax mass, and relative mass lost during



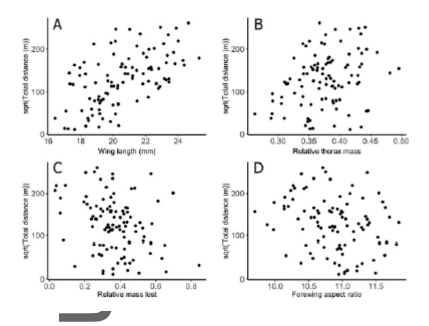


Fig. 3 Scatter plots for significant factors impacting maximum speed attained by male gypsy moth, including (A) wing length and (B) forewing aspect ratio. Raw data are plotted to visualize effects of wing length and forewing aspect ratio on maximum speed. (C) Boxplot depicting relationship between mean proportion of sustained flights initiated and wing length.

Author

