Flight in slow motion: aerodynamics of the pterosaur wing

Colin Palmer*

Department of Earth Sciences, University of Bristol, Bristol, UK

The flight of pterosaurs and the extreme sizes of some taxa have long perplexed evolutionary biologists. Past reconstructions of flight capability were handicapped by the available aerodynamic data, which was unrepresentative of possible pterosaur wing profiles. I report wind tunnel tests on a range of possible pterosaur wing sections and quantify the likely performance for the first time. These sections have substantially higher profile drag and maximum lift coefficients than those assumed before, suggesting that large pterosaurs were aerodynamically less efficient and could fly more slowly than previously estimated. In order to achieve higher efficiency, the wing bones must be faired, which implies extensive regions of pneumatized tissue. Whether faired or not, the pterosaur wings were adapted to low-speed flight, unsuited to marine style dynamic soaring but adapted for thermal/slope soaring and controlled, low-speed landing. Because their thin-walled bones were susceptible to impact damage, slow flight would have helped to avoid injury and may have contributed to their attaining much larger sizes than fossil or extant birds. The trade-off would have been an extreme vulnerability to strong or turbulent winds both in flight and on the ground, akin to modern-day paragliders.

Keywords: flight; wing membrane; pterosaurs; aerodynamics

1. INTRODUCTION

Pterosaurs were flying reptiles that lived alongside the dinosaurs throughout most of the Mesozoic (ca 220–65 Ma) [1]. Unlike their living functional counterparts, birds and bats, pterosaurs possessed a flexible, membranous wing that was supported by a single, super-elongate wing finger [1–3]. As far as is known, no other flying vertebrates have ever adopted this gross morphology and there are no direct analogues in mechanical aerodynamics, the closest being the mainsail of a sailboat [4].

Although we know from fossils with preserved soft tissues that the pterosaur wing membrane was thin (most likely of varying elasticity), with internal reinforcing fibres—perhaps surface fibres (pycnofibres) [3]—and possibly pneumatized in the regions closest to the wing bones [5], it remains difficult to determine the nature of the membrane attachment to the bony skeleton. It was probably either midway between the dorsal and ventral sides of the wing bones [6], as in bats [7], or attached entirely to their dorsal surfaces [8].

Reconstructions of pterosaur flight capabilities have varied hugely because of structural and biomechanical unknowns (e.g. [9–17]). Most have been based on assumptions of wing bone morphology from fossil evidence, combined with wing section data drawn from the pre-1950s aerodynamic literature, which are necessarily unrepresentative of actual pterosaur wing profiles. Results from more recent theoretical and experimental work on low-speed aerofoils [18] and sail boats [19,20] can improve our understanding of the relevant airflow phenomena (see electronic supplementary material), but still do not provide results that can be directly applied to possible pterosaur wing sections. Only two specific investigations [11,21] of such sections have been published, but both contain anomalous results (see electronic supplementary material). Consequently, it is not possible to quantify the effects of varying the location of the wing bone relative to the lifting surface, varying the camber of the wing section or of different wing bone cross-sections. In the absence of such data it is not possible to predict the overall flight performance with any certainty. To address this deficit, two-dimensional models of a range of wing sections were made and tested in a low-speed wind tunnel and the results used to produce comparative flight performance curves for large, generic ornithocheiridiform (Pteranodontidae, Isthiodactylidae, Anhangueridae, etc.) pterosaurs. The sources of information are described in more detail in the electronic supplementary material, but the models were made intentionally generic so that the results would be widely applicable and not species-specific.

2. METHODS

(a) Model tests

Two-dimensional rigid and flexible models of wing sections representative of different cross-section locations (figure 1) along the wing of a generic, 5.8-m-wingspan derived pterosaur were constructed and tested in a wind tunnel at appropriate values of Reynolds number ($Re_c$ see electronic supplementary material). The rigid models were made from thin, curved sheets of epoxy resin/carbon fibre composite (figure 2). Since the actual camber of the pterosaur wing membranes cannot be known with any precision, models were made with three different camber values to provide a range of results. Pterosaur wings comprised two distinct regions—the proximal region where a propatagium was present and the distal region where it was absent [1,2]. The distal region of the main wing membrane was supported by wing bones situated along the anterior margin, whereas in
was modelled with oval sections of two different sizes, used to model possible muscle tissue. The first wing phalanx reconstructions. Faired and unfaired geometries were the anterior margin, to represent different propatagium the ventral side at 20 and 40 per cent of the wing chord from region was therefore modelled with the wing bone (ulna) on orientation and point of location of the pteroid. The proximal depends upon the assumed angle of the elbow joint, and

the proximal region the wing bones lay within the margins of the membrane (figure 1). The local width of the propatagium depends upon the assumed angle of the elbow joint, and orientation and point of location of the pteroid. The proximal region was therefore modelled with the wing bone (ulna) on the ventral side at 20 and 40 per cent of the wing chord from the anterior margin, to represent different propatagium width reconstructions. Faired and unfaired geometries were used to model possible muscle tissue. The first wing phalanx (WP1) was modelled with oval sections of two different sizes,
in the two-dimensional tests, the parasitic drag of the body and the induced drag (drag owing to lift). The resulting lift and drag values were then used to calculate a glide polar curve (the variation of sink speed with forward speed [31,32]).

The parasitic drag of the body was calculated using the methodology of Bramwell & Whitfield [10] in order that comparisons could be made with their work. The induced drag was calculated using the standard aerodynamic formulation [30]:

\[ C_D = \frac{e^2}{(e^2 + A R)}, \]

where \( e \) is a constant dependent upon the geometry of the wing planform, \( A R \) the aspect ratio of the wings and \( C_L \) the wing lift coefficient. A value of \( e = 0.9 \) was used, applicable to a highly tapered wing [4]. \( A R \) was calculated from \( A R = B^2/S \) (where \( B = \) the total wing span and \( S \) the wing area).

The polar curve was calculated using the identity

\[ C_T = W/(0.5\rho S V_C^2), \]

where \( C_T \) is the resultant of \( C_L \) and \( C_D \) (the wing lift and drag coefficients), \( W \) the weight of the animal, \( \rho \) the mass density of air and \( V_C \) the airspeed. This is solved to give the airspeed vector, from which the horizontal and vertical speed components can be derived.

The resultant curve is an inverted U shape, and the maximum of the curve is the point of minimum sink. The point where it is tangential to a line through the axis is the maximum aerodynamic efficiency (\( L/D_{\text{max}} \)) and also the maximum range in still air. It is important to be aware of this distinction. At minimum sink, time in the air is maximized, whereas at \( L/D_{\text{max}} \) the range is maximized. The former matters more for behaviour that relies on soaring in rising air [31]. Minimum sink can be improved in two ways: by increasing the \( L/D_{\text{max}} \) and/or by reducing the flight speed, as moving towards the top left of the graph improves the sink rate without an increase in aerodynamic efficiency. Indeed, it is possible to improve minimum sink even as \( L/D_{\text{max}} \) reduces.

One other region of the polar curve deserves attention—the bottom left of the graph. This is where speed becomes very low and flight can only be sustained by achieving high values of lift coefficient. A precipitous drop in the polar curve reflects a sudden stall and probable loss of control. A more rounded shape reflects a more gradual and controlled transition from flight to stall.

3. RESULTS

The ulna positioned at 40 per cent of wing chord (faired or unfaired) reduced maximum lift but did not increase minimum drag, resulting in only a small reduction in aerodynamic efficiency. A more anterior (20% chord) location for this wing bone resulted in a greater reduction in performance, but in both cases the effect of the fairing was very small.

Tests with the WP1 and WP2 phalanges gave the following results. (i) With the sections on the ventral side, drag increased substantially (when compared with a cambered section alone), with little difference between the two shapes. There was almost no effect on the maximum lift (in fact it increased a little with the WP2 phalanx). (ii) A phalanx located on the dorsal side of the wing reduced the section performance substantially, increasing drag and decreasing the maximum lift coefficient. (iii) The larger the bone section relative to the width of the wing section, the greater the drag. (iv) Fairing the phalanx sections to the extent previously suggested [8] did not reduce the drag or influence the maximum lift. (v) A more extensive fairing, designed to minimize separation, reduced the drag by 35 per cent with no change in maximum lift. (vi) The flexible section had similar minimum drag but greatly increased maximum lift (>25%) (at the cost of high drag). (vii) The maximum lift coefficients produced by the rigid sections approached 2.0 and were still higher for the flexible section, values considerably in excess of those reported for birds [32,33].

4. DISCUSSION

Pterosaur wing section performance is sensitive to wing bone location and size because the bones trigger flow separation (which creates drag) but this effect is reduced when they are positioned posterior to the wing margin (on the ventral side). The humerus and ulna are the largest-diameter bones in the pterosaur wing and would have been surrounded by soft tissue, further increasing their size; so proximal regions of the wing would have potentially suffered substantial loss of performance owing to these bones. However, this effect was mitigated by the propatagium, which positioned the bones posterior to the anterior margin of the membrane, and so acted as a drag reduction device (contra [21]). Such a thin leading edge section is very sensitive to the local angle of incident flow [34], which might explain the role of the pterosaur pteroid—to vary the local incidence angle of the propatagium as the overall angle of attack of the wing changed, and thus maintain the optimum flow conditions.

When a phalanx was positioned on the anterodorsal wing margin, the performance was highly degraded. Similar results have been seen in sail/mast tests [20,35,36], together providing strong support for ventral wing finger positioning [8].

The glide polars constructed from these wing section results show the influences on the overall flight performance (figure 4). Flight efficiency is significantly inferior to previous estimates [10,12], and somewhat lower than some extant soaring birds [37–40]. However, owing to the low flight speed, the minimum sink rate (approx. 1.0 m s\(^{-1}\)) was comparable to extant birds [32,37–40] and bats [41]. As wing bone size increases (relative to the wing chord), the sink rate increases with little effect upon flight speed.

The flight performance was improved by extensive fairing of the wing bones, and, while the presence of such a fairing is entirely speculative, it may have been provided by pneumatized tissue [5]. The fairing increased the aerodynamic efficiency and flight speed, but had only a limited effect upon the minimum sink speed, the parameter that determines the loitering and thermal/slope soaring capability.

With a flexible membrane, the flight envelope was extended to lower speeds owing to the enhanced high lift capability and progressive stall of these sections. Since the animals presumably had some control over the wing camber [3,22], the envelope curve around the results with the flexible membrane best shows the full range of performance. When compared with the rigid wing section results, the low-speed flight capability is extended and combined with a softer stall, which would have enhanced control during landing manoeuvres when low speed, high drag and high lift are required.

In conclusion, these tests have quantified the two-dimensional characteristics of possible pterosaur wing
sections for the first time. They show that these creatures were significantly less aerodynamically efficient and more capable of flying at lower speeds than previously estimated. In order to achieve higher efficiency, the wing bones must be faired, which either implies a substantial weight penalty or, more likely, a high degree of pneumaticity in the fairing tissue.

Whether faired or not, the pterosaur wing sections were adapted to a low-speed flight regime that minimizes the sink rate. This regime is unsuited to marine-style dynamic soaring adopted by procellariiform birds, the sink rate. This regime is unsuited to marine-style dynamic soaring adopted by procellariiform birds, which requires high flight speed coupled with high aerodynamic soaring adopted by procellariiform birds.

The pterosaurs: from deep time
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