**The Effects of Vein Morphology on Elastic Wing Deformation**

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Submission date: September 2022

**Abstract:**

Unlike flying vertebrates, insect flight relies on the passive, elastic deformation of their wings to generate lift. To do so, wing aerofoil shape is distorted by the aerodynamic and inertial forces acting on their unique and intricate mechanical properties. The complex interplay between these factors complicate previously proposed mathematical models that try to understand the benefits native to more rigid or more elastic wings. In our study we focus on elastic deformation and lift generation as a function of vein morphology and wing flexural rigidity. By synthetically replicating the wings of the Protaetia Cuprea and solely fortifying their veinal network, we are able to directly assess the implications of flexural stiffness on wing deformation and its aerodynamic influences. Our results conclude that a change to vein cross-sectional area directly effects wing rigidity and thus, deformation capabilities. Furthermore, our results provide insight pertaining to important factors to take into consideration when synthetically remodeling authentic flapping wings.

**1. Introduction:**

For millions of years, wings have been subject to evolution through natural and sexual selection. Their divergence and gradual morphological advances directly impact both maneuverability and the energetic efficiency of flight in different species1. Unlike flying vertebrates, insects lack musculature in their wings which otherwise offer active and precise, aerodynamically favorable alterations during flight2. Instead, during flapping, insects rely on the more passive elastic deformation of their thin, membranous wings. In this case, the aerofoil shape of the malleable insect wings is controlled by their interactions with inertial and aerodynamic forces as they flap3.

Passive deformation in insects is further regulated by mechanical properties throughout the wings such as a relatively rigid veinal network, used to provide deformation-supporting and deformation-limiting areas2. Different venation patterns thus translate into differences in wing twist and camber, pushing species to adopt different niches depending on what their wings are better suited for. One such example was seen when comparing between interspecific wing-vein arrangements and the resulting elastic deformation within the scarab beetle family. Rose chafers, which require precision-landing to feed on flowers, were found to have a subtle difference in vein arrangement leading to a more compliant trailing edge (TE). In contrast, dung beetles, who use fast flight to reach a general feeding ground area, expressed a shift in wing-vein placement resulting in a more locally rigid wing with less deformation1,4. It is at this intersection that controversy still remains whether these elastic deformations improve or stunt aerodynamic output in comparison to a more rigid alternative.

In an attempt to understand the functional significance of wing venation on flexural rigidity (EI), Combes and Daniels took a quantitative approach using a simple cantilever beam model to measure the flexural rigidity of insect wings. They concluded that flexural rigidity, and consequently, span-wise and chord-wise deflection scales with the size of the individual5. Studies by Yonatan Meresman and Gal Ribak, however, debated that morphological parameters do not change evenly with changes to body size. This renders simplistic mathematical models unfavorable when trying to inspect the tradeoffs between having a more rigid or compliant wing3.

Here, we aimed to circumvent the lack of isometric growth in free-flying models in order to determine the effects of flexural rigidity on wing deformation and lift generation. By replicating the wings of Protaetia Cuprea (Rose Chafer beetle) and only modifying their veinal cross-sectional areas, we were able to directly manipulate wing rigidity while nullifying all other mechanical properties. Our study thus steers away from generic and oversimplified mathematical predictions in the hopes of understanding the complexities behind compliant versus rigid wings and their aerodynamic influence.

**2. Materials and Methods**:

**2.1 Wings**

3 sets of 3D-printed, scaled-up replicas (1:6) of Rose Chafer wings, fitted with mounting-stalks specific to the flapper device, (see section 2.2) were obtained from Or Filc in collaboration with the School of Engineering. Each wing-set represented a different veinal cross-sectional area, herein; horizontal, round, and vertical (labeled H, O and V respectively) geometry. The rest of the wings’ morphological and mechanical properties remained identical. Six natural morphological landmarks comprising wing edge-vein junctions and intervein joints: the wing tip (wt), marginal-joint (mj), radius-posterior (RP), media-posterior (MP), cubitis-anterior (CuA) and analis-anterior (AA) were marked with white dots on each wing using acrylic paint (see figure 1a.) to allow their accurate tracking using high-speed films (obstruction to flapping kinematics due to paint was considered negligible in this experiment). The seventh point, referred to as the wing-base (wb), was considered to be the gear of the flapper directly above the wing clasp.

**2.2 Flapper and Load Cell**

The Flapper (figure 1d.), a custom-made contraption whose motor is powered by a battery, was provided by Leeor Mordoch in collaboration with the School of Engineering and was secured onto the Zemic L6B-L load cell. Clasps on both sides were custom fitted to secure the synthetic wings such that the wings were sustained by screws to ensure no variation in the wing offset or wing pitch. The flapper was made to flutter the synthetic wings back and forth at 4 speeds powered by 0.3V, 0.5V, 0.7V and 0.9V while the Zemic L6B-L load cell simultaneously recorded the lift force generated by the flapping. A screwdriver was used to gently press down on the load cell at the start of each recording to indicate the starting point. Frame numbers of the videos were then converted to time stamps in relation to that of the load cell (which measured weight every 10 milliseconds) and the lift force measurements between the starting frame to the end frame were averaged.

**2.3 Filming System**

Flapping was recorded using 2 synchronized high-speed cameras (Fastcam SA-3\_120K, Photron inc.) fitted with an 85mm lens and set to film at 500 fps (Exp 1/5000, resolution 1024x1024 pixels). The cameras were positioned above the flapper on opposite ends. Two infrared LED floodlights (ELIMEC Ltd) covered by a diffusive screen were placed underneath the flapper to provide better visual contrast in the recorded films. The cameras were spatially calibrated using an 8 cm-long wand and the “easywand5” software file from the MATLAB archives.

**2.4 Data Analysis**

**Digitization**

Digitalization of the seven points in each film frame was performed with the DLTdv5 software from MATLAB. Three landmarks on the leading edge (wb, mj, and wt) were used to define a relatively constant plane throughout 3D space while four landmarks (RP, MP, CuA, and AA) defined the trailing edge (figure 2a). Digitized landmarks helped track the position and orientation of trailing edge relative to the leading edge to extract flapping kinematics and wing deformation magnitudes.

**Normalized Flapping Cycle and Lift Force**

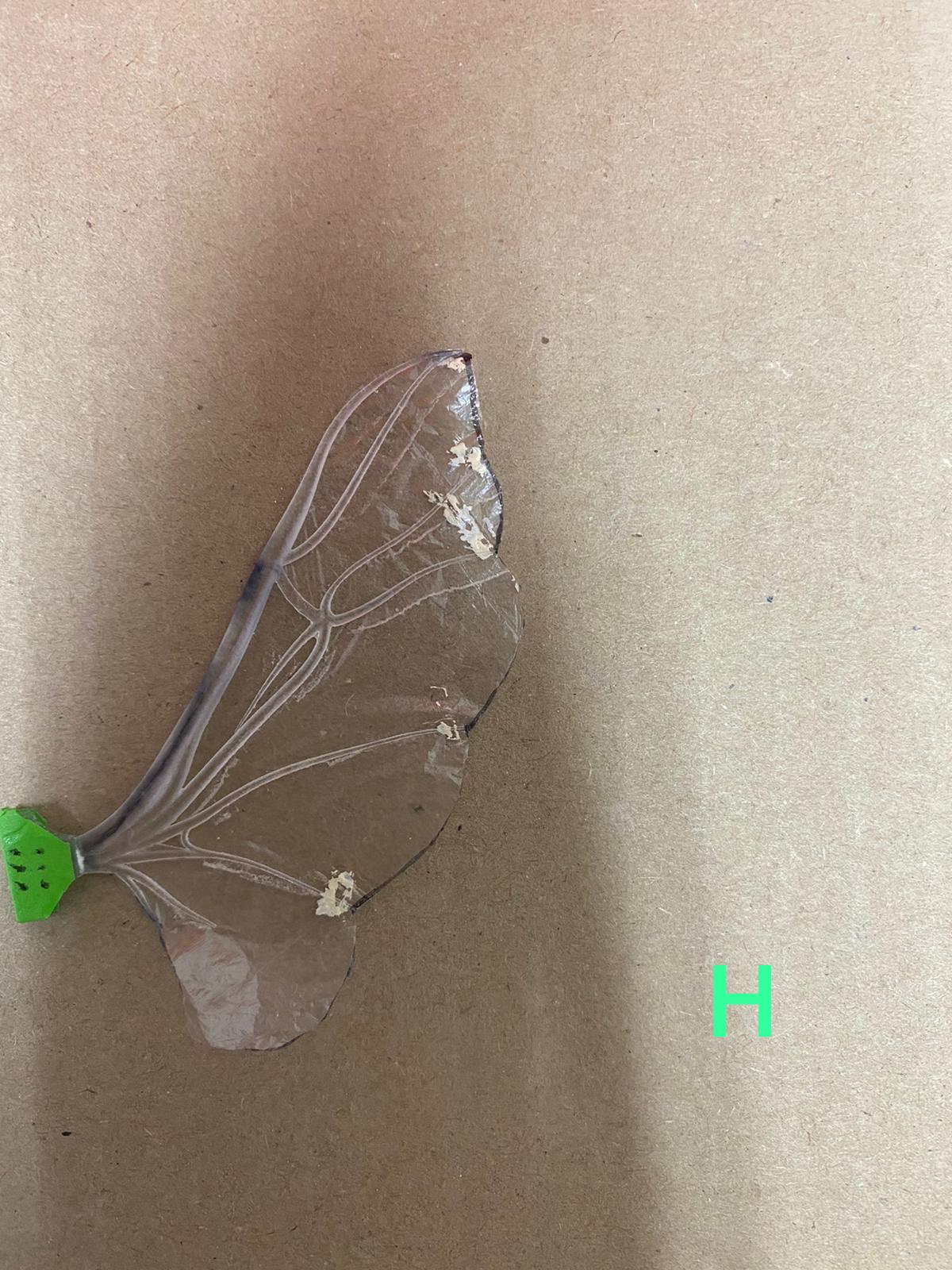
All measurements were based on three flapping cycles. To control for variance in cycle duration, we divided the serial numbers of each film frame by the number of frames within a flapping cycle to give a non-dimensional time scale of flapping cycles labeled 0-3. Lift force was calculated by the load cell measuring the weight every 10 milliseconds. All weights measured over the duration of the 3 flapping cycles were averaged together and converted to Newtons.

**Wing Deformation**

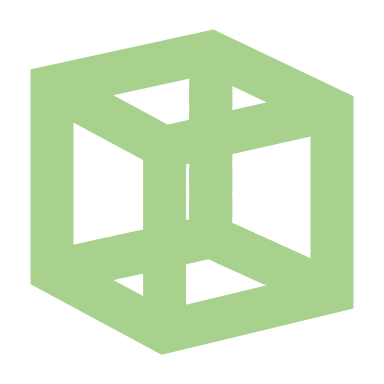
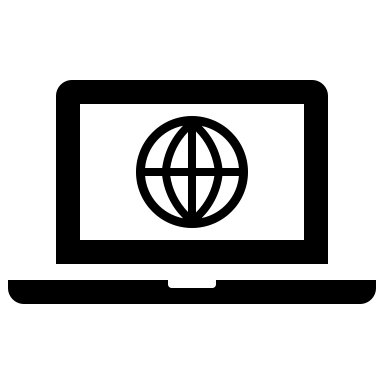
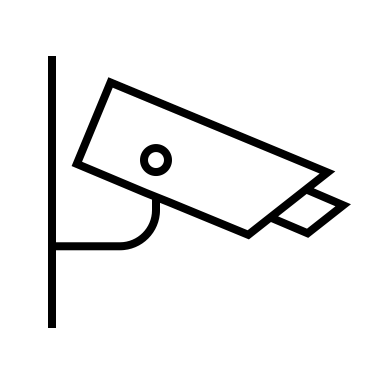
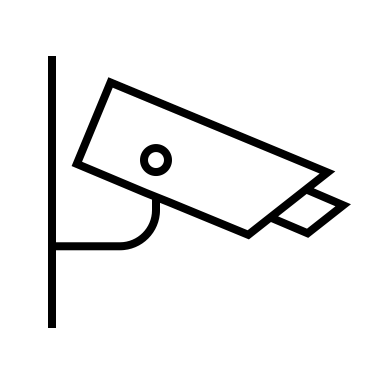
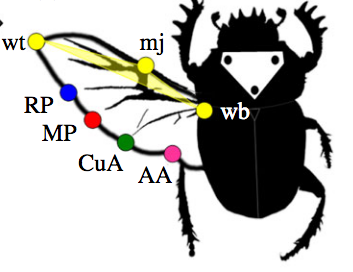
Wing deformation was measured only on the right wing in every trial (each trial consisted of flapping a pair of wings), relative to the leading edge. We defined a fixed Cartesian coordinate system using three leading-edge landmarks (wb, mj, and wt) to represent an instantaneous plane in 3D space given that the leading edge is relatively rigid. The perpendicular distance of the four trailing edge points was then calculated in relation to the leading edge to find the chordwise deflection. The deflection from the leading edge defined wing camber while the difference in magnitude between each point’s deflection define the wing-twist.

To define the coordinate system, two vectors were made. One connecting points wb to wt and one connecting wb to mj. The normal force axis (Z) was then defined by the cross-product: . Due to inaccuracies in depth perception, the DLTdv5 defined the wt vector as negative in relation to the mj vector. The resulting cross-product was thus a negative meaning that the positive normal force direction is defined as the direction of the ventral side of the wing while the negative normal force is defined as going towards to dorsal side. The chordwise, lateral axis (Y) was then defined by the cross-product: . All vectors were then converted to unit vectors. These vectors were then used as a rotation matrix to transform all data point on the wing to the wing frame of reference defined by the leading edge. The transformation of a given point P from the frame of reference of the camera to the frame of reference of the leading edge (P’) is:

The procedure was repeated for each film frame. By definition, the component of the transformed data points is the deflection out of the plane of the rigid leading edge.



c.



IR

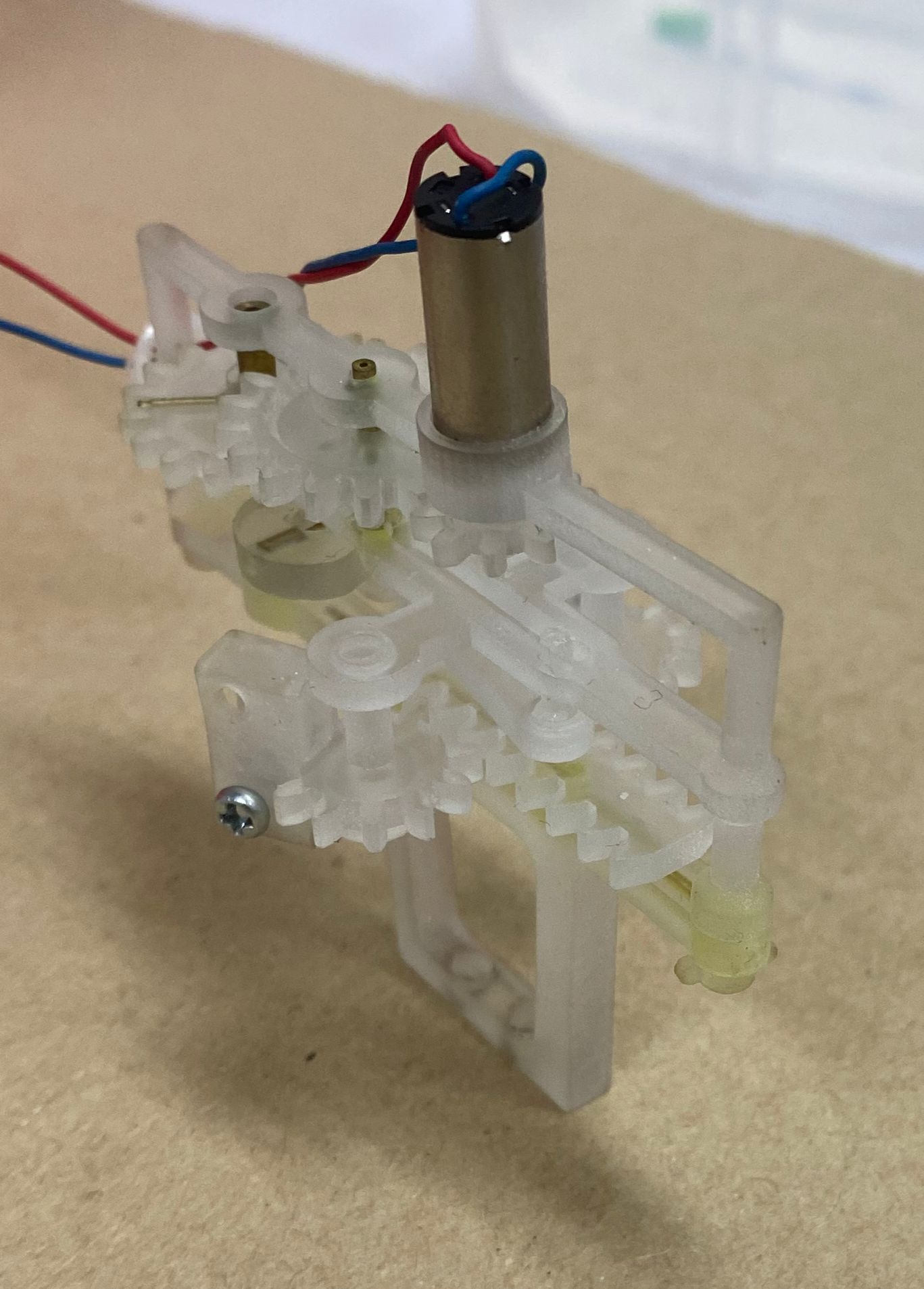
IR

DS

flapper

a.

b.



d.

**Figure 1. Methodology. (a)** An illustration obtained from Y. Meresman and G. Ribak’s paper presenting the Protaetia Cuprea’s digitized landmarks used to extract flapping kinematics and wing deformation of the trailing edge. Leading edge as the instantaneous 3D plane comprised of wb (wing-base; in our case, the gear of the flapper), mj (marginal-joint), and wt (wing tip). Trailing edge was defined as AA (Analis-Anterior vein), Cua (Cubitus-anterior vein), MP (Media-posterior vein) and RP (Radius-posterior vein). **(b)** Schematic representation of the experimental set-up. **(c)** Synthetic wing replica 1:6 scale-up of Protaetia Cuprea wing. **(d)** The flapper instrument, without the wings attached.

**3. Results:**

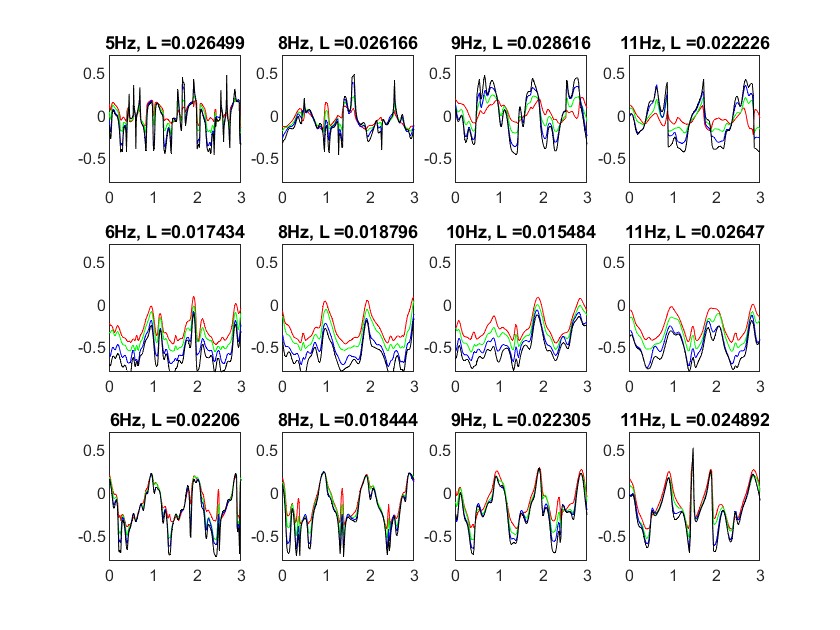
**3.1 Wing Deformation** –

Throughout the flapping cycle of all trials, the four TE landmarks (RP, MP, CuA, and AA seen in figures 1a) exhibited a clear cyclic deformation pattern consistent with the flapping kinematics. Apart from the H1-wing flapping at 8Hz (Figure 2a, first row second column) whose deflection followed the direction of the wing flap resulting in positive camber, the remainder of the trials upon downstroke (defined here as flapping in a forward direction relative to the flapper) showed TE landmarks to be deflected towards the dorsal side of the wing while upon upstroke (defined here as flapping in a backward direction towards the flapper) the landmarks deflected towards the ventral side of the wing. Chordwise deformation was thus found to always be opposite to the wing’s direction of movement resulting in negative camber irrespective of the wing types, except for the H1-wing at 8Hz. Additionally, only in H1 wings of 9Hz and 11Hz (figure 2a, row 1 columns 3 and 4) was there a symmetrical reflection about the y-axis to denote equal elasticity of both the ventral and dorsal sides of the specific wing as opposed to the remainder of the trials whose deflection never fully surpassed the LE to the ventral side.

Magnitude of the wing deflection for each point at mid-stroke relative to the LE, seen by the deviation from the y-axis 0 mark, demonstrated a clear spanwise gradient (wing twist) within the chordwise deformation. The landmark AA (represented by the black line) had the greatest deviation while RP (represented by the red line) had the least. This indicates ascending spanwise and chordwise rigidity from the proximal point AA being the least rigid to the more distal point, RP, being the most rigid. When comparing between wing sets, horizontally veined (H) wings from both trials displayed the greatest resistance to chordwise deformation while those with vertical (V-wings) and rounded (O-wings) geometry veins showed no significant differences regarding their compliance. H and O wings did, however, seem to translate to having greater spanwise gradients seen by the more robust separations between relative deformation coefficients of each landmark. This suggests a steeper increase in spanwise rigidity relative to each point in H and O wings when ascending from the proximal to distal edge.

**3.2 Wingbeat Frequency and Lift–**

Raising the wingbeat frequency (presented at the top of each graph in Hz) resulted in inconclusive fluctuations of generated lift-force hereby suggesting that wingbeat frequency had no significant effect on any of the wings’ lift generation capabilities. Even upon positive camber as noted in the H1 wing at 8Hz, there was no apparent increase to lift generation. In addition, wingbeat frequency had no apparent effect on chordwise or spanwise wing deflection when comparing between the same sets (i.e. H1 wing deformation at all frequencies in figure 1 fluctuates about the 0.5 mark).



H1-wings

O1-wings

V1-wings

- RP

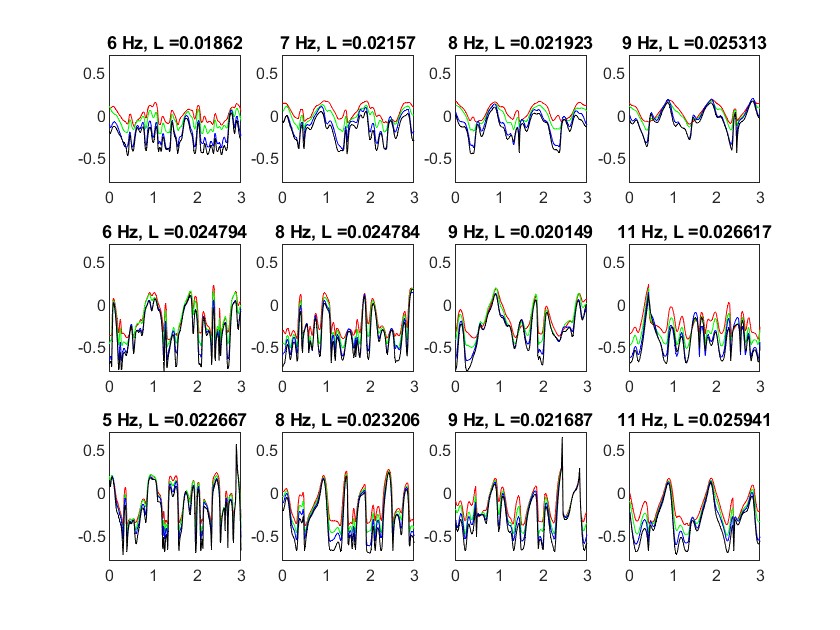
- MP

- CuA

- AA

a.

**Figure 2a. and 2b.** Trials 1 and 2 refer to flapping of different batches of the same wing sets (i.e. trial 1 uses one batch of H wings and trial 2 uses another batch of H wings). Each individual graph displays normalized deformation of the trailing edge relative to the leading edge as a function of wing set. Each row depicts a different wing set (Horizontal (H), Round (O), or Vertical (V)) and each column depicts ascending wingbeat frequency (seen on top of each individual graph in Hz). Deformation of each landmark is color coded as; RP in red, MP in green, CuA in blue, and AA in black. Lift force (L) on the top of each individual graph represents the lift force measured in Newtons.



b.

H2-wings

O2-wings

V2-wings

**4. Discussion and Conclusion:**

As mentioned in previous studies by Gal Ribak, oversimplified mathematical models incorrectly assume that larger individuals of the same species have an isometric magnification of wings3,5 . This inability to predict results by simple scaling of the model makes it difficult to calculate the change in forces that emerge from variations of wing rigidity. In an attempt to gain insight as to the benefits of wing elasticity versus wing rigidity, we recreated 3 isometric models of P. cupreia wings, where, by solely modifying the veinal cross-sectional area, we provided a method for controlled manipulation to directly alter wing compliance.

In our study, our hypothesis held true in that we observed noteworthy differences in rigidity when comparing between normalized TE deviations in relation to the LE of each wing set. Nevertheless, we found no sufficient evidence to infer a positive correlation between wingbeat kinematics coupled with varying degrees of rigidity, on lift-force generation. When raising the wingbeat frequency within the same set of wings, we would expect a steady squared proportionate increase in lift force generated, granted all other variables remain constant, seen by the lift force equation: (where L = lift force, = lift coefficient, = air density, = velocity, and A = area of the wing). This was not observed however, when comparing the frequency differences within a set, most likely due to the loss of aerofoil shape as the TE buckled under the air’s force translating to a negative camber. Unlike the documented positive wing camber in free-flying beetles which was achieved via TE deflection in the direction of the flapping to promote lift, our graphs depict an opposite trend in the synthetic wings3,4. Wing buckling thus significantly reduced the overall wing area exposed to the air, greatly impacting aerodynamic qualities and also rendered our lift force measurements inconclusive and inaccurate. On the other hand, however, the H1 wing flapping at 8Hz was an outlier suggesting that negative camber may not be the culprit for the lack of lift-generation. Even though it showed positive camber, the lift-force measured did not show any particular improvement.

Furthermore, aside from H1 wings at 9Hz and 11Hz, uniform reflection of the deformation coefficients about the y-axis was not observed. This entails dorso-ventral asymmetry potentially preventing the formation of functional aerofoil structures to promote lift both during upstroke and downstroke which is not entirely compliant with data depicting flexural stiffness of free-flying rose chafer wings1,4. It is important to note that specifically for the H-wing sets, two different batches, one of which was older, were used. This possibly caused degradation of the synthetic material in the older batch, effecting its resilience and thus also tampered with the frequency, lift and deformation we recorded between the H-wing sets.

Overall, our findings did show that veins and flexural rigidity play a significant role in wing deformation. Moreover, in order to better mimic the kinematics of free-flying rose chafer wings we see that solely manipulating the rigidity through vein geometry is insufficient. Wootton mentions the accountability that the wing cuticle as well as vein corrugation holds on wing twist and wing camber. He also stresses the necessity of experimentation using physical replicas and refining them to better resemble results recorded in free-flight experiments to overcome the limitations mathematical predictions have2,6. This study thus highlights the importance of the interaction between the complex mechanical properties found in actual wings. Ultimately, in order to find more viable and reproducible measurements pertaining to the lift force generated in response to wing rigidity, it would be beneficial to improve the capacity of the synthetic wings to resist negative camber. Likewise, improvements to the flapper should be made to make it more durable as it frequently broke down, hindering the maximum wingbeat frequency we can provide and the flapping symmetry between sets.

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