

## Ventral-aspect radar cross sections and polarization patterns of insects at X band and their relation to size and form

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Footnote

<sup>1</sup>Retired.

[Note on supplemental files. These two files are Excel workbooks (.xlsx, 59 and 19 kB) each with a single sheet containing a table of data and associated notes. ]

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### ABSTRACT

A dataset of ventral-aspect insect radar cross-sections (RCSs) and polarization patterns, measured at X band (9.4 GHz, linear polarization) in laboratory rigs, has been collated from a number of sources. The data have been analysed to identify relationships between RCS parameters (one representing size and two the polarization-pattern shape) and the insects' masses and morphological dimensions and forms. An improved mass-estimation relationship, with appropriate asymptotes for very small and very large insects, is presented. This relationship draws only on the RCS size parameter and it is shown that incorporating one or both of the RCS shape parameters provides little additional benefit. Small insects have polarization-pattern shapes that fall within a relatively limited region of the range of parameter values allowed by electromagnetic scattering theory. Larger insects have shapes that extend beyond this region, following a broad trajectory as size and mass increases; at masses above ~0.6 g the pattern becomes 'perpendicular', with maxima when the E-field is orthogonal to the body axis rather than parallel to it. RCS shape can be used to infer morphological form for small insects (<80 mg), but not for larger ones. These results are consistent with observations from X-band vertical-beam entomological radars and provide a basis for identification, at least to broad taxon classes, of the targets detected by such radars.

Keywords: insect; radar cross-section; polarization pattern; mass estimation; morphological form

### 1. Introduction

A variety of radar types have proved to be effective at detecting insects in flight,

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2  
3 especially when the insects have risen above terrain features and vegetation (Drake and  
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5 Reynolds 2012). Flights at these higher altitudes usually constitute migrations (Dingle  
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7 2014) and may lead to invasions of growing crops and consequent economic losses  
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9 (Drake and Gatehouse 1995; Koul, Cuperus, and Elliott 2008). Radar provides the only  
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11 practical means of acquiring information about such flights (e.g. Chapman, Reynolds, et  
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13 al. 2002; Chapman et al. 2005, 2006; Drake and Reynolds 2012, chapters 12, 13; Drake  
14  
15 and Wang 2013). In addition, radar observation has proved critical to recent research  
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17 into the cues that insects use to determine their orientation (heading direction) while  
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19 flying hundreds of metres above the surface, often in darkness, and into the effects these  
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21 orientation behaviours have on the insects' flight trajectories (Chapman et al. 2010; Hu,  
22  
23 Lim, Reynolds, et al. 2016; Reynolds et al. 2016).

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27 A radar configuration that has proved particularly useful for observing insect  
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29 migrations employs a vertical beam that incorporates both rotating linear polarization  
30  
31 and a very narrow angle conical scan (Chapman, Smith, et al. 2002; Drake 2002). With  
32  
33 this 'ZLC configuration' (also known as a 'VLR' or *vertical-looking radar*), insects are  
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35 interrogated by the beam modulations during the few seconds that it takes for them to  
36  
37 complete a beam transit. Analysis of the rather complicated echo signal time series  
38  
39 allows retrieval of information not only about the individual insect's trajectory (its  
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41 height, its speed and direction of movement, and its heading direction) but also about its  
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43 identity. The latter takes the form of four parameters characterising the target: one  
44  
45 measure of size, two of shape, and the wing-beat frequency (though for some echoes no  
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47 wing-beating is detectable). Some or all of these parameters have been used to  
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49 discriminate target types and, in association with other data (e.g. trap catches), make  
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51 identifications of the species undertaking the migrations (e.g. Chapman, Reynolds, et al.  
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53 2002; Chapman et al., 2005, 2006, 2010; Drake and Wang 2013). A study of a large  
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3 sample of echoes for which all four parameters were retrieved, obtained over eight  
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5 months from a single site in inland eastern Australia using an X-band ZLC radar,  
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7 revealed several distinct target classes which occurred with varying frequency from  
8  
9 night to night and from month to month (Drake 2016).  
10

11 From the viewpoint of a biologist user, mass is a much more readily  
12  
13 interpretable measure of target size than any directly radar-observable quantity. It is the  
14  
15 most obvious identification character to use when relating radar observations of  
16  
17 airborne insects to samples obtained from aerial or ground-level traps, in part because  
18  
19 the range of insect masses is very large (>4 orders of magnitude). Mass values also feed  
20  
21 directly into estimates of 'bioflow' and consideration of aero-ecological processes (Isard  
22  
23 and Gage 2001; Frick et al. 2013; Hu, Lim, Horvitz, et al. 2016), so their estimation has  
24  
25 intrinsic biological value. Methods for estimating the masses of insect targets from their  
26  
27 radar properties, especially the *radar cross section* (RCS, the radar measure of target  
28  
29 size), have been developed previously and are considered again here. It would  
30  
31 obviously be useful if further quantities could be identified that would allow  
32  
33 discrimination between targets with similar masses; these would constitute information  
34  
35 dimensions orthogonal to that provided by the measure of radar size. The obvious  
36  
37 candidates for such quantities are the wing-beat frequency (see below) and the two  
38  
39 shape parameters, either singly or in combination. Because the radar wavelength is  
40  
41 comparable to the size of the insect targets, the shape parameters can be expected to  
42  
43 provide information only about large-scale features. The obvious candidate is the  
44  
45 length:width ratio, termed here the 'form'  $f = l/w$ . The primary aim of this study has been  
46  
47 to identify relationships linking the three radar-target parameters to  $m$  and  $f$ , and if these  
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49 are informative to develop equations for routine estimation of these two characters from  
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51 ZLC-radar observation data.  
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3 Association of target properties with RCS size and shape values can be achieved  
4 through calculation using electromagnetics theory or by measurement. Application of  
5 theory to biological targets presents many challenges and so far has been confined to a  
6 single example (a bat; Mirkovic et al., 2016). Laboratory measurements of the RCSs of  
7 insects for which the species identification is known have been made on several  
8 occasions (summarized in Drake and Reynolds 2012, chapter 4). For this paper we have  
9 compiled a dataset containing all available vertical-incidence measurements made at X-  
10 band frequencies (~9.4 GHz) for which both size and shape parameters were obtained;  
11 we have then used this empirical data to examine the relationships between the masses  
12 and morphological properties of insects and their observable radar properties.  
13 Associations between radar parameters and taxonomic group will also be searched for.  
14 Some of the values in the dataset are from published research, some are from archived  
15 data for which only selected results had been published previously, and some are new  
16 measurements.

17  
18 The fourth identification parameter retrievable from ZLC echoes, the wing-beat  
19 frequency, may also have utility both for inferring target mass (as larger insects  
20 generally have lower wing-beat frequencies; Dudley 2000, chapter 3) and as an  
21 orthogonal dimension for discriminating between targets of similar masses (Drake  
22 2016). However, it was not determined for the specimens measured and is not  
23 considered further in this report.

## 2. Parameters, datasets, and measurements

### 2.1. Size, shape, and the polarization pattern

24 Both the ZLC observations and the laboratory measurements determine the  
25 backscattering RCS of a single insect oriented with its underside directed towards the  
26 beam ('ventral aspect'), and the variation of this with polarization angle (i.e. with the  
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3 direction of the electric field of a linear-polarized electromagnetic wave). The variation  
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5 of the RCS with polarization angle is termed the 'copolar linear polarization pattern'  
6  
7 (CLPP). In the case of the laboratory measurements, the specimen was mounted  
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9 horizontally and upright with the underside down, in the expectation that this  
10  
11 approximates a normal flight attitude, and the beam was directed vertically. The zero  
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13 angle of the CLPP is defined to be aligned with the main axis of the insect's body. For  
14  
15 the radar observations, it is not certain that the targets have adopted an upright and  
16  
17 horizontal attitude (see below), so it should only be said that the RCS values relate to  
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19 zenith incidence.  
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23 A generally steady flight is to be expected during migration, and an absence of  
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25 manoeuvring during the beam transit is assumed in the analysis of the echo-signal time  
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27 series. This assumption is essentially validated whenever a good quality fit to an echo  
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29 signal is obtained, as required for successful retrieval of the flight and target-  
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31 identification parameters (Drake and Reynolds 2012, pp. 156-159). A steady flight  
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33 suggests an upright stance (i.e. zero roll angle) but it does not require the insect's body  
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35 to be horizontal and there is now evidence that some migrating insects (particularly  
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37 small ones) fly with a non-zero (and positive, i.e. head above tail) pitch angle  
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39 (Melnikov, Istok, and Westbrook 2015). The effect of a pitched attitude cannot be  
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41 explored with the dataset available for this study and is not considered further, except to  
42  
43 note that: (1) consideration of electromagnetic scattering processes suggests pitch  
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45 effects will be slight if the pitch is only moderate, at least for targets that are not much  
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47 longer than a wavelength (32 mm for the X band radars of concern here); and (2) radar-  
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49 observed CLPPs (Dean and Drake 2005) appear compatible with the laboratory  
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51 measurements reported here.  
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3 For copolar-linear observations, the variation of the RCS  $\sigma$  with polarization  
4 angle  $\phi$  has the general form

$$\sigma(\phi) = a_0[1 + \alpha_2 \cos 2(\phi - \theta_2) + \alpha_4 \cos 4(\phi - \theta_4)], \quad (1)$$

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10 where  $a_0$  is the zenith-incidence polarization-averaged RCS and  $\alpha_2$  and  $\alpha_4$  are  
11 dimensionless parameters with positive values (Aldhous 1989; Dean and Drake 2005).  
12 The parameter  $a_0$  is a measure of the target's size, with dimensions  $L^2$  (i.e. area; units  $m^2$   
13 or, more commonly for insect targets,  $cm^2$ ; often expressed logarithmically as decibels  
14 relative to  $1 cm^2$ , denoted dBsc). In comparison with RCSs provided by other radar  
15 configurations,  $a_0$  has the considerable advantage of relating to a consistent aspect and  
16 of being independent of the insect's orientation. Relationships between  $a_0$  and mass  
17 have been developed previously (Aldhous 1989; Chapman, Smith, et al. 2002; Wang  
18 2008); an improved relationship incorporating the additional data available to this study  
19 is presented here.

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32 The expression in square brackets in Equation (1) is the CLPP. For a bilaterally  
33 symmetric target, which an insect in unbanked flight will constitute to a good  
34 approximation, the CLPP must also exhibit bilateral symmetry and therefore  $\theta_2$  and  $\theta_4$   
35 must be equal. Symmetry considerations also require that the common value, denoted  
36  $\theta_0$ , must represent either the target's body axis or the perpendicular to it. Both radar  
37 observations (e.g. Dean and Drake 2005) and laboratory measurements (Hobbs and  
38 Aldhous 2006) are generally consistent with a common value for  $\theta_2$  and  $\theta_4$ , and this  
39 simpler CLPP form has been used in the analyses presented here. The  $\alpha_2$  term in the  
40 CLPP produces elongation (of a basic circular form) and  $\alpha_4$  contributes a cruciform  
41 element. Consideration of electromagnetic scattering mechanisms indicates that for  
42 insects that are small relative to a wavelength the elongation direction of the CLPP will  
43 coincide with the body axis. Previous analyses of measurements at X band have  
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confirmed this and have also demonstrated that for very large insects (e.g. female migratory locusts *Locusta migratoria*, typical mass ~3 g) CLPP elongation occurs at right angles to the body axis (Riley 1985). Following Dean and Drake (2005), these two CLPP types will be referred to as 'parallel' and 'perpendicular' respectively. For the measurements, the insects were oriented in the rig with their heads towards  $\theta_0 = 0$ . For observations made with a radar, determining whether  $\theta_0$  represents the body axis or the normal to it requires additional, *a priori*, information (such as that the observations are being made in a region where the aerial fauna includes very few large insects). Even if the  $90^\circ$  uncertainty is resolved, ambiguity about whether the heading direction is  $\theta_0$  or  $\theta_0 + 180^\circ$  (for the parallel case) will remain.

According to electromagnetic theory, the parameters  $\alpha_2$  and  $\alpha_4$  are positive and are constrained by

$$\alpha_4 \leq \left(1 \pm \sqrt{1 - \alpha_2^2/2}\right)/2 \quad (2)$$

(Dean and Drake 2005). This equation defines a semi-elliptical boundary with  $0 \leq \alpha_2 \leq \sqrt{2}$  and  $0 \leq \alpha_4 \leq 1$ , the ellipse centre being at  $(\alpha_2, \alpha_4) = (0, 1/2)$ . In the measurement data, the angle  $\theta_0$  for targets with perpendicular CLPPs will be retrieved as  $\sim 90^\circ$ . An alternative scheme for representing perpendicular targets is to retain  $\theta_0$  as representing the body axis (i.e.  $\theta_0 = 0$ ), but to make  $\alpha_2$  negative (Dean 2007); this moves these shapes into the left-hand half of the constraint ellipse so that they are clearly separated from similarly-shaped CLPPs with parallel alignment. This reset is straightforward for the laboratory measurements, for which the true alignment is known, and is adopted in this work. It is of course problematic for observation data unless some means of distinguishing the parallel and perpendicular cases is available.

Various transformations or combinations of the variables  $a_0$ ,  $\alpha_2$ ,  $\alpha_4$  and  $\theta_0$  are available and some have interpretative value in particular circumstances (Aldhous 1989; Hobbs and Aldhous 2006; Dean 2007; Drake 2016). The 'harmonic model' (Hobbs and Aldhous 2006) of Equation (1) appears natural and straightforward and has been used in a previous investigation of insect target shapes (Dean and Drake 2005); it will form the basis for the present analysis. There will however be some discussion of the parallel  $\sigma_{xx} = \sigma(\phi - \theta_0 = 0)$  and transverse  $\sigma_{yy} = \sigma(\phi - \theta_0 = 90^\circ)$  RCS values, which can be obtained from the transformations (Aldhous 1989, p. 21)

$$\sigma_{xx} = a_0(1 + \alpha_2 + \alpha_4) \quad (3a)$$

$$\sigma_{yy} = a_0(1 - \alpha_2 + \alpha_4) . \quad (3b)$$

The ratio of these two RCSs,  $r_{sh} = \sigma_{xx}/\sigma_{yy}$ , contours of which form straight lines in the  $(\alpha_2, \alpha_4)$  plane, will receive particular attention. Use will also be made of the orthogonal variables  $(p, q)$  introduced by Drake (2016). Contours of  $p$  are concentric ellipses, with  $p = 1$  defining the constraint boundary of Equation (2) and  $p = 0$  the central point at  $(\alpha_2, \alpha_4) = (0, 0.5)$ . Contours of  $q$  are orthogonal to these and form curves that run from the centre to the periphery, with  $q = 0$  coincident with the  $\alpha_4$  axis,  $q = 1$  intersecting the boundary at  $\alpha_2 \approx 0.9$ , and  $q = 2$  intersecting at  $\alpha_2 \approx 1.4$ , just below the boundary's right-hand maximum. The  $(p, q)$  system becomes unsatisfactory for  $\alpha_4 \geq 0.5$ , but such targets occur infrequently and there is only one example in the measurement dataset.

## 2.2. The dataset and data sources

The CLPP data used in this analysis comprise 156 specimens of 40 species (Table 1). They originate from four sources: (1) measurements made in 1979 by JRR and associates at the Centre for Overseas Pest Research, U.K., denoted here by C; (2) measurements by Aldhous (1989), denoted A; (3) measurements by Wolf et al. (1993),

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3 denoted W; and (4) measurements made in 2006 by ADS and DRR at Rothamsted  
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5 Research, U.K., denoted S. The species included are mainly grasshoppers and locusts  
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7 (Orthoptera) and moths and butterflies (Lepidoptera), two groups that include numerous  
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9 migratory species (Johnson 1969); a few beetles (Coleoptera), bees and wasps  
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11 (Hymenoptera), and crane flies (Diptera) are also present. All specimens occur once only  
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13 in the dataset. There are five species for which measurements are available from two  
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15 sources and 11 species for which measurements of five or more specimens are  
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17 available; however, intra-specific variation is not considered in this paper.  
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21 The specimens derive from Europe (southern England, C, A, S) and North  
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23 America (USA, W), or from laboratory cultures of African and North American species  
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25 (C, A, and probably W). Many of the species measured are known migrants and some  
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27 have been the subject of entomological radar observations (Drake and Reynolds 2012,  
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29 chapters 10-13). A few probably non-migratory species appear to have been included  
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31 because of their ready availability when measurements were being made; these add  
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33 diversity to the sample and in some cases provide examples of particular body forms  
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35 and sizes. Most of the species represented occur only in one continent and the dataset  
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37 will therefore not be representative of the migrating insect fauna at any actual ZLC-  
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39 radar observing site. This is especially the case for the orthopteran subsample, which  
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41 comprises only six species of which four, with 33 specimens (69% of the subsample),  
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43 are of unusually large types (locusts, all with  $m > 1$  g and all but one, an outlier, with  
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45 perpendicular CLPPs). Because such large insects are not found in the migrant faunas at  
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47 the two localities (in England and Australia) where most ZLC-radar observations have  
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49 been made, some of the analyses reported here have been repeated with these specimens  
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51 excluded. The lepidopteran specimens comprise 28 species, many known to be  
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53 migratory; the mass range  $100 < m \leq 300$  mg, with 54 specimens (57% of the  
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lepidopteran subsample), is perhaps disproportionately represented. Masses (for all taxa) range from 9 mg to 4.1 g, the latter probably close to the upper limit for migrant species; the range 550-1050 mg is unfortunately represented by only a single specimen.

*Insert Table 1 near here*

Selected results from C appeared in Riley (1985), but the values used here have been recalculated from the original data and for  $\alpha_2$  and  $\alpha_4$  are mostly new, as are all the S data. For A and W, the values used here are those recalculated by Hobbs and Aldhous (2006), with  $\alpha_2$  reset negative for targets with  $\theta_0 \approx 90^\circ$  from the body axis (i.e. the largest targets); further, because these authors used a different convention, the  $\sigma_{xx}$  and  $\sigma_{yy}$  values for these large targets have been exchanged.

The values used in this analysis were obtained by the SM3 method of Hobbs and Aldhous (2006), which ensures that  $\alpha_2$  and  $\alpha_4$  fall within (or on) the constraint boundary of Equation (2). This in turn ensures that the RCS remains  $\geq 0$  at all polarization angles, as required by definition. For C and S, SM3 estimates were obtained with the *nlm* (Non-Linear Minimization) facility of the R data analysis system (R Development Core Team 2008); the quantity minimized was the sum of squares of the differences between the measured and fitted RCSs, over the full  $360^\circ$  range of polarization angles  $\phi$ . The fitted RCSs were calculated from the SM3 formula with fit variables  $\sigma_{xx}$ ,  $\varepsilon$ ,  $\beta$  and  $\theta_0$  (Hobbs and Aldhous 2006, their equation 4;  $\varepsilon^2 = \sigma_{yy}/\sigma_{xx}$  and  $\beta$  is a phase). Values of the parameters  $a_0$ ,  $\alpha_2$  and  $\alpha_4$  of Equation (1) were recovered as

$$a_0 = \sigma_{xx}(3 + 2\varepsilon \cos \beta + 3\varepsilon^2)/8 \quad (4a)$$

$$\alpha_2 = \sigma_{xx}(1 - \varepsilon^2)/2 a_0 \quad (4b)$$

$$\alpha_4 = \sigma_{xx}(1 - 2\varepsilon \cos \beta + \varepsilon^2)/8 a_0 \quad (4c)$$

(Aldhous 1989, p. 21). Note that Equation (4b) automatically delivers the negative values of  $\alpha_2$  discussed above for the targets for which  $\sigma_{yy} > \sigma_{xx}$ . Solutions were also

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3 obtained via Equation (1) for verification purposes. They were mostly within 1% of the  
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5 SM3 values for  $a_0$  and  $\alpha_2$  and often also for  $\alpha_4$ , although there were some large  
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7 differences for this last parameter, usually when its value was  $<0.1$ . These 'harmonic'  
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9 solutions also helped to identify false minima in a small number of the SM3 fits, which  
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11 were then recalculated satisfactorily using different starting values. Some statistics for  
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13 the parameter values are provided in Table 2.  
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17 *Insert Table 2 near here*  
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19 In addition to these CLPP measurements, in order to extend the analysis of the  
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21 mass vs RCS relationship to low masses, limited use has been made of available values  
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23 of  $\sigma_{xx}$  and  $\sigma_{yy}$  for smaller insects, denoted L. Measurements ( $\sigma_{xx}$  only) of planthoppers  
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25 and aphids (both Hemiptera) made by JRR and G. A. Bent respectively were recovered  
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27 by digitising a graphic in Riley (1985). Measurements of ladybirds and other beetles,  
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29 lacewings (Neuroptera), moths, and hoverflies (Diptera) were made between 2000 and  
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31 2009 by JWC and ADS, working at Rothamsted Research; statistics or example values  
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33 for some of these have appeared previously (Chapman, Reynolds, et al. 2002; Chapman  
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35 et al. 2005, 2006). After exclusion of two large specimens that fall well within the range  
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37 for which CLPP measurements are available, L comprises 77 specimens of 15 species  
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39 and covers a mass range of 0.2–65 mg; there are mass values for all of these, but only  
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41 37 (48%) also have lengths and widths.  
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### 45 46 47 **2.3. Measurements**

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49 All CLPP measurements were made in laboratory rigs with the target at a distance of  
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51  $\sim 1$  m from the antenna, which had a small aperture so that the target was in the far field.  
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53 For W a pulse-chirp radar system was used and only echo from the target's range was  
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55 recorded; the radar operated over frequencies ranging from 8 to 12 GHz and the results  
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57 are averages thought to represent a frequency of  $\sim 10$  GHz. The other measurement  
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3 systems used continuous transmission at 9.4 GHz, the frequency used by almost all X-  
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5 band entomological radars; they relied on absorptive materials, and for A a sky  
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7 background, plus cancellation of non-target echo by passively returning an anti-phase  
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9 signal via a directional coupler. Calibration was with metal spheres with RCSs  
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11 extending over the range of RCSs being measured and placed at the target position; a  
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13 calibration curve was fitted and its parameters used to convert measured echo intensities  
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15 (recorded as a voltage) to RCSs. Measurements were made at  $\phi$  intervals of  $10^\circ$  in C,  
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17  $15^\circ$  in W, and  $<1^\circ$  in A and S. The S data subset includes repeated measurements of a  
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19 small metal rod (length 16 mm, i.e.  $\sim\lambda/2$ , and diameter  $\sim 1$  mm) or 'dipole' that served to  
20  
21 verify that the system performance had not altered. Further details for A, C, and W are  
22  
23 available in the original publications (Aldhous 1989; Riley 1985; Wolf et al. 1993;  
24  
25 Hobbs and Aldhous 2006); for S, the rig used was an upgraded version of that for C,  
26  
27 with automated data acquisition at fine angular resolution.  
28  
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31

32 The smaller insects in the L data subset were measured with a transmission-line  
33  
34 (or 'rail-line') rig (Riley 1985). It was not practicable to measure the insect's angular  
35  
36 position, and the  $\sigma_{xx}$  and  $\sigma_{yy}$  values were obtained by gluing the insect first vertically  
37  
38 ( $\sigma_{yy}$  measurement) and then horizontally ( $\sigma_{xx}$  measurement) to a taut attachment line  
39  
40 passing between the rails, with the insect oriented perpendicular to the rails (and parallel  
41  
42 to the E-field) in the latter case.  
43  
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45

46 Measurements were made with specimens that were either anaesthetized (A, W),  
47  
48 freshly dead (some S, probably C), or freshly thawed after having been frozen (some S).  
49  
50 As water is the main radio-reflective component of an insect it is important to measure  
51  
52 both mass and the RCS properties before dehydration occurs; it appears that this was  
53  
54 generally achieved. The legs of some of the large locusts in C drooped when the insect  
55  
56 was mounted in the measuring rig, and for some of these the measurements were made  
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3 with the hind legs tied up with light thread to approximate the posture adopted by  
4 locusts in 'cruising flight'. An investigation of a small sample indicated that specimens  
5 with their legs drooping had CLPPs that were parallel, and thus very different from the  
6 perpendicular patterns obtained from the same specimens with their legs tied up.  
7  
8 Measurements made with the legs down, or with one or both legs missing, have not  
9  
10 been used. Masses and body lengths were measured for all 156 specimens and abdomen  
11  
12 widths for all but one.  
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### 20 **3. Results**

21  
22 The RCS parameters for each specimen, along with specimen identifications, masses,  
23 and dimensions, are provided in a supplementary spreadsheet file (Excel 2007,  
24 Microsoft Corporation, Redmond, WA, USA; available via Figshare [Note to Editor:  
25 please insert appropriate citation and link.]) containing two tables: the first with the  
26 CLPP parameters (C, A, S, and W data) and the second with  $\sigma_{xx}$  and  $\sigma_{yy}$  values (L). The  
27  
28 following sections present analyses and interpretations of this ensemble of results.  
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#### 36 **3.1. Estimation of target mass**

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38 With more measurements now available, it is appropriate to re-examine the relationship  
39 of target mass to ZLC-observable size and shape parameters. As the parameter  $a_0$  is a  
40 measure of target size, it is the obvious quantity on which to base an estimate of mass  
41 and a method depending on it alone is developed first. The possibility that improved  
42 mass estimates can be obtained by additionally incorporating the shape parameters  $\alpha_2$   
43 and  $\alpha_4$  is also explored.  
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##### 53 **3.1.1. Polarization-averaged RCS and mass**

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55 For a particular radar frequency and a particular target shape, the variation of the  
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3 target's RCS with its size passes through three regimes (Knott, Shaeffer, and Tuley  
4  
5 2004). In the 'Rayleigh region', where the target's longest dimension  $l$  is much smaller  
6  
7 than the radio wavelength  $\lambda$ , the RCS  $\sigma$  increases with the sixth power of  $l$ , i.e.  $\sigma \propto l^6$ .  
8  
9 At the opposite extreme, in the 'optical region',  $\sigma \propto l^2$ . Between these two is the  
10  
11 transitional 'Mie region', in which the RCS passes through a small number of maxima  
12  
13 and minima; these oscillations continue, with diminishing intensity, in the initial part of  
14  
15 the optical region. This variation is well documented for water spheres (e.g. Drake &  
16  
17 Reynolds 2012, pp. 51-54), for which the transition region is approximately  
18  
19  $\lambda/5 \leq l \leq 5\lambda$ . If this applies also to insects, then the two smallest specimens in the CLPP  
20  
21 dataset ( $l = 5$  mm, supplementary file) and about half of those in L just fall into the  
22  
23 Rayleigh region and the remainder ( $8 \leq l \leq 65$  mm) are Mie targets. The volume  $V$  of a  
24  
25 target will vary as  $l^3$  (again assuming a particular target shape). If migratory insects all  
26  
27 have approximately the same density, so that their mass  $m \propto V$ , then  $a_0 \propto m^2$  in the  
28  
29 Rayleigh region; for (hypothetical) optical-region mega-insects,  $a_0 \propto m^{2/3}$ .  
30  
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35 The variation of mass  $m$  with  $a_0$  for the combined datasets is shown in Figure 1  
36  
37 using logarithmic scales for both variables. For the L data points, the  $\sigma_{xx}$  values were  
38  
39 divided by a factor of 2.19, the average ratio of  $\sigma_{xx}/a_0$  for the targets in the main dataset  
40  
41 with  $a_0 < 0.25$  cm<sup>2</sup> ( $n = 18$ , s. d. 0.24). It is apparent that there is a broad spread, with a  
42  
43 range in  $\log_{10}m$  of  $\sim 0.5$  at most  $a_0$  values. For a 1-cm<sup>2</sup> target, for example, this  
44  
45 corresponds to a mass range of  $60 \leq m \leq 180$  mg. This can be attributed to variations in  
46  
47 shape and density between species, with a contribution also from experimental error.  
48  
49 There is no evidence of maxima and minima, except perhaps in the extent of the spread  
50  
51 which could originate from maxima and minima occurring at different  $a_0$  values for  
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53 insects with different morphological forms.  
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*Insert Figure 1 near here*



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3 In order to estimate the masses of the targets detected by a ZLC radar, it is  
4 necessary to develop a mathematical relationship between  $m$  and  $a_0$ . From Figure 1, it is  
5 evident that a relatively simple, smooth, curve will suffice, and indeed is all that the  
6 data warrant. At low RCS values, the curve should exhibit the asymptotic Rayleigh  $m^2$   
7 dependence (i.e.  $m \propto a_0^{1/2}$ ) and at high values it should approach  $m \propto a_0^{3/2}$ . It therefore  
8 can appropriately be represented by a spline, with two linear regions (on a log-log plot)  
9 joined by a curved form. For the latter, a simple polynomial may be sufficient. At the  
10 two transition points, the values of both functions and both first derivatives should be  
11 equal. (If they are not, discontinuities or obvious distortions will appear in distributions  
12 of mass values estimated from the relationship.) These requirements place constraints  
13 on the curve parameters, two of which, the slopes of the linear sections, are already  
14 known. The result is that if a second-order polynomial is employed to describe the link  
15 section, only two parameters plus the two transition points are free variables. Choice of  
16 transition points is not entirely free as they should conform approximately with the  
17 limits of the Rayleigh and optical regions.

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36 Chapman, Smith, et al. (2002) have previously proposed an  $m$  vs  $a_0$  relationship  
37 (Table 3), with parameters estimated from analyses of the C, A, and W data subsets, that  
38 conforms with these requirements to a good approximation. It uses a second-order curve  
39 in the Mie region, and while it does not employ an optical-region asymptote explicitly,  
40 the slope at the position of the largest target ( $\log_{10}a_0 \approx 1.2$ ) is  $\sim 1.6$ , which is not much  
41 different from the asymptotic value of 1.5, so it is broadly consistent with this  
42 requirement. To determine the free Rayleigh-region parameter, they took account also  
43 of those elements of the L data subset that were then available. For the study dataset, the  
44 transition to the Rayleigh-region was fixed at  $a_0 = 0.25 \text{ cm}^2$ , the value used by  
45 Chapman, Smith, et al. (2002), as this appears consistent with both theory and the data  
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3 (Figure 1), and the theoretical slope values for both asymptotic regions were adopted as  
4 constants. The spline parameters were then obtained by treating the  $\log_{10}a_0$  values as  
5 independent variables and minimising the square of the residual between the  $\log_{10}$ -  
6 transformed estimated and measured masses (Table 3, Figure 1). The constant,  
7 quadratic, and (for the third-order case) cubic coefficients in the linking polynomial  
8 were free and the remaining coefficients and the upper transition point determined from  
9 their values. Minimization was carried out using the Solver function in Excel 2007,  
10 using the default options; for the third-order case, it was necessary to identify the  
11 appropriate solution of a quadratic equation for the upper transition point. The  
12 minimizations converged without difficulty.

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*Insert Table 3 near here*

The new relationship with a second-order linking section differs from that of Chapman, Smith, et al. (2002) mainly by an offset in the estimated  $\log_{10}m$  of from 0.11 to 0.17. The two curves run almost parallel to each other, crossing only at  $a_0 \approx 15$  dBsc ( $m \approx 20$  g), which is larger than any of the measured values (or any likely migrant species). The effect is to increase the estimates of  $m$  by 25% for small insects and up to 39% for the largest species. The root-mean-square deviation for  $\log_{10}m$  falls from 0.23 to 0.20, which is comparable with the range of variation apparent in Figure 1 (see above) and corresponds to an uncertainty in mass of around  $\pm 50\%$ . Introducing a third order term to the polynomial reduces the r.m.s. deviation to 0.19; as the course of the curve (not shown) changes only slightly and a slight sinuosity introduced in it appears spurious, and as the upper transition point becomes unreasonably low, this more complicated formula has not been adopted.

### 3.1.2. Mass estimation incorporating shape parameters

To explore whether the shape parameters  $\alpha_2$  and  $\alpha_4$  can contribute to the estimation of mass, we have undertaken a multiple-regression analysis with  $\log_{10}m$  as the dependent variable and  $\log_{10}a_0$ ,  $\alpha_2$  and  $\alpha_4$  as independent variables. This is of course possible only for targets for which the full CLPP was measured, i.e. for the C, A, S, and W data. No attempt has been made to merge the estimation relationship into the expected asymptotic forms in the Rayleigh and optical regions. However, the dataset includes some of the largest insects known to migrate and extends down to an  $a_0$  value of  $0.005 \text{ cm}^2$  ( $-23 \text{ dBsc}$ , mass  $9 \text{ mg}$ ; Table 2), which corresponds quite closely to the lower limit of target sizes that are detectable (Chapman, Smith, et al. 2002) or commonly detected (Drake 2013, 2016) with ZLC radars, so this is not a major defect.

A sequence of multiple linear regressions was undertaken, starting with a constant term plus a term linear in  $\log_{10}a_0$  and then adding terms in  $(\log_{10}a_0)^2$ ,  $\alpha_2$ , and  $\alpha_4$ . The quadratic term was included because the  $\log_{10}m$  vs  $\log_{10}a_0$  relationship must change slope over the  $a_0$  range encompassed by the dataset (as discussed in the previous subsection where a quadratic relationship was also employed). Regressions were carried out using the *lm* (Linear Models) facility of R (R Development Core Team 2008) and assessed using analysis of variance (with R's *anova* facility). The results are summarized in Table 4. They show that a linear term in  $\log_{10}a_0$  has the greatest explanatory power, and an  $\alpha_2$  term is more effective than a quadratic term in  $\log_{10}a_0$  at further reducing the residual sum of squares and increasing the  $R^2$ . Adding either a quadratic term in  $\log_{10}a_0$  or an  $\alpha_4$  term to the combination of  $\log_{10}a_0$  (linear) and  $\alpha_2$  provides no significant further improvement. A scatterplot of  $\alpha_2$  vs  $a_0$  (not shown) reveals that  $\alpha_2 > 0.5$  for  $a_0 < 0 \text{ dBsc}$  but it decreases steadily above this point and is negative for  $a_0 > 3 \text{ dBsc}$ , so the value of this parameter provides information on the

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3 higher masses that occur above the Rayleigh region (i.e. the upturn in slope in Figure 1  
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5 at  $a_0 \approx 0$  dBsc). A linear term in  $\alpha_2$  apparently does this more effectively than a  
6  
7 quadratic term in  $\log_{10}a_0$ , with the former giving an uncertainty on the mass estimates of  
8  
9  $\pm 40\%$  (residual standard error on  $\log_{10}m$  of 0.16) and the latter  $\pm 60\%$  (0.24). The  $\alpha_4$  vs  
10  
11  $a_0$  plot (also not shown) exhibits a less clear and more scattered dependence of  $\alpha_4$  on  $a_0$ ,  
12  
13 so it is unsurprising that this parameter makes almost no additional contribution.  
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16  
17 *Insert Table 4 near here*  
18

19 Aldhous (1989), using only the A data subset, explored how mass varied with  
20  
21 several RCS quantities and found approximately linear relationships for  $\log_{10}m$  vs  
22  
23  $\log_{10}\sigma_{le}$ , where  $\sigma_{le}$  is the lesser of  $\sigma_{xx}$  and  $\sigma_{yy}$ , and for  $\log_{10}m$  vs  $\log_{10}(\sigma_{xx}/\sigma_{yy})$ . (These  
24  
25  $\sigma_{xx}$  and  $\sigma_{yy}$  are as defined here rather than in the original publication; see section 2.2.)  
26  
27 Scatterplots (not shown) demonstrate that these relationships retain their quasi-linear  
28  
29 form with the full CLPP dataset. We have therefore explored including these terms in  
30  
31 the regression as an alternative to  $\alpha_2$  (Table 4). It was found that  $\log_{10}\sigma_{le}$  provides only  
32  
33 modest additional explanatory power but  $\log_{10}(\sigma_{xx}/\sigma_{yy})$  performs almost as well as  $\alpha_2$ .  
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### 38 **3.2. Shape as an identification character**

#### 39 **3.2.1. Radar shape and taxonomic group**

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41 The distribution of the shape parameter pairs ( $\alpha_2$ ,  $\alpha_4$ ) over the allowed region of the  
42  
43 ( $\alpha_2$ ,  $\alpha_4$ ) plane is shown for the full CLPP dataset in Figure 2(a). It can be seen that  
44  
45  $\alpha_4 < 0.5$  for all but one specimen and that pairs are found across much of the right-hand  
46  
47 half ( $\alpha_2 > 0$ ) of the  $\alpha_4 < 0.5$  part of the allowed area but are confined to a more  
48  
49 restricted region ( $\alpha_4 < 0.3$ ) of the left-hand half. There is a concentration of targets on or  
50  
51 close to the boundary where  $\alpha_2 \approx 1.0$ , a region corresponding approximately to the  
52  
53 'main cluster' identified in analyses of ZLC-radar observations of target shape (Dean  
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3 and Drake 2005, Drake 2016). There are relatively few targets in the region  $0 < \alpha_2 \leq 0.5$   
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5 and only one of these has  $\alpha_4 < 0.2$ ; the equivalent region of the left-hand half, however,  
6  
7 is crowded.  
8

9  
10 *Insert Figure 2 near here*  
11

12 Lepidopteran specimens extend throughout the right-hand half of the figure but  
13  
14 are most numerous in or close to the main-cluster region; there is only one specimen  
15  
16 with  $\alpha_2 < 0$ . Orthopterans occur in both halves but only two fall into the main cluster;  
17  
18 orthopterans in the right-hand half mostly have  $\alpha_4 > 0.2$ . The specimens from other  
19  
20 orders all fall into the right-hand half, some in the main cluster but others well away  
21  
22 from the boundary; this is a morphologically mixed group so a variety of  $(\alpha_2, \alpha_4)$   
23  
24 combinations is to be expected. The figure also shows the results for the dipole; they  
25  
26 fall along the boundary where  $\alpha_2$  is approaching its maximum possible value and,  
27  
28 reassuringly, show little spread.  
29  
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31

32 In order to explore these patterns more precisely, a main-cluster region (MCR)  
33  
34 will be defined in terms of the orthogonal variables  $(p, q)$  as these delineate it very  
35  
36 nicely (Figure 2(b)). There are 79 specimens in the MCR, of which 2 are Orthoptera, 68  
37  
38 are Lepidoptera, and 9 are other taxa; of the 77 specimens falling outside the MCR, 46  
39  
40 are Orthoptera, 26 Lepidoptera, and 5 other. The uneven distribution of orthopterans  
41  
42 and lepidopterans is significant (G-test for a model-I  $2 \times 2$  contingency table with Yates  
43  
44 correction,  $G = 69.3$ , 1 d.f.,  $P < 0.001$ ; Sokal and Rolf 1995, ch. 17); it remains  
45  
46 significant, though at a lower level, if the specimens with perpendicular CLPPs (32  
47  
48 locusts, 1 moth) are excluded ( $G = 21.8$ ,  $P < 0.001$ ).  
49  
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### 52 53 54 3.2.2. *Relation of radar shape to mass and morphology* 55

56 The location of a specimen on the  $(\alpha_2, \alpha_4)$  plane will be determined by its size and  
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3 general morphology, with these represented by the parameters  $m$ ,  $l$ ,  $w$ , and  $f$ . A  
4  
5 relatively clear pattern emerges when the symbols for the points on the  $(\alpha_2, \alpha_4)$  plane  
6  
7 indicate the specimen's mass (Figure 2(b)). All specimens in the left-hand half of the  
8  
9 allowed region have  $m > 600$  mg and only one specimen (an outlier – see below) in the  
10  
11 right-hand half exceeds this mass. All but two of the lightest specimens ( $m \leq 100$  mg,  
12  
13  $n = 29$ ) occur in the MCR, as do 49 (73%) of the intermediate-mass insects  
14  
15 ( $100 < m \leq 300$  mg,  $n = 67$ ). The remaining population of the MCR comprises 3 insects  
16  
17 with  $300 < m \leq 400$  mg, which constitutes 5% of the heaviest specimens ( $m > 300$  mg,  
18  
19  $n = 60$ ). The mass difference between specimens in the MCR and those falling outside it  
20  
21 is significant (Wilcoxon rank sum test, two-sided,  $n_1 = 79$ ,  $n_2 = 77$ ,  $W = 333$ ,  $P$   
22  
23  $< 0.001$ ). Inspection of Figure 2(b) suggests that in the MCR and further inwards, in the  
24  
25 adjoining 'MCR Extension' (MCRE; Figure 2(b)), there is a general increase of target  
26  
27 mass with distance from the boundary. Further, for specimens outside the MCR, there  
28  
29 appears to be a trend of mass increasing from top-right ( $\alpha_2 \approx 1.4$ ,  $\alpha_4 \approx 0.5$ ) to bottom-  
30  
31 mid-left ( $\alpha_2 \approx -0.7$ ,  $\alpha_4 \approx 0.05$ ). The latter phenomenon is the same as that noted in  
32  
33 section 3.1 where  $\alpha_2$  (if signed as here) was found to be a useful predictor of mass. As is  
34  
35 to be expected from the analysis of section 3.1, there is also a significant difference in  
36  
37  $a_0$  values between the specimens in the MCR and outside it ( $W = 705$ ,  $P < 0.001$ ). The  
38  
39 pattern of  $a_0$  values across the  $(\alpha_2, \alpha_4)$  plane (not shown) is similar to that for mass,  
40  
41 though the variance appears greater and this makes the trends somewhat less evident.  
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49 Regression analyses (Table 5) show that the trends are significant. However, the  
50  
51 decrease of mass with increasing  $p$  within the combined MCR and MCRE regions has a  
52  
53 modest adjusted  $r^2$  and accounts for only 29% of the sum of squares. Four points with  
54  
55 very low mass ( $9 \leq m \leq 14$  mg) appear as outliers, but this is essentially an artefact of  
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57 the measurement sample in which low masses are under-represented: it appears likely  
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3 that inclusion of more insects with  $m < 40$  mg would further reduce the  $r^2$  and the sum  
4  
5 of squares accounted for. Adding a  $q$  term into the regression produces negligible  
6  
7 improvement. For the non-MCR specimens, the adjusted  $r^2$  indicates a strong  
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9 dependence of mass on  $\alpha_2$ , with this parameter accounting for 86% of the sum of  
10  
11 squares; two outlier points here are due to specimens that arguably belong in the MCR  
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13 as they have masses (49, 82 mg) lower than the rest ( $\geq 173$  mg) and  $q$  values that only  
14  
15 just fall outside the nominal MCR limits. An added  $\alpha_4$  term is marginally significant but  
16  
17 accounts for only a further 1% of the sum of squares. When  $\log_{10}a_0$  rather than  $\log_{10}m$  is  
18  
19 made the dependent variable, the results (Table 5) are similar, though the adjusted  $r^2$   
20  
21 values are lower and smaller proportions of the sum of squares are accounted for. These  
22  
23 weaker statistics reflect the less strong trends and greater variance discerned through  
24  
25 inspection of the plots (previous paragraph). The trend statistics for the non-MCR  
26  
27 specimens are obviously strengthened by the large number of  $m > 1$  g specimens (all  
28  
29 locusts) with their extreme values of both  $\alpha_2$  and the response variables  $m$  and  $a_0$ .  
30  
31 Excluding these leaves a still significant relationship for  $\log_{10}m$  but not for  $\log_{10}a_0$   
32  
33 (Table 5). If the two low-mass specimens that arguably belong in the MCR are also  
34  
35 excluded, the adjusted  $r^2$  for  $\log_{10}m$  rises to 0.77 (though that for  $\log_{10}a_0$  is not  
36  
37 improved). Similar weak trends for  $\log_{10}a_0$  in these two regions were noted in an  
38  
39 analysis of a single night of ZLC-radar observations at an Australian site (Dean and  
40  
41 Drake 2005).  
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48 *Insert Table 5 near here*

49  
50 Differences between the MCR and non-MCR specimens are also evident when  
51  
52 the specimen character indicated is the length, width, or form (Figure 3; detailed results  
53  
54 not presented), and these are statistically significant (Wilcoxon test,  $n_2 = 77$  for  $l$  but 76  
55  
56 for  $w$  and  $f$ ,  $P < 0.001$ ). Regressions of  $l$ ,  $w$ , and  $f$  against  $p$  for the specimens in the  
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3 combined MCR and MCRE account respectively for 16, 5, and 1% of the sum of  
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5 squares, with the  $p$  coefficient significant at the  $P < 0.001$  level for  $l$ , at  $< 0.05$  for  $w$ , and  
6  
7 not significant ( $\sim 0.3$ ) for  $f$ . For the non-MCR specimens regressed against  $\alpha_2$ , the  
8  
9 proportions are 81, 10, and 37% respectively with  $P$  values  $< 0.001$ , 0.007, and  $< 0.001$ ;  
10  
11 with the  $m > 1$  g locusts excluded, the proportions are 36, 30, and 9% with  $P$  values  
12  
13  $< 0.001$ ,  $< 0.001$ , and 0.05. Thus the length shows differences and trends similar to those  
14  
15 for  $m$  and  $a_0$ , but for the width and form the trends are less clear.

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17  
18 *Insert Figure 3 near here*  
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20  
21 Two additional regression analyses have been undertaken for the form  $f$  to  
22  
23 examine its potential for providing information orthogonal to that from  $a_0$ . For the  
24  
25 CLPP data, adding  $q$  into the regression of  $f$  against  $p$  in the combined MCR and MCRE  
26  
27 increases the sum of squares accounted for, but only to 2% and neither parameter is  
28  
29 significant. Adding  $\alpha_4$  into the regression of  $f$  against  $\alpha_2$  for all non-MCR specimens  
30  
31 provides negligible improvement; however, with the locusts excluded the proportion  
32  
33 accounted for increases to 20% and this is also achieved with  $\alpha_4$  alone.

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35  
36 For the small specimens in the L subsample, the available radar measure of  
37  
38 shape is the ratio  $r_{sh}$  (i.e.  $\sigma_{xx}/\sigma_{yy}$ ). The form  $f$  shows a significant relationship with  $r_{sh}$  for  
39  
40 these specimens (Figure 4(a);  $n = 39$ , coefficient  $0.224 \pm 0.028$ , intercept  $1.28 \pm 0.34$ ,  
41  
42 64% of variance accounted for,  $P < 0.001$ ). Two clusters apparent in this plot are due to  
43  
44 single-species groups, but other species show a broad spread of  $r_{sh}$  values (Figure 4(a)).  
45  
46 A similar analysis for the mostly larger CLPP specimens, with  $r_{sh}$  calculated from  $\alpha_2$   
47  
48 and  $\alpha_4$  via Equations 3 (and after excluding two specimens with  $r_{sh} > 50$ ), shows no  
49  
50 relationship (Figure 4(b);  $n = 121$ ,  $P \approx 0.8$ ). This is still the case if only specimens with  
51  
52  $m \leq 80$  mg, i.e. in the same mass range as those in the L subsample, are included  
53  
54 ( $n = 15$ ,  $P \approx 0.6$ ), and also if the range is extended slightly to  $m \leq 100$  mg in order to  
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enlarge the sample ( $n = 27$ ,  $P \approx 0.6$ ). Thus for targets with  $a_0 \leq 0.25 \text{ cm}^2$  (which from Table 3 corresponds to  $m \leq 80 \text{ mg}$ ),  $f$  can be estimated as

$$f = 0.224r_{\text{sh}} + 1.28 . \quad (5)$$

Because of the uncertainties on these parameters, the 1-standard deviation uncertainty on  $f$  will increase from  $\pm 0.3$  to  $\pm 0.9$  over the range  $0 < r_{\text{sh}} \leq 20$ . The data provide no basis for estimating  $f$  for targets with  $a_0 > 0.25 \text{ cm}^2$ .

*Insert Figure 4 near here*

### 3.2.3. Mass and morphology associated with different radar shapes

An alternative way of representing these relationships is to determine where insects of different sizes or shapes are located in the  $(\alpha_2, \alpha_4)$  plane. In the case of mass (Figure 5), insects with  $m \leq 150 \text{ mg}$  are found entirely in the MCR; those with  $150 < m \leq 250 \text{ mg}$  occupy both the MCR and the MCRE and are also found in the adjacent area to the right ( $q > 1.5$ ) of these; those with  $250 < m \leq 400 \text{ mg}$  are predominantly in the MCRE; those with  $400 < m \leq 1000 \text{ mg}$  are in the area adjacent and to the left of the MCRE (with  $0 < q \leq 0.8$ ), and almost all of those with  $m > 1 \text{ g}$  fall into the left-hand half of the diagram (where  $q$  is considered negative), with the largest ( $m > 2 \text{ g}$ ) slightly further to the left. All but three of the 13 lightest insects ( $m \leq 70 \text{ mg}$ ) occur along the outer periphery of the MCR, with  $p > 0.95$ . A trajectory traced through the plane by a notional 'typical insect' as its mass increases can thus be identified, though it is broad in extent (Figure 6). (It should be noted that variance in mass occurs along the trajectory, on the scale indicated by the distance across the ellipses, as well as laterally.) The mass ranges associated with the regions of Figure 6, and the number of specimens falling within or near them, are presented in Table 6. Region C, which contains only a small number of specimens, is tentatively identified as a spur off the main trajectory.

1  
2  
3 *Insert Figure 5 near here*  
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5 *Insert Figure 6 near here*  
6

7 Similar trends are evident for RCS  $a_0$  and length  $l$  (Table 6), although these  
8 characters partition the specimens into the regions less precisely than does  $m$ . As  
9 specimens with similar widths  $w$  occur in both the left- and right-hand sides of the plane  
10 (not shown), there is no equivalent trend for this character. For the form  $f$ , specimens  
11 with  $f \leq 6$  ( $n = 102$ ) fall within regions A, B, D and E while those with  $f > 6$  fall  
12 predominantly ( $n = 31$ ) into the left-hand half of the plane with the remainder ( $n = 22$ )  
13 scattered widely across the right hand half. The specimens in region C and close to the  
14 constraint boundary ( $n = 5$ ) had narrow bodies ( $2 < w \leq 3$  mm) and lengths in the range  
15  $20 < l < 22$  mm (giving  $6 < f < 8$ ); all were *Chorthippus brunneus*, a grasshopper. Their  
16 location was the closest to that of the metal dipoles (Figure 2), which are also very  
17 narrow ( $\sim 1$  mm) and made to a length, 16 mm, that will produce resonance when  
18 aligned with the E field. The RCS values of the five *C. brunneus* specimens were  $\sim 1$  dB  
19 higher than predicted (from relationship of Table 5) for their  $\sim 200$  mg mass, so it is  
20 possible that they were at or near resonance too. Two smaller specimens of this species,  
21 with  $l \approx 16$  mm and  $f \approx 8$ , fell into region B. Several of the small L specimens have  
22  $r_{sh} \geq 10$  (and  $f \geq 2.6$ , Figure 4), and from this it can be inferred that they lie in the right-  
23 hand parts of the A, B, or C regions (Figure 6, which shows contours of  $f$ ). The ladybird  
24 beetles, with  $f \approx 1.3$ , have  $r_{sh} \leq 4.3$  and therefore must fall to the left of A and B; their  
25  $\alpha_4$  values are unknown but given their low masses ( $11 \leq m \leq 32$  mg), it seems likely  
26 (see Figure 5) that they lie close to the constraint boundary.  
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52 *Insert Table 6 near here*  
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### 3.2.4. Identification of perpendicular CLPPs

As noted above, when targets are observed with a ZLC-radar (as opposed to in a laboratory rig), there is a possibility of a 90° error in the retrieved alignment. The question of whether this can be resolved from the radar observations themselves is examined here by direct reference to the dataset. In this sample, and after eliminating one outlier point, the transition to perpendicular CLPPs occurs around  $m \approx 600$  mg (Figure 2b) and in the zone  $3 < a_0 \leq 4$  dBsc. Unfortunately the dataset includes only one specimen in the  $500 < m < 1000$  mg transition region, so the transition behaviour cannot be discerned in any detail.

## 4. Discussion

The measure of size provided by a ZLC radar is the polarization-averaged RCS  $a_0$  and an established monotonic relationship between this and mass has been verified, and its parameters refined, in this work. Masses estimated from  $a_0$  have an uncertainty of  $\sim \pm 60\%$ , but given the wide range of insect sizes this still provides useful discrimination. If the shape parameter  $\alpha_2$  is incorporated into the mass-estimation procedure the uncertainty can be reduced to  $\sim \pm 40\%$ , but seamless linking into the lower and upper asymptotic regions will then be lost so this is feasible only when small or very large insects are absent or of no interest. Use of a spline quadratic in  $\log_{10} a_0$  to link appropriate linear asymptotic slopes (Table 3, line 2) appears adequate and robust and leaves the shape parameters as independent identification variables; this relationship is therefore recommended for general use.

An important finding from this research is that the values of  $\alpha_2$  and  $\alpha_4$ , the primary shape parameters, are determined mainly by the size of the insect, as represented either by its mass or its length. Small and medium-sized insects, with

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2  
3  $m \leq 150$  mg and  $l \leq \sim 20$  mm, are almost completely confined to regions A and B of the  
4  
5  $(\alpha_2, \alpha_4)$  plane (Figure 6), an area also known as the MCR and identified previously  
6  
7 from ZLC observations of nocturnal insect migration. As insect size increases beyond  
8  
9 150 mg the CLPP parameters follow a broad trajectory, moving initially away from the  
10  
11 constraint boundary (i.e.  $\alpha_4$  increasing) into region D and then leftwards ( $\alpha_2$   
12  
13 decreasing) into E, passing into the left-hand half of the plane ( $\alpha_2$  negative, CLPP  
14  
15 perpendicular) at a mass of around 600 mg (region F) and getting close to the lower left-  
16  
17 hand boundary ( $\alpha_2$  and  $\alpha_4$  both decreasing, region E) at  $\sim 4$  g. Narrow ( $w \leq 3$  mm)  
18  
19 insects with  $m \approx 200$  mg and  $l \approx 20$  mm may be found off this main trajectory in region  
20  
21 C.  
22  
23  
24

25  
26 These major variations of  $\alpha_2$  and  $\alpha_4$  with  $m$ , and hence also with  $a_0$ , mean that  
27  
28 these parameters do not form a naturally orthogonal system with  $a_0$ . The trajectory is  
29  
30 broad, and the obvious candidate for orthogonality is the transverse direction. This  
31  
32 changes as  $m$  increases, meaning that different relationships would have to be sought in  
33  
34 different regions. Symmetry considerations require that a perfectly round insect will  
35  
36 have a circular CLPP, i.e. one with  $\alpha_2 = \alpha_4 = 0$ , and basic consideration of  
37  
38 electromagnetic scattering leads to the expectation that the CLPPs of elongated insects  
39  
40 are unlikely to be circular, and that the patterns for insects of the same mass but  
41  
42 differing degrees of elongation (i.e. different forms) will not be the same, so some  
43  
44 degree of orthogonality seems assured. The obvious relation to test, therefore, is one  
45  
46 between form and a quantity representing the transverse direction; the latter could be  $\alpha_2$   
47  
48 or  $q$  in regions A, B and D and  $\alpha_4$  in regions C-G. Regression analyses found no  
49  
50 contribution from  $q$  in the former region and only a moderate one from  $\alpha_4$  in the latter.  
51  
52  
53  
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55  
56 In contrast to these results which suggest there is little prospect of estimating  $f$   
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58 from the radar data for the larger insects ( $m > \sim 80$  mg) predominating the CLPP sample,  
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1  
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3 Chapman, Reynolds, et al. (2002) and Chapman et al. (2005, 2006) were able to use  
4  
5 shape to discriminate between targets when observing very small ( $m \leq 13$  mg) species.  
6  
7 They used the ratio  $r_{sh} = \sigma_{xx}/\sigma_{yy}$  as their identification parameter, with values  
8  
9 determined by laboratory measurements (and included here in L) of around 10, 10, and  
10  
11 20 for diamondback moths, carabid beetles, and lacewings respectively. In this study, a  
12  
13 linear relationship between  $r_{sh}$  and  $f$  was found for the full L sample ( $m \leq 81$  mg), but  
14  
15 not for a smaller sample of the specimens of similar size in the CLPP sample. That a  
16  
17 relationship between radar shape and morphological form should be more apparent for  
18  
19 small insects (targets falling into the Rayleigh region, or close to it) than for larger ones  
20  
21 (Mie region) is not altogether unexpected given the more complicated radio-wave  
22  
23 scattering processes involved with the latter which lead, at higher masses, to the  
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25 transition to perpendicular CLPPs. Given that  $r_{sh}$  broadly increases with  $\alpha_2$  (Figure 6),  
26  
27 there is no reason to expect it to have any advantage as a discriminating variable for  
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29 specimens larger than those in L.  
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35 The majority of targets detected so far with ZLC-radars have  $a_0 < 3$  dBsc (e.g.  
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37 Drake 2016), which the results presented here indicate is in the range where CLPPs are  
38  
39 parallel rather than perpendicular. Thus target alignments can confidently be attributed  
40  
41 to the direction of the maximum axis of the CLPP. An attempt to detect this transition in  
42  
43 a single night's observations at an Australian site was unsuccessful, even though the  
44  
45 radar seemed to indicate that a small number of sufficiently large targets were present  
46  
47 (Dean and Drake 2005). Operation of a unit in a region where large locusts, or other  
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49 species of similar size, are known to be present in the aerial fauna may be necessary to  
50  
51 verify that orientations of such targets are at right angles to the observed CLPP axis, and  
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53 to better determine the  $a_0$  value at which the transition occurs.  
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3 The measurement sample analyzed in this work includes many migrant species,  
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5 but it is not representative of the migrant fauna at any particular location. However, a  
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7 study of radar characters of insects detected at the Australian site over an eight-month  
8  
9 period identified peaks in the A, B, D and E regions of the  $(\alpha_2, \alpha_4)$  plane, but not in C  
10  
11 or the region to the left of A and B (Drake 2016). Moreover, the peaks found by that  
12  
13 study in regions A and B (the MCR) were almost all centred at  $a_0 \leq -5$  dBsc, with  
14  
15 some extending up to 0 dBsc (masses of ~90 and ~200 mg respectively, Figure 1),  
16  
17 which agrees reasonably with the  $m \leq 150$  mg range determined here (Table 6). Peaks in  
18  
19 the D and E regions in the Australian study almost all had  $0 < a_0 \leq 5$  dBsc (i.e.  
20  
21  $200 < m < 850$  mg), which again is consistent with this work. The Australian data  
22  
23 suggest that the possibly resonant behaviour of the *C. brunneus* specimens in the  
24  
25 measurement dataset may be relatively unusual. They also indicate that the  
26  
27 measurement sample, with the large locusts ( $m > 1$  g) excluded, is reasonably  
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29 representative of the Australian migrant fauna (even though the great majority of the  
30  
31 species measured do not occur in Australia). Perhaps migrant insect faunas do not differ  
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33 morphologically very much from one continent to another: while the species differ, at  
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35 higher taxonomic levels there is much commonality (e.g. Johnson 1969) and as the  
36  
37 phenomena of nocturnal migration are broadly similar from region to region (e.g. Drake  
38  
39 & Gatehouse 1996, Drake & Reynolds 2012) it can be expected that the adaptations  
40  
41 leading to them will be too. The results from this study should therefore have wide  
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43 applicability.  
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Table 1. Composition and sources of the CLPP dataset<sup>a</sup>.

| Taxonomic group     | Source |        |       |        |         |
|---------------------|--------|--------|-------|--------|---------|
|                     | C      | A      | W     | S      | All     |
| Orthoptera          | 5, 30  | 2, 18  | 0, 0  | 0, 0   | 6, 48   |
| Lepidoptera         | 1, 13  | 15, 34 | 3, 5  | 13, 42 | 28, 94  |
| others <sup>b</sup> | 1, 1   | 1, 2   | 3, 9  | 1, 2   | 6, 14   |
| All                 | 7, 44  | 18, 54 | 6, 14 | 14, 44 | 40, 156 |

<sup>a</sup>Number of species, number of specimens. See text (section 2.2) for details of sources C, A, W and S.

<sup>b</sup>Bees and wasps (Hymenoptera) (2, 7), beetles (Coleoptera) (2, 3), crane flies (Diptera) (2, 4).

Table 2. Summary of masses and RCS parameter values for the CLPP dataset.

| Taxonomic group     | Mass               | RCS parameter         |                        |                      |
|---------------------|--------------------|-----------------------|------------------------|----------------------|
|                     | $m$<br>(mg)        | $a_0$<br>(dBsc)       | $\alpha_2$             | $\alpha_4$           |
| Orthoptera          | 1597<br>(68, 4120) | 4.6<br>(0.39, 16.8)   | -0.03<br>(-0.90, 1.31) | 0.20<br>(0.01, 0.56) |
| Lepidoptera         | 194<br>(9, 648)    | 0.98<br>(0.013, 2.72) | 0.91<br>(-0.10, 1.22)  | 0.23<br>(0.08, 0.43) |
| others <sup>b</sup> | 114<br>(10, 220)   | 0.51<br>(0.005, 1.48) | 0.94<br>(0.62, 1.27)   | 0.22<br>(0.09, 0.48) |
| All taxa            | 618<br>(9, 4120)   | 2.04<br>(0.005, 16.8) | 0.62<br>(-0.90, 1.31)  | 0.22<br>(0.01, 0.56) |

<sup>a</sup>Average (minimum, maximum).

<sup>b</sup>As in Table 1.

Table 3. Parameters of mass vs RCS relationships<sup>a</sup>

| Origin<br>( $\log_{10}m$ r.m.s.)    | Region                  |                                     |               |  |
|-------------------------------------|-------------------------|-------------------------------------|---------------|--|
|                                     | Lower <sup>b</sup>      | Linking                             | Upper         | Upper transition <sup>c</sup>                |
| Chapman et al. (2002)<br>(0.23)     | 2.097<br>0.5            | 2.205<br>0.8729<br>0.3323           | –             | –  |
| This study, second order.<br>(0.20) | 2.207<br>0.5<br>(81 mg) | 2.331<br>0.9124<br>0.3425           | 2.079<br>1.5  | 0.8580<br>(7.21 cm <sup>2</sup><br>2.32 g)   |
| This study, third order.<br>(0.19)  | 2.243<br>0.5<br>(88 mg) | 2.249<br>0.8028<br>0.9583<br>0.7826 | 2.1459<br>1.5 | 0.2605<br>(1.87 cm <sup>2</sup> ,<br>359 mg) |

<sup>a</sup>Parameters are for the relationship of  $\log_{10}m$  to  $\log_{10}a_0$ . (Note  $a_0$  (dBsc) is  $10\log_{10}a_0$ .)

The first value in each list is the constant term, the second the coefficient of  $\log_{10}a_0$ , and so on for higher powers of  $\log_{10}a_0$  if these are present.

<sup>b</sup>Transition from the lower to the linking curve was fixed at  $\log_{10}a_0 = -0.60206$  (i.e.  $a_0 = 0.25$  cm<sup>2</sup>) for all curves. Value in parenthesis is the corresponding mass  $m$ .

<sup>c</sup>First value is  $\log_{10}a_0$ ; values in parenthesis are  $a_0$  and the corresponding mass  $m$ .

Table 4. Results of multiple linear regression analyses of  $m$  vs ( $a_0$ ,  $\alpha_2$ ,  $\alpha_4$ ) relationships

| Model <sup>a</sup><br>(note)         | Coefficients | ANOVA sum<br>of squares, ( $P$ ),<br>[residual]. | Adjusted $R^2$ ,<br>residual<br>standard error. | Degrees of<br>freedom,<br>$F$ , ( $P$ ). |
|--------------------------------------|--------------|--|---|--|
| Constant                             | 2.418        | –  | 0.759   | 1, 154                                   |
| $\log_{10}a_0$                       | 0.829        | 35.93 (<0.001)<br>[11.29]                        | 0.271   | 490 (<0.001)                             |
| Constant                             | 2.345        | –  | 0.813   | 2, 153                                   |
| $\log_{10}a_0$                       | 0.9378       | 35.93 (<0.001)                                   | 0.239   | 338 (<0.001)                             |
| $(\log_{10}a_0)^2$                   | 0.2156       | 2.578 (<0.001)<br>[8.709]                        |   |  |
| Constant                             | 2.712        | –  | 0.915   | 2, 153                                   |
| $\log_{10}a_0$                       | 0.5068       | 35.93 (<0.001)                                   | 0.161   | 837 (<0.001)                             |
| $\alpha_2$                           | –0.4655      | 7.334 (<0.001)<br>[3.953]                        |   |  |
| (See note b)                         |              |  |   |  |
| Constant                             | 2.645        | –  | 0.819   | 2, 153                                   |
| $\log_{10}a_0$                       | 0.2591       | 35.93 (<0.001)                                   | 0.235   | 351 (<0.001)                             |
| $\log_{10}\sigma_e$                  | 0.4343       | 2.831 (<0.001)<br>[8.457]                        |   |  |
| Constant                             | 2.705        | –  | 0.908   | 2, 153                                   |
| $\log_{10}a_0$                       | 0.5106       | 35.93 (<0.001)                                   | 0.167   | 768 (<0.001)                             |
| $\log_{10}(\sigma_{xx}/\sigma_{yy})$ | –0.4315      | 7.009 (<0.001)<br>[4.278]                        |   |  |

<sup>a</sup>Dependent variable is  $\log_{10}m$ .

<sup>b</sup>Adding  $(\log_{10}a_0)^2$  reduces sum of squares by 0.07 (0.2%), coefficient not significant ( $P \sim 0.09$ ). Adding  $\alpha_4$  reduces sum of squares by 0.04 (0.1%), coefficient not significant ( $P \sim 0.2$ ).

Table 5. Results of regression analyses of  $m$  and  $a_0$  vs  $(p, q)$  and  $(\alpha_2, \alpha_4)$ .

| Region and dependent variable <sup>a</sup>                | Model [2nd model] <sup>b</sup>                      | Coefficients  | Sum of squares, ( $P$ ) <sup>c</sup> , [residual]. | Adjusted $r^2$ , residual standard error. | Degrees of freedom, $F$ , ( $P$ ). |
|---|---|---------------|--|---|------------------------------------|
| MCR and MCRE<br>$\log_{10}m$                              | Constant<br>$p$<br>[ $q$ , 0.9, 0.02%]              | 3.35<br>-1.39 | -<br>2.88 (<0.001)<br>[6.95]                       | 0.29<br>0.27                              | 1, 98<br>41 (<0.001)               |
| MCR and MCRE<br>$\log_{10}a_0$                            | Constant<br>$p$<br>[ $q$ , 0.2, 1%]                 | 1.07<br>-1.50 | -<br>3.36 (<0.001)<br>[21.34]                      | 0.13<br>0.47                              | 1, 98<br>15 (<0.001)               |
| All except MCR<br>$\log_{10}m$                            | Constant<br>$\alpha_2$<br>[ $\alpha_4$ , 0.02, 1%]  | 2.97<br>-0.66 | -<br>14.57 (<0.001)<br>[2.39]                      | 0.86<br>0.18                              | 1, 75<br>456 (<0.001)              |
| All except MCR<br>$\log_{10}a_0$                          | Constant<br>$\alpha_2$<br>[ $\alpha_4$ , 0.2, 0.9%] | 0.47<br>-0.46 | -<br>7.02 (<0.001)<br>[4.41]                       | 0.61<br>0.24                              | 1, 75<br>120 (<0.001)              |
| All except MCR<br>( $m > 1$ g excluded)<br>$\log_{10}m$   | Constant<br>$\alpha_2$<br>[ $\alpha_4$ , 0.003, 9%] | 2.74<br>-0.41 | -<br>0.99 (<0.001)<br>[0.89]                       | 0.52<br>0.15                              | 1, 42<br>47 (<0.001)               |
| All except MCR<br>( $m > 1$ g excluded)<br>$\log_{10}a_0$ | Constant<br>$\alpha_2$<br>[ $\alpha_4$ , 0.6, 0.6%] | 0.22<br>-0.18 | -<br>0.19 (0.07)<br>[2.22]                         | 0.06<br>0.23                              | 1, 42<br>3.6 (0.07)                |

<sup>a</sup>MCR, MCRE – Main Cluster Region, MCR Extension (see sections 3.2.1, 3.2.2).

<sup>b</sup>The second model incorporates an additional independent variable; this variable, its  $P$  value, and the additional reduction in the sum of squares it produces (expressed as a proportion), are provided.

<sup>c</sup>From ANOVA; sum of squares accounted for by the variable, and  $P$  value for that variable.

Table 6. Predominant regions of the  $(\alpha_2, \alpha_4)$  plane for different insect characters

| Mass $m$ (mg)        |                     |                     | RCS $a_0$ (dBsc) |        |               | Length $l$ (mm) |        |         |
|----------------------|---------------------|---------------------|------------------|--------|---------------|-----------------|--------|---------|
| Range                | Number <sup>a</sup> | Region <sup>b</sup> | Range            | Number | Region        | Range           | Number | Region  |
| <70                  | 13                  | A                   |                  |        |               | <12             | 6      | A       |
| 70-150               | 46                  | A, B                | <-2 <sup>c</sup> | 45     | A, B          |                 |        |         |
| 150-250 <sup>x</sup> | 32                  | B, C                | -2-0             | 25     | B, D          | 12-22           | 87     | A, B, C |
| 250-400              | 22                  | D                   | 0-4 <sup>d</sup> | 50     | B, C,<br>D, E | 22-25           | 17     | D       |
| 400-1000             | 10                  | E                   |                  |        |               | 25-30           | 12     | E       |
| 1000-<br>2000        | 15                  | F                   | >4               | 35     | F, G          | >30             | 33     | F, G    |
| >2000                | 17                  | G                   |                  |        |               |                 |        |         |

<sup>a</sup>Total 155 as specimen with no width measurement excluded from analysis.

<sup>b</sup>See Figure 6, but note that the ellipses there are schematic and indicate a general area.



Figure captions

Figure 1. Relation of measured insect masses  $m$  (mg) to polarization-averaged RCSs  $a_0$  (dBsc) for the study dataset. Key: • – CLPP averages; + – estimated from  $\sigma_{xx}$ ; × – water-filled capillary tube, estimated from  $\sigma_{xx}$ ; main curve – fitted spline (Table 3), with linear sections dashed and the second-order section continuous; short-dashed lines – Chapman, Smith, et al. (2002) fit, of which for clarity only the ends are shown.

Figure 2. (a) Positions of the CLPP dataset specimens on the  $(\alpha_2, \alpha_4)$  plane;  $n = 156$ . The solid line indicates the constraint boundary (Equation (2)). The area within the dashed rectangle is shown enlarged at right. Key: ○ Orthoptera ( $n = 48$ ), + Lepidoptera ( $n = 94$ ), Δ other orders ( $n = 14$ ), □ dipole ( $n = 7$ ). (b) As (a) but with the dipoles omitted and symbols representing the specimen's mass (see key on plot) and the MCR (Main Cluster Region,  $0.8 < p \leq 1$ ,  $0.7 < q \leq 1.5$ ) and MCRE (MCR Extension,  $0.5 < p \leq 0.8$ , same  $q$  range) indicated.

Figure 3. As Figure 2b but with size of symbol indicating the specimen's (a) length, (b) width, and (c) form. For (b) and (c),  $n = 155$ .

Figure 4. Scatterplots of the form  $f$  against the ratio  $r_{sh} = \sigma_{xx}/\sigma_{yy}$  for (a) the L subsample (with regression line shown) and (b) the combined C, A, W, and S subsamples (less two specimens with  $r_{sh} > 50$ ). Clusters A and B in (a) are of ladybirds and lacewings respectively; regions C, D, and E are occupied by carabid beetles, hoverflies, and moths respectively. Specimens with  $m \leq 80$  mg in (b) are marked with a +.

Figure 5. Positions on the  $(\alpha_2, \alpha_4)$  plane of CLPP dataset specimens in different mass ranges. The symbols indicate the specimens' forms  $f$  (see key at top left).

Figure 6. Schematic trajectory on the  $(\alpha_2, \alpha_4)$  plane traced by a notional 'typical insect' as its mass increases (grey line). The ellipses show the approximate area of the plane occupied by CLPP specimens for each of the mass ranges of Figure 5 (with the two similar distributions for  $70 < m < 100$  mg and  $100 < m < 150$  mg combined, and see Table 6); they provide an indication of the lateral spread of points from the notional trajectory line. The dashed lines are contours of  $r_{sh} = \sigma_{xx}/\sigma_{yy}$  with values 2, 5, 10, and 20.

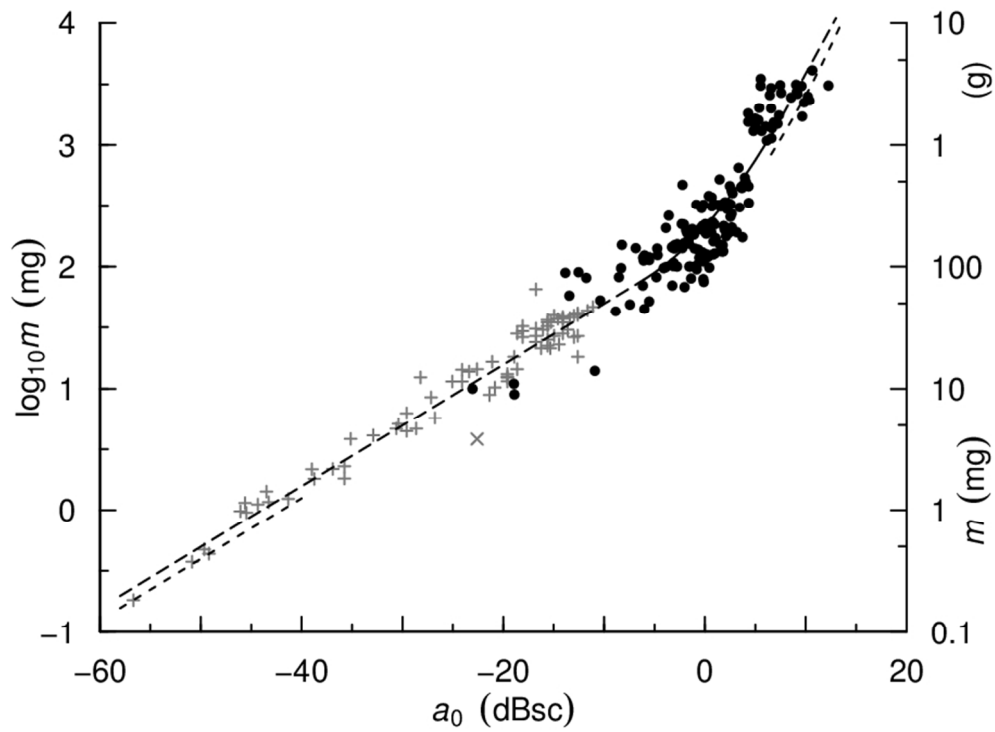


Figure 1  
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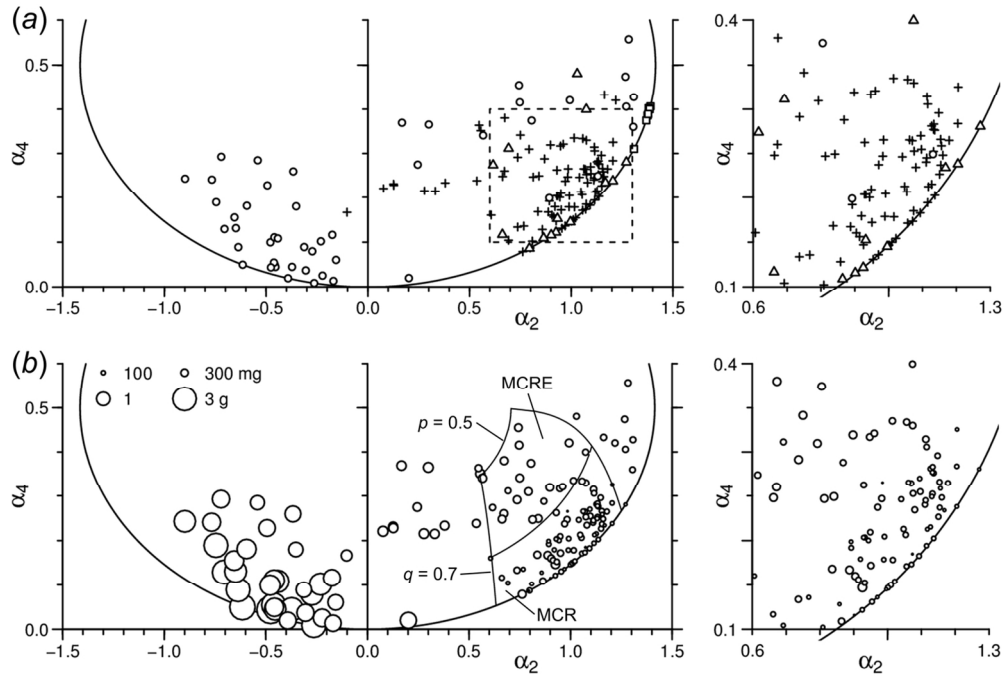


Figure 2  
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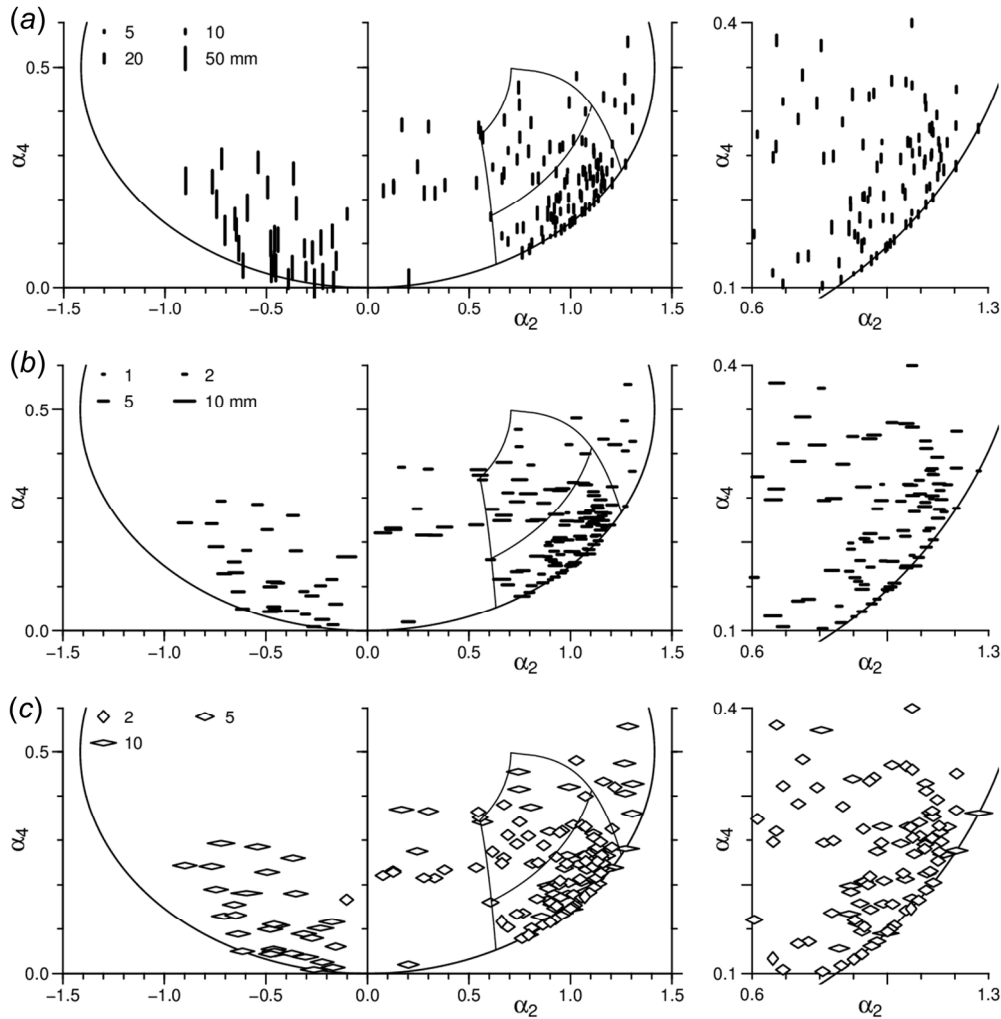


Figure 3  
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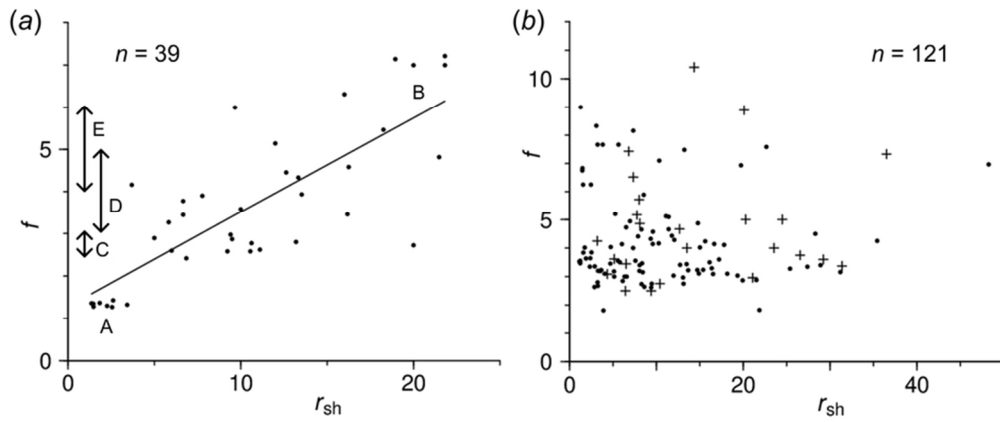


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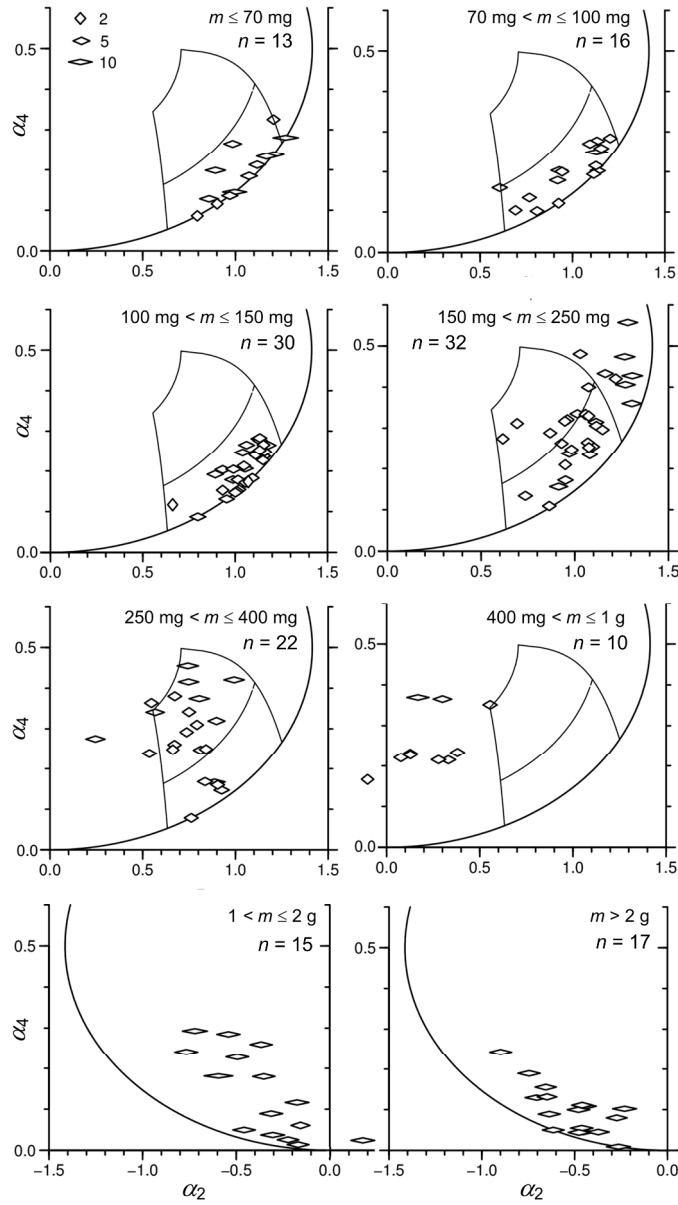


Figure 5  
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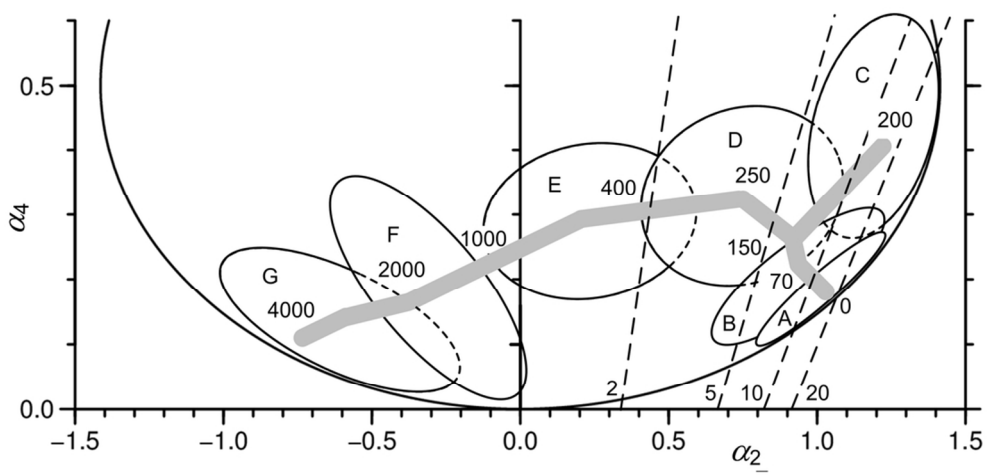


Figure 6  
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