# FLIGHT ENERGETICS OF EUGLOSSINE BEES IN RELATION TO MORPHOLOGY AND WING STROKE FREQUENCY 

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## SUMMARY

Mass-specific oxygen consumption of euglossine bees during free hovering flight is inversely related to body mass, varying from $66 \mathrm{ml} \mathrm{O}_{2} \mathrm{~g}^{-1} \mathrm{~h}^{-1}$ in a $1 \cdot 0-\mathrm{g}$ bee to $154 \mathrm{ml} \mathrm{O}_{2} \mathrm{~g}^{-1} \mathrm{~h}^{-1}$ in a $0 \cdot 10-\mathrm{g}$ bee. Individuals of Eulaema and Eufreisea spp. have smaller wings and higher wing stroke frequency and energy metabolism at any given mass than bees of Euglossa spp. or Exaerete frontalis. Calculated aerodynamic power requirements represent only a small fraction of the energy metabolism, and apparent flight efficiency [aerodynamic power ( $=$ induced + profile power)/power input] decreases as size declines. If efficiency of flight muscle $=0 \cdot 2$, the mechanical power output of hovering bees varies inversely with body mass from about 480 to $1130 \mathrm{~W} \mathrm{~kg}^{-1}$ of muscle. These values are 1.9 to 4.5 times greater than previous predictions of maximum mechanical power output (Weis-Fogh \& Alexander, 1977; see also Ellington, 1984c). Mass-specific energy expenditure per wing stroke is independent of body mass and essentially the same for all euglossines. Differences in energy metabolism among bees having similar body mass is primarily related to differences in wing stroke frequency. Scaling of energy metabolism in relation to mass is generally similar to the relationship for sphingid moths despite the fact that bees have asynchronous flight muscle whereas moths have synchronous muscle.

## INTRODUCTION

The mechanical power requirements of insects during hovering flight have been analysed in detail (Weis-Fogh, 1972, 1973; Ellington, 1984c). Hovering flight is a particularly costly activity because all downward movement of air necessary to support body weight must be supplied by the beating wings. In addition, inertial
forces associated with wing movement are large (Sotavalta, 1952) and, unless they are counteracted by elastic torques, may place an additional energetic burden on the flight muscle (Weis-Fogh, 1973). Energy metabolism, or power input ( $\mathrm{P}_{\mathrm{i}}$ ), during hovering should reflect mechanical power requirements (the product of metabolism and muscle efficiency should yield the total mechanical power output, $\mathrm{P}_{\mathrm{o}, \mathrm{m}}$ ). Calculation of all mechanical power requires extremely detailed kinematic and morphometric data and relies on a variety of unproven assumptions about how aerodynamic forces are generated (see Ellington, 1984a, b for detailed discussion; also Zarnack, 1972; WeisFogh, 1973). Estimates of power requirements based on metabolic data require fewer measurements and assumptions and therefore are useful for providing an independent estimate of mechanical values of flight cost.
The energy metabolism of insects during hovering flight represents the highest rate of sustained aerobic energy expenditure in the animal kingdom. The flight metabolism of Lepidoptera, which have synchronous muscle (i.e. a constant phase relationship between impulses from the central nervous system and contraction of flight muscle; Josephson, 1981) has been relatively well characterized (Zebe, 1954; Heinrich, 1971; Heinrich \& Casey, 1973; Casey, 1976a,b, 1980, 1981a,b; Bartholomew \& Casey, 1978). Few data are available, however, for the flight metabolism of hovering insects having asynchronous muscle. Performance of these insects is of particular interest for comparison with Lepidoptera because their wing stroke frequency is mechanically determined by the morphology of the wings and the thorax (Sotavalta, 1952; Greenewalt, 1960). Among Hymenoptera, hovering metabolism has been measured in bumble-bees (Heinrich, 1975), honey-bees (Heinrich, 1980; Withers, 1981) and carpenter bees (Nicholson \& Louw, 1982; Chappell, 1982), but little concurrent information on other flight characteristics or morphology is available from the same studies. Consequently, few generalizations are currently possible concerning the effects of morphology on the flight energetics of these insects.

The euglossine (orchid) bees are particularly interesting because males routinely hover for long periods and they vary in mass from less than 60 mg to well over 1 g . The euglossines are structurally rather similar except for size and degree of pubescence, but the tribe includes several genera that vary in details of ecology and in morphology (Kimsey, 1982; May \& Casey, 1983). The present study was conducted to determine the effects of body size on flight metabolism in a group of taxonomically related insects having asynchronous flight muscle and to examine the interrelationships between energy metabolism, body morphology, wing stroke frequency and mechanical power requirements.

## MATERIALS AND METHODS

This study was conducted at the Barro Colorado Island research station $\left(9^{\circ} 10^{\prime} \mathrm{N}\right.$, $79^{\circ} 50^{\prime}$ W) of the Smithsonian Tropical Research Institute, Panama, between 15 May and 5 June, 1981. All data were obtained from male bees within 1 h after they were netted at cardboard squares impregnated with cineole, methyl salicylate or skatol.

## Morphometrics

Body mass ( M ) and thoracic mass ( $\mathrm{M}_{\mathrm{th}}$ ) were measured to the nearest 0.1 mg on an analytical balance. Side-to-side thoracic diameter ( $\mathrm{L}_{\mathrm{th}}$ ) was measured to the nearest 0.1 mm with Vernier calipers. To measure wing length ( $\mathrm{L}_{w}$ ) and area ( $\mathrm{A}_{w}$ ) one pair of wings was placed between two glass slides and projected at known magnification using a photographic enlarger. The outline of the wing was traced onto paper and the wing length was measured with a ruler. The wing outlines were then cut out and the area measured to the nearest $0.1 \mathrm{~cm}^{2}$ with a Li-Cor area meter.

## Oxygen consumption

The bees were placed in a small glass jar and were allowed to warm up (determined by vigorous abdominal pumping movements) spontaneously or after mild agitation. When the bees took off they were placed in a larger jar ( 218,452 or 825 ml , depending on the size of the bee) where they flew for at least 2 min, timed by stopwatch. In most cases, the bees hovered continuously for the desired period. Occasionally, a bee was reluctant to remain aloft and the jar was shaken to cause it to resume flight.
Hypodermic syringes were used to remove gas samples from the flight jar through a three-way stopcock cemented to the lid. We lined the inside of the lid with Parafilm ${ }^{\text {® }}$ to ensure an airtight seal. Oxygen concentration of the gas sample was determined using a Scholander $0 \cdot 5-\mathrm{ml}$ gas analysis apparatus (Scholander, 1947). All values for $\dot{\mathrm{V}}_{\mathrm{O}_{2}}$ were converted to STPD. Energy expenditure was calculated assuming an energy equivalent of $20.1 \mathrm{~J} \mathrm{ml} \mathrm{O}_{2}^{-1}$. We assume that resting metabolic rate represents a negligible fraction of the energy metabolism during hovering.

## Wing stroke frequency

Immediately after gas samples had been taken from the jar and while the bees continued to hover, the lid was removed and the jar top covered with an insect net. The flight tone was recorded on a Sony TCM-600 cassette recorder by holding a directional microphone at the mouth of the jar. Wing stroke frequency $(n)$ was determined by playing the tape back on a Tektronix 5113 storage oscilloscope.

## Data analysis

Linear regression analysis was performed on the data using either the statistical applications package for a DEC MINC (LSI-11) computer or an SAS regression procedure on the Rutgers University IBM 370 . Slopes and elevations of regression were compared by analysis of covariance. Although data pertain to a single tribe of the Apidae, the morphology of different bee species in our sample was not homogeneous (Kimsey, 1982; May \& Casey, 1983). We therefore first analysed the interrelations among morphological and energetic parameters for all bees, then performed similar analyses for subgroups of bees having close taxonomic and/or morphological affinity. For Exaerete frontalis too few data over too narrow a size range were available to make
regression analysis meaningful. We present average data for this species and include it with the genus Euglossa in some regression analyses to differentiate these bees from the morphologically different bees of the genera Eulaema and Eufriesea.

## RESULTS

## Morphology

The euglossine bees that we examined fall into one of two superficially distinct types. Individuals of the genera Eulaema or Eufriesea are black or black and yellow and have pubescent thoraces, resembling bumble-bees rather closely. Individuals of the genera Euglossa and Exaerete lack external pubescence and are brightly coloured metallic blue or green. This grouping appears to reflect taxonomic affinity (Kimsey, 1982) and is reflected in our morphometric analysis (see below).

The range of body mass of the genera in our sample is not the same. The Euglossa spp. range in size from about 50 to 250 mg . Exaerete frontalis varies in mass from about 600 to 800 mg . In the genera Eulaema and Eufriesea, mass varies from about 350 mg to well over 1 g and does not overlap the mass of individuals of the genus Euglossa (Table 1). As a consequence of the lack of overlap in size, we are forced to compare characteristics of different bees by extending their respective regression lines well beyond their normal size range (e.g. Fig. 1). While this sometimes indicates morphological or functional differences, it is often difficult to demonstrate statistically significant differences due to the large increases in $95 \%$ confidence interval for the equation beyond the size range for which data are available.

In both groups of bees thoracic mass is approximately a constant fraction of body mass (Table 2) and is proportionately slightly larger in the glabrous bees ( $42 \%$ of body mass) than in the pubescent bees ( $38 \%$ of body mass). Since thoracic mass varies directly with body mass, we assume that mass of flight muscle is a constant $15 \%$ of total body mass, based on data presented by Greenewalt (1962).

For a given body mass, the wings of the Eulaema-Eufriesea group are shorter and smaller in area than those of the glabrous species (Fig. 1; Table 2). In neither group is geometrical similarity apparent; both groups exhibit significantly higher slopes of wing length in relation to body mass than the predicted value of 0.33 . Exaerete frontalis had longer wings than either Eulaema type bees of the same size or predicted values for Euglossa (Fig. 1A).

The difference in wing area between Euglossa spp. and the Eulaema-Eufriesea group is inversely related to body mass. Consequently, if the relationship of wing area to mass for Euglossa is extrapolated into the Eulaema size range, wing area of the two groups is similar at a body mass of about 900 mg and the regressions intersect at the middle of the Eulaema distribution (Fig. 1B). If the Eulaema regression is extrapolated to the Euglossa spp. size range, however, it falls below all points of the Euglossa distribution, i.e. the predicted wing area of the former is smaller than the latter in the same range. Exaerete frontalis has larger wing areas than observed values for Eulaema or predicted values for Euglossa (Fig. 1B). We point out these patterns
here despite the fact that they require extrapolation beyond the range of data, because they are closely correlated with patterns of variation in wing stroke frequency and flight metabolism as discussed below.

The relationships of wing size to body mass indicate that shape changes with mass, both among the euglossines as a whole and within the various sub-groups. If geometrical similarity were preserved, wing length should vary with the $1 / 3$ power of body mass and wing area with the $2 / 3$ power of mass. When all bees are considered together, both wing length and area scale according to a significantly higher power than predicted (Table 2). Similar trends in scaling exist for all the groupings in Table 2, and these trends are significant for wing area in Eulaema-Eufriesea and for both area and length when Euglossa and Exaerete are considered together. It appears that large euglossine bees have larger wings than would be predicted by the assumption that they are geometrically similar to related small bees. Wing loading (the ratio of

Table 1. Morphometric parameters, wing stroke frequency and oxygen consumption of euglossine bees

body mass to wing area) is not significantly correlated with body mass within groups or in all bees (Table 2).

## Wing stroke frequency

The relatively small wings of the euglossine bees operate at high frequencies $(80-250 \mathrm{~Hz})$ during hovering. In both the Eulaema group and in Euglossa, $n$ is strongly inversely related to wing length and body mass and is not correlated with wing loading (Table 3). The magnitude and scaling of $n$ on wing length are generally similar to the values for insects in Greenewalt's (1962) group I classification (strongest fliers).

In both glabrous and pubescent bees, $n$ is tightly coupled with each morphological parameter except wing loading and sufficiently different between the two groups that the regression for all bees is usually less well correlated and shows a lower slope than the regression for a single group (Table 3). For example, Fig. 2A illustrates the relationship of frequency to body mass for all bees, regardless of taxon. Fig. 2B shows the same data except that regressions are calculated separately for different groups.

Table 2. Linear regression for log-transformed values of morphometric parameters in relation to body mass where $\log y=m \log x+\log b$

| Taxa | Slope (+s.e.) | Intercept | $r^{2}$ | $P$ | $N$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Thoracic diameter (cm) |  |  |  |  |  |
| Eulaema, Eufriesea | $0.284(+0.021)$ | -0.022 | 0.918 | $0 \cdot 0001$ | 19 |
| Euglossa | $0.320(+0.016)$ | -0.048 | 0.975 | 0.0001 | 12 |
| Euglossa, Exaerete | $0.386(+0.010)$ | 0.014 | 0.990 | 0.0001 | 16 |
| All bees | $0.361(+0.0082)$ | -0.005 | 0.983 | 0.0001 | 35 |
| Thoracic mass (g) |  |  |  |  |  |
| Eulaema, Eufriesea | $0.935(+0.047)$ | $-0.388$ | 0.958 | 0.0001 | 19 |
| Euglossa | $1.045(+0.066)$ | -0.388 | 0.966 | 0.0001 | 11 |
| Euglossa, Exaerete | $1.034(+0.034)$ | -0.380 | 0.967 | 0.0001 | 15 |
| All bees | $1.062(+0.017)$ | -0.364 | 0.992 | $0 \cdot 0001$ | 34 |
| Wing length (cm) |  |  |  |  |  |
| Eulaema, Eufriesea | $0.457(+0.037)$ | $0 \cdot 362$ | 0.900 | 0.0001 | 19 |
| Euglossa | $0.419(+0.039)$ | $0 \cdot 370$ | 0.922 | 0.0001 | 12 |
| Euglossa, Exaerete | $0.535(+0.034)$ | 0.455 | 0.946 | 0.001 | 16 |
| All bees | $0.422(+0.016)$ | 0.370 | 0.954 | $0 \cdot 0001$ | 35 |
| Wing area ( $\mathrm{cm}^{2}$ ) |  |  |  |  |  |
| Eulaema, Eufriesea | $1.012(+0.090)$ | 0.559 | 0.881 | $0 \cdot 0001$ | 19 |
| Euglossa | $0.788(+0.040)$ | 0.484 | 0.926 | 0.0001 | 12 |
| Euglossa, Exaerete | $1.050(+0.038)$ | 0.717 | 0.991 | 0.0001 | 16 |
| All bees | $0.900(+0.036)$ | 0.569 | 0.949 | $0 \cdot 0001$ | 35 |
| Wing loading ( $\mathrm{g} \mathrm{cm}^{-2}$ ) |  |  |  |  |  |
| Eulaema, Eufriesea | $-0.037(+0.101)$ | -0.561 | 0.008 | NS | 19 |
| Euglossa | $0.173(+0.064)$ | -0.506 | 0.422 | 0.023 | 12 |
| Euglossa, Exaerete | $-0.051(+0.038)$ | $-0.717$ | $0 \cdot 115$ | NS | 16 |
| All bees | $0.100(+0.038)$ | -0.566 | $0 \cdot 169$ | $0 \cdot 014$ | 35 |
| NS, not significant. |  |  |  |  |  |



Fig. 1. (A) The relationship of wing length to body mass in Euglossa spp. (O), Eulaema-Eufriesea spp. ( $O$ ), and Exaerete frontalis ( $\star$ ). Each point in this and subsequent figures represents a separate individual. Lines are fitted by the method of least squares; Solid line, Euglossa spp.; long-dashed line, Euglossa spp. plus Exaerete frontalis; short-dashed line, Eulaema and Eufriesea spp. (B) The relationship of wing area to body mass (symbols as in Fig. 1A).

Table 3. Linear regression for log-transformed data relating wing stroke frequency to various morphological parameters

| Taxa | Slope (+s.e.) | Intercept | $r^{2}$ | $P$ | $N$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Wing length (cm) |  |  |  |  |  |
| Eulaema, Eufriesea | $-1.197(+0.079)$ | $2 \cdot 413$ | 0.931 | 0.0001 | 15 |
| Euglossa | $-0.681(+0.150)$ | $2 \cdot 291$ | 0.674 | 0.0011 | 12 |
| Euglossa, Exaerete | $-0.914(+0.44)$ | 2.28 | 0.96 | 0.0001 | 16 |
| All bees | $-0.863(+0.048)$ | $2 \cdot 304$ | 0.902 | 0.0001 | 35 |
|  | Body mass (g) |  |  |  |  |
| Eulaema, Eufriesea | $-0.535(+0.0064)$ | 1.982 | 0.802 | 0.0001 | 15 |
| Euglossa | $-0.260(+0.080)$ | 2.062 | 0.515 | 0.0086 | 12 |
| Euglossa, Exaerete | $-0.451(+0.03)$ | 1.88 | 0.93 | 0.001 | 16 |
| All beer | $-0.347(+0.032)$ | 1.992 | 0.785 | 0.0001 | 35 |
|  | Wing loading ( $\left.\mathrm{g} \mathrm{cm}^{\mathbf{- 2}}\right)$ ) |  |  |  |  |
| Eulaema, Eufriesea | $-0.600(+0.315)$ | $2 \cdot 430$ | 0.176 | NS | 15 |
| Euglossa | $-0.126(+0.427)$ | 2.213 | 0.009 | NS | 12 |
| Euglossa, Exaerete | $1.86(+0.69)$ | $3 \cdot 46$ | 0.316 |  | 16 |
| All bees | $0.0083(+0.282)$ | $2 \cdot 15$ | $0 \cdot 000027$ | NS | 35 |
| S, not significant. |  |  |  |  |  |



Fig. 2. (A) The relationship of wing stroke frequency to body mass in euglossine bees. (B) The relationship of wing stroke frequency to body mass in Eulaema-Eufriesea app. (\%) and in Euglossa spp. and Exaerete frontalis ( $O$ ). Regression equations and statistics describing these relations are given in Table 3.

The regression in Fig. 2A best describes the relationship of frequency to body mass for the bees as a group, but it is clear that significant differences exist in stroke frequency between groups (Table 3; Fig. 2B). For example, analysis of covariance


Fig. 3. The relationship of oxygen consumption to body mass in euglossine bees during free hovering flight. Symbols as in Fig. 2B.
comparing Euglossa spp. with the Eulaema-Eufriesea group shows that the slopes of $\log n$ vs $\log$ mass are not statistically different but the Euglossa regression is significantly lower ( $P<0 \cdot 05$ ). The difference is even more marked if Exaerete is included with Euglossa. Differences in wing stroke frequencies of various bees are closely related to differences in their respective wing morphology. Exaerete frontalis with its large wings has the lowest $n$ (Fig. 2B). The glabrous bees have lower frequencies than the pubescent bees (Fig. 2B) and as body mass increases, predicted $n$ for Euglossa becomes more similar to observed values for Eulaema spp.

## Energy metabolism

Oxygen consumption during continuous hovering flight is strongly correlated with body mass to the $0 \cdot 64$ power if all bees are considered together (Fig. 3; Table 4). The exponent is significantly different from $0.75(P<0.05)$. The slope of $\log \dot{\mathrm{V}}_{\mathrm{O}_{2}}$ on $\log \mathrm{M}$ for Euglossa is higher than for Eulaema-Eufriesea. The difference is not significant but its magnitude is sufficient that the possibility of different scaling should be left
open. The elevations of the regressions do differ significantly ( $P<0 \cdot 05$ ), confirming that at a common mass the pubescent species would be expected to have higher $\dot{\mathrm{V}}_{\mathrm{O}_{2}}$. Exaerete frontalis is not statistically distinguishable from Eulaema-Eufriesea, although data for all Exaerete individuals fall below the predicted value for the Eulaema group.

Energy metabolism during flight is a function of wing stroke frequency and the energy expended per wing stroke ( $\mathrm{E}_{n}$ ). At any given body mass, wing stroke frequency and energy metabolism are greater in the pubescent bees than in the others (Figs 2B, 3). Stroke energy, obtained by dividing energy metabolism by wing stroke frequency, is of similar magnitude in both groups. In addition, in both groups the stroke energy scales approximately with the first power of body mass (Fig. 4; Table 5; slopes not significantly different from $1 \cdot 0$ ). Mass-specific energy expenditure for all bees amounts to $3.94 \mathrm{~mJ} \mathrm{~g}^{-1}$ stroke ${ }^{-1}$ ( +1.05 s.D.). The Eulaema-Eufriesea group had slightly lower values $(3 \cdot 78+0 \cdot 92)$ than the glabrous bees $(4 \cdot 14+1 \cdot 19)$. These data indicate that the differences in energy metabolism of bees are the result primarily of differences in wing stroke frequency.

## Power output and muscle efficiency

The power output of the flight muscles of these bees appears to be very high. If mechanical efficiency of flight muscle is a constant 0.2 and flight muscle mass is $15 \%$ of total mass, mass-specific mechanical power output of the flight muscle is inversely related to body mass and ranges from about $1130 \mathrm{~W} \mathrm{~kg}^{-1}$ of muscle $\left(170 \mathrm{~W} \mathrm{~kg}^{-1}\right.$ of body mass) in a $0 \cdot 1-\mathrm{g}$ bee to about $480 \mathrm{Wkg}^{-1}$ of muscle ( $72 \mathrm{Wkg}^{-1}$ body mass) in a $1 \cdot 0-\mathrm{g}$ bee (Table 5). These data are generally similar to those of hovering sphinx

Table 4. Linear regression for log-transformed data relating oxygen consumption to various morphological parameters

| Taxa | Slope | Intercept | $r^{2}$ | $P$ |
| :--- | :---: | :---: | :---: | :---: |
|  |  | Body mass (g) |  |  |
| Eulaema, Eufriesea | $0.365(+0.119)$ | 1.789 | 0.356 | 0.007 |
| Euglossa | $0.522(+0.246)$ | 1.700 | 0.310 | 0.060 |
| Euglossa, Exaerete | $0.580(+0.09)$ | 1.75 | 0.76 | - |
| All bees | $0.637(+0.050)$ | 1.824 | 0.833 | 0.0001 |
|  |  |  |  |  |
|  |  | Wing length (cm) |  |  |
| Eulaema, Eufriesea | $0.761(+0.247)$ | 1.509 | 0.359 | 0.0067 |
| Euglossa | $1.300(+0.541)$ | 1.240 | 0.366 | 0.038 |
| Euglossa, Exaerete | $1.17(+0.117)$ | 1.24 | 0.77 | 0.001 |
| All bees | $1.424(+0.132)$ | 1.281 | 0.778 | 0.0001 |
|  |  |  |  |  |
|  |  | Wing loading (g cm |  |  |
|  |  | 1.538 | 0.045 | NS |
| Eulaema, Eufriesea | $-0.311(+0.347)$ | 1.844 | 0.070 | NS |
| Euglossa | $0.929(+1.071)$ | 0.32 | 0.12 | NS |
| Euglossa, Exaerete | $-1.50(+1 \cdot 10)$ | 2.291 | 0.183 | 0.011 |
| All bees | $1.233(+0.454)$ |  |  |  |
| NS, not significant. |  |  |  |  |



Fig. 4. The relationship of energy expenditure per wing stroke (obtained as energy metabolısm/wing stroke frequency) to body mass; $\mathrm{m}=$ slope, $\mathrm{b}=\mathrm{y}$ intercept, $r=$ correlation coefficient. Symbols as in Fig. 2B.
moths of similar mass (Casey, 1981c) and substantially greater than estimates of maximum mechanical power output (Fig. 5) based on mechanical considerations (Weis-Fogh \& Alexander, 1977).

A variety of studies of the aerodynamic power requirements of hovering insects suggest that mass-specific induced power requirements (whether calculated by momentum theory, blade-element theory or vortex theory) are independent of body mass. Profile power output is generally similar to induced power in both magnitude and scaling (Weis-Fogh, 1972, 1973; Ellington, 1984c). Since mass-specific aerodynamic power requirements ( $\mathrm{P}_{\mathrm{o}, \mathrm{a}}$ ) are apparently size independent and massspecific flight metabolism $\left(P_{i}\right)$ is inversely related to body mass, the overall efficiency of hovering ( $\mathrm{P}_{\mathrm{o}, \mathrm{a}} / \mathrm{P}_{1}$ ) decreases markedly with body size. Over the range of body mass that we examined, flight efficiency varies from about $3.3 \%$ in a $0 \cdot 1-\mathrm{g}$ bee to about $7.7 \%$ in a $1.0-\mathrm{g}$ bee.

## DISCUSSION

## Metabolism, mass and morphology

Until recently, data for metabolism of Hymenoptera during free, hovering flight were not available. Earlier data from tethered insects (Hocking, 1953) are less than $50 \%$ of our values. Several recent measurements of the rates of energy metabolism of hovering bees are similar to ours and also support our observation that mass-specific metabolism of bees during hovering is inversely related to body mass (Table 6). Likewise, Chappell (1982) has shown that mass-specific flight metabolism of hovering carpenter bees is inversely related to body mass and similar in magnitude to our values for bees of similar body mass. Honey-bees, however, have appreciably lower $\dot{\mathrm{V}}_{\mathrm{O}_{2}}$ than euglossines of similar mass (Table 6).

Although data for metabolism of Hymenoptera during hovering flight are as high as or higher than previous data for strenuous activity of most insects and vertebrates, several lines of evidence suggest that these values are not unreasonable. First, data on heat transfer coefficients and body temperatures in the field indicate that rates of heat loss are comparable to heat production estimated from $\dot{\mathrm{V}}_{\mathrm{O}_{2}}$ measurements. For instance, in Euglossa imperialis (at $\mathrm{M}=0 \cdot 162 \mathrm{~g}$, the smallest species for which complete data on heat transfer coefficients are available), heat loss in the field would be equivalent to about $98 \mathrm{ml} \mathrm{O}_{2} \mathrm{~g}^{-1} \mathrm{~h}^{-1}$, assuming an air flow from forward flight or induced wind of $1.0 \mathrm{~m} \mathrm{~s}^{-1}$, i.e. about $90 \%$ of the expected $\dot{\mathrm{V}}_{\mathrm{O}_{2}}$ during hovering (May \& Casey, 1983). In addition, three different methods of gas analysis (paramagnetic analyser, Heinrich, 1975; polarographic analyser, Withers, 1981; Chappell, 1982;

Table 5. Allometry of various derived energetic parameters

| Derived parameter | Slope | Y-intercept | Predicted $(0.1 \mathrm{~g})$ | Predicted $(0 \cdot 1 \mathrm{~g})$ |
| :---: | :---: | :---: | :---: | :---: |
| Euglossa, Exaerete |  |  |  |  |
| Body mass-specific metabolism ( $\mathrm{W} \mathrm{kg}^{-1}$ ) | -0.42 | $2 \cdot 50$ | 832 | 316 |
| Muscle mass-specific metabolism ( $\mathrm{Wkg}^{-1}$ ) | -0.42 | $3 \cdot 32$ | 5495 | 2089 |
| Mass-specific 'mechanıcal' power output ( $\mathrm{Wkg}^{-1}$ ) | -0.42 | $1 \cdot 80$ | 166 | 63 |
| Muscle mass-specific 'mechanical' power output ( $\mathrm{Wkg}^{-1}$ ) | -0.42 | $2 \cdot 62$ | 1099 | 417 |
| Energy expenditure per wingstroke (mJ) | 1.03 | $0 \cdot 62$ | $0 \cdot 39$ | 4-17 |
| Eulaema, Eufriesea |  |  |  |  |
| Body mass-specific metabolism ( $\mathrm{Wkg}^{-1}$ ) | -0.65 | $2 \cdot 53$ | 1513 | 339 |
| Muscle mass-specific metabolism ( $\mathrm{Wkg}^{-1}$ ) | -0.65 | $3 \cdot 35$ | 10115 | 2260 |
| Mass-specific 'mechanical' power output ( $\mathrm{Wkg}^{-1}$ ) | -0.65 | 1.83 | $302 \cdot 6$ | $67 \cdot 8$ |
| Muscle mass-specific 'mechanical' power output ( $\mathrm{Wkg}^{-1}$ ) | -0.65 | $2 \cdot 65$ | 2017 | 452 |
| Energy expenditure per wingstroke (mJ) | 0.87 | $0 \cdot 54$ | $0 \cdot 47$ | $3 \cdot 47$ |
| All bees |  |  |  |  |
| Body mass-specific metabolism ( $\mathrm{W} \mathrm{kg}^{-1}$ ) | -0.37 | $2 \cdot 56$ | 851 | 363 |
| Muscle mass-specific metabolism ( $\mathrm{Wkg}^{-1}$ ) | -0.37 | $3 \cdot 38$ | 5623 | 2398 |
| Mass-specific 'mechanical' power output ( $\mathrm{Wkg}^{-1}$ ) | -0.37 | 1.86 | $170 \cdot 3$ | $72 \cdot 6$ |
| Muscle mass-specific 'mechanical' power output ( $\mathrm{Wkg}^{-1}$ ) | -0.37 | $2 \cdot 68$ | 1133 | 484 |
| Energy expenditure per wingstroke (mJ) | 0.97 | $0 \cdot 57$ | $0 \cdot 40$ | $3 \cdot 71$ |
| Predicted values for $0 \cdot 1-\mathrm{g}$ and $1 \cdot 0-\mathrm{g}$ bees are included. All other symbols as in Table 1. <br> Mechanical power calculated assuming a muscle efficiency of |  |  |  |  |

Nicholson \& Louw, 1982; Scholander apparatus, present study) all yield more or less similar results for different bees.

Differences in flight metabolism in the different subgroups of euglossine bees are generally consistent with differences in their respective morphological or functional characteristics. Higher levels of flight metabolism at any given mass in EulaemaEufriesea spp. are associated with higher wing stroke frequencies than in Euglossa or Exaerete, which in turn are a consequence of relatively smaller wings. It is tempting to suggest that the concordant departures from geometrical similarity within both groups of euglossines and the differences between groups in the magnitude of morphological and functional parameters reflect the division of the euglossines into two different 'adaptive types', corresponding to the pubescent and glabrous groups of genera. However, Exaerete frontalis has larger wings, lower $n$ and lower $\dot{\mathrm{V}}_{\mathrm{O}_{2}}$ than predicted from extrapolations of Euglossa spp. regressions (Figs 1A, B, 2B, 3) and may well form a third type. The ecological and behavioural significance of these differences awaits investigation.

It is interesting that wing loading of euglossines is a poor indicator of their wing stroke frequency and energetics. While the Eulaema-Eufriesea group have higher


Fig. 5. The relationship of mechanical power output (obtained from metabolic data assuming muscle efficiency $=0.2$ ) to body mass. Dashed line represents the relationship derived for sphinx moths (Casey, 1981c). Solid line labelled $\mathrm{P}_{\mathrm{o}, \mathrm{m}} \max$ represents the maximum mechanical power output estimated by Weis-Fogh \& Alexander (1977). Symbols as in Fig. 2B.
wing loading than the glabrous species (Table 2), within-group wing loading is independent of body mass and invariably less well correlated with $n$ or $\dot{\mathrm{V}}_{\mathrm{O}_{2}}$ than other morphological parameters (Tables 3, 4). Previous studies have attempted to explain differences in flight energetics as a result of differences in wing loading (Heinrich, 1971; Bartholomew \& Heinrich, 1973; Casey, 1976a; May, 1981). Our data suggest that both within and between closely related taxa of asynchronous fliers, wing stroke frequency is strongly coupled to energy metabolism, and that wing size and shape rather than wing loading are the primary determinants of $n$.

## Power requirements and muscle power output

Mechanical power output is related to flight metabolism by the efficiency with which the flight muscle converts energy into mechanical work. Insect flight muscle, like vertebrate skeletal muscle, has generally been assumed to have a mechanical efficiency of about $0 \cdot 2$, yielding mechanical power output ranging from 400 to $1300 \mathrm{~W} \mathrm{~kg}^{-1}$ of muscle in euglossines (Fig. 5). However, Weis-Fogh \& Alexander (1977) have calculated that the maximum mechanical power output ( $\mathrm{P}_{\mathrm{o}, \mathrm{m}} \mathrm{max}$ ) of muscle, irrespective of size, is about $250 \mathrm{~W} \mathrm{~kg}^{-1}$ of muscle, only a fraction of the values above. These authors used maximal measured metabolic rates for vertebrates and flying locusts (assuming $20 \%$ efficiency) and found that those values were in basic agreement with the calculations from their mechanical theory. Unfortunately, those metabolic rates were well below the values obtained for freely hovering insects (Kammer \& Heinrich, 1978). Based on a much wider survey of flight metabolism, Ellington (1984c) noted, as we do, the large discrepancy between $\mathrm{P}_{\mathrm{o}, \mathrm{m}}$ as calculated from metabolic rate and values based on mechanical considerations. He concluded that insect flight probably is appreciably less than $20 \%$ efficient.

If Weis-Fogh \& Alexander's (1977) model is quantitatively correct and $250 \mathrm{~W} \mathrm{~kg}^{-1}$ is a reasonable estimate of maximum mechanical power output, then the mechanical efficiency $\left(\mathrm{P}_{\mathrm{o}, \mathrm{m}} / \mathrm{P}_{\mathrm{l}}\right)$ of the flight muscle must decrease progressively with size in the Euglossa spp., varying from about $10.4 \%$ in a 1.0 -g bee to about $4.4 \%$ in a 0.1 -g bee. However, in small insects which operate at high $n$ the ability of the mitochondria to

Table 6. A comparison of oxygen consumption during hovering flight of euglossine bees with that of other Hymenoptera

|  | Body <br> mass <br> $(\mathrm{g})$ | Mean <br> $\dot{V}_{\mathbf{O}_{2}}$ <br> $\left(\mathrm{mlg}^{-1} \mathrm{~h}^{-1}\right)$ | Euglossine $^{\mathrm{b}}$ <br> $\left(\mathrm{mlg}_{\mathrm{V}^{-1}} \mathrm{~h}^{-1}\right)$ | Source |
| :--- | :---: | :---: | :---: | :---: |

[^0]Table 7. A comparison of calculated mechanical power requirements (Ellington, 1984c) for honey-bee ( 0.094 g ) and bumble-bee ( 0.205 g ) with measured energy metabolism of euglossine bees of similar body mass

|  | $\begin{array}{c}\text { Honey-bee } \\ \left(\mathrm{Wkg}^{-1}\right)\end{array}$ |  | Bumble-bee |  |
| :--- | :---: | :---: | :---: | :---: |
| $\left(\mathrm{Wkgg}^{-1}\right.$ muscle) |  |  |  |  |$)$

supply power to the contractile apparatus rather than the maximum rate at which the fibrils can work is probably the limiting factor for maximum mechanical power output (Weis-Fogh \& Alexander, 1977). Since metabolic rates are very high at small size, our data suggest that current values of $\mathrm{P}_{\mathrm{o}, \mathrm{m}}$ max may be underestimated by their approach.

If muscle efficiency rather than $\mathrm{P}_{\mathrm{o}, \mathrm{m}}$ is constant over this size range, then what work is being done? We do not have sufficiently detailed kinematic data for full analysis of aerodynamic and inertial forces in euglossines. Ellington (1984c) has calculated these parameters for a honey-bee and a bumble-bee, however. In order to compare metabolic data with mechanical power estimates (Table 7), we calculated $\dot{\mathrm{V}}_{\mathrm{O}_{2}}$ based on our equation for all bees. For bees of similar mass to the bumble-bee ( $0 \cdot 200$ ) and honey-bee ( 0.095 ) in Ellington's analysis, predicted values are 120 and 149 ml $\mathrm{O}_{2} \mathrm{~g}^{-1} \mathrm{~h}^{-1}$ (derived from values in Table 6). Aerodynamic power requirements for these animals are far below $20 \%$ of $\mathrm{P}_{\mathrm{i}}$ and are essentially size-independent. The profile power component of the total aerodynamic power may be underestimated by the blade element analysis of Weis-Fogh (1973). However, until new data are available to evaluate this suggestion, these calculations represent the best estimates of aerodynamic power currently available (see Ellington, 1984a,b,c). Despite the uncertainties in calculating the magnitude of aerodynamic work, it seems unlikely that such errors could account for this large discrepancy. Addition of inertial power brings calculated values much closer to our values predicted as $0.2 \mathrm{P}_{\mathrm{i}}$, although the scaling of mechanical power still does not conform to our lines.

Cogent arguments have been advanced to suggest that inertial torque of the wings is largely recovered through elastic storage (Weis-Fogh, 1972, 1973; Alexander \& Bennett-Clark, 1977; Ellington, 1984c). One of Weis-Fogh's (1973) arguments is questionable, however, in the light of new data. He calculated that aerodynamic plus inertial power significantly exceeded $20 \%$ of $P_{i}$, so elastic storage must be present to reduce inertial power requirements and keep efficiencies within reasonable limits. However, Weis-Fogh did not use metabolic data from freely hovering insects and he therefore severely underestimated energy expenditure during hovering. Based on
more recent data it appears that elastic storage need not be invoked to explain flight energetics since aerodynamic plus inertial power is always less than $20 \%$ of the power input. Casey (1981c) showed that predicted inertial power output, in the absence of elastic storage, would almost exactly account for the difference between calculated $\mathrm{P}_{\mathrm{o}, \mathrm{a}}$ and $20 \%$ of $\mathrm{P}_{1}$ in sphinx moths. On the other hand, if maximum power output is fixed at a level well below $20 \%$ of the metabolic rate and the mechanical efficiency of the muscle is reduced, Weis-Fogh's conclusion that elastic storage is important in flying insects is supported. The actual inertial power that could be supplied during hovering would then be represented by $\mathrm{P}_{\mathrm{o}, \mathrm{m}} \max -\mathrm{P}_{\mathrm{o}, \mathrm{a}}$. As shown in Table 7, aerodynamic power requirements for honey-bees and bumble-bees are about $185 \mathrm{~W} \mathrm{~kg}^{-1}$ of muscle, i.e. about three-quarters of the maximum aerodynamic power output predicted by Weis-Fogh \& Alexander (1977). Under these circumstances, inertial power would only be about $30 \%$ of the aerodynamic power requirements, indicating that about $90 \%$ of the energy cost of accelerating the wing mass is reclaimed by elastic storage (see Ellington, 1984 c for further discussion). If this is correct, however, the reason that power input and power output and their components scale so differently is unclear. Further information on in vivo muscle efficiency is clearly needed.

It is of interest that a similar discrepancy occurs between mechanical power requirements and metabolic rate of running vertebrates. Mechanical power requirements of runners are size-independent while mass-specific metabolism is inversely related to body mass. Taylor, Heglund, McMahon \& Looney (1980) suggested that much of the energy is used in generating force in the muscle without any change in muscle length and hence without producing work. Flight muscles might generate force without work during wing deceleration in each half stroke (Ellington, 1984c) or if there were appreciable overlap in activity of antagonistic muscles (Neville \& Weis-Fogh, 1963). The second explanation seems improbable since Neville \& WeisFogh showed little overlap at normal flight temperature even in locusts, and highly endothermic insects should experience even less (Heinrich, 1974). It remains possible that more frequent activation of the flight muscles, even without increased power output, could increase metabolic rate in small insects either due to increased cycling of actin-myosin crossbridges or increased muscle activation power requirements (Heglund, Fedak, Taylor \& Cavagna, 1982). Another alternative, as suggested by Ellington (1984c), is that elastic recovery of energy may be less efficient in small insects, Here, too, more data on the functioning of the flight system are required.

## Comparison of Hymenoptera with Lepidoptera

The flight muscles of Hymenoptera are asynchronous while those of the Lepidoptera are synchronous. Consequently, a comparison of the flight energetics of these two groups is particularly interesting given the differences in muscle biochemistry, histology and contraction mechanics (Kammer \& Rheuben, 1981; Josephson, 1981; Pringle, 1981). Wing stroke frequency of insects is closely related to wing length regardless of whether each muscle contraction is determined by impulses from the central nervous system (synchronous muscle) or by the mechanical resonant characteristics of the wing and thorax (asynchronous) (Greenewalt, 1962).

The relationship of energy metabolism to body mass for all euglossine bees is very similar to that exhibited by sphinx moths (Fig. 6). Over the range of size where these groups overlap (approximately 300 mg to 1.0 g ) their energy metabolism differs by less than $10 \%$. Both the bees and the sphingids are high performance fliers that operate at wing stroke frequencies that are higher than most insects of comparable mass (e.g. see Greenewalt, 1962) and that routinely hover for extended periods in nature. Both groups have $\dot{\mathrm{V}}_{\mathrm{O}_{2}}$ values about $30 \%$ higher than the saturniid moths, which have larger wings, lower stroke frequency and which hover less readily.

While energy metabolism of the sphingids and euglossines is essentially the same, differences in morphology between the groups result in differences in required wing stroke frequencies. Although sphingids operate at high wing stroke frequencies $(25-66 \mathrm{~Hz})$ relative to other synchronous fliers, the bees with their smaller wings operate at still higher frequencies (about $80-240 \mathrm{~Hz}$; Table 1; Fig. 2) over the same range of body mass. For the sphingids, $\mathrm{E}_{n}$ is about $12-25 \mathrm{~mJ} \mathrm{~g}^{-1}$ body mass while for


Fig. 6. Energy metabolism of hovering insects in relation to body mass. Individual data shown only for moths: closed circles are sphinx (hawk) moths, open circles are saturniids, open squares are other taxa; Diagonal slashes indicate data from Zebe (1954), all other data are from Bartholomew \& Casey (1978).
bees it is only about $4 \mathrm{~mJ} \mathrm{~g}^{-1}$. The trade-off between frequency and $\mathrm{E}_{n}$ in these two groups is a consequence both of mechanical differences caused by different wing morphology and physiological limitations that apparently preclude insects with synchronous, non-fibrillar flight muscle from attaining stroke frequencies much above 120 Hz . Thus sphingids are precluded from operating at very high stroke frequencies but their relatively larger wings compared with the bees make such high frequencies unnecessary since more air can be accelerated per wing stroke.

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[^0]:    - Mean values except $X$. capensis.
    ${ }^{\text {b }}$ Predicted from body mass based on equation for all bees, Table 4.
    ${ }^{c} P$. C. Withers, personal communication.
    ${ }^{\mathrm{d}}$ Assuming thoracic mass $=0.35 \times$ body mass.
    ${ }^{\circ}$ Estimate for $1 \cdot 5-\mathrm{g}$ bee.

