Elytral surface structure in *Poecilus lepidus* (Coleoptera: Carabidae): What about the nature of its inheritance?

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**Key words.** Coleoptera, Carabidae, matt and bright elytral surface, colour impression, epigenetics, sex-limited inheritance, geographical differences

**Abstract.** The cuticle of the ground beetle *Poecilus lepidus* (Leske) (Coleoptera: Carabidae) displays a complex variety of colour morphs that are genetically determined. Besides the colour of the elytra, there is a remarkable intersexual dimorphic difference in its surface structure between males and females, as there is in most *Poecilus* species. At least in Central European populations of *P. lepidus*, all males exhibit a bright elytral surface, whereas all females exhibit a matt surface; this difference is due to a nearly plain surface in males but a knobbly sculptured one in females. At first glance, this phenomenon may be easily interpreted as being a result of secondary sexual character inheritance. However, *P. lepidus* specimens are known from two other regions that differ markedly: females display the same bright elytral surfaces as males in the Italian Apennine Mountain and all over Bulgaria. In the present study, cross-breeding experiments with German and Bulgarian specimens resulted in a dominance of matt over bright in the classical Mendelian F₃ : 1 ratio, although only in females. All the males showed a bright elytral surface. These findings throw into question the interpretation as given above. Because classical modes of inheritance do not provide a sound interpretation of the data as here detailed, we discuss two forms of epigenetic mechanisms which might be responsible for the observed sexual dimorphism: (1) silencing of a gene by genomic imprinting, and (2) haplo-insufficient dominance of one allele to its counterpart. Ultimately, the observed pattern is interpreted by sex-limited inheritance that depends on the presence or absence of a particular allele.

**INTRODUCTION**

The impression of a structural colour as perceived when observing an insect cuticle depends on both the multiple transparent layers within the cuticle as well as the structure of the cuticular surface. Like other insects, many beetles display bright metallic colours due to the presence of multilayer systems. This term covers two modes of colour-producing layers: (i) due to a stack of alternating layers of different refractive indices (and thickness) with a wide range in the numbers of such double layers (quarter wavelength plates); and (ii) a helicoidal order of cholesteric liquid crystals within the layer in which the orientation of chitin micelles rotates (half wave plates). Such helicoidal systems exhibit optical activity; they are nearly restricted to scarab beetles (Lamellicornia) and were first described in detail by Neville & Caveney (1969). Parker (2005) in his review *Simple classical optics in animals – Reflectors and antireflectors* only refers to the former type as “multilayer systems”.

Multilayer colours may be superimposed by effects of structures on the cuticular surface. They can occur due to different forms of scattering at surface structures of the cuticle with a dimension of ~ 1 µm (Hinton, 1976; antireflectors in Heteroptera: Miyamoto & Kosaku, 2002 and Lepidoptera: Yoshida, 2005), or by depressions of the surface of diameter ~ 10 µm that lead to different angles of reflected light at different parts of the depression (e.g. dimples in Cicindelids: Seago et al., 2008). The more or less planar surface of the hexagonal depression or dimple in *Cicindela* produces the main contribution to the colour effect that is superimposed by the reflectance of the dimple’s walls (Seago et al., 2008).

Studies on the heredity of colour morphs of carabid beetles are scarce (Puisségur, 1964; Liebherr, 1983). The complex variety of structural colour morphs of the ground beetle *Poecilus lepidus* (Leske) (Coleoptera: Carabidae) is genetically determined (Mossakowski et al., 2008; Paarmann et al., 2008). In the present study, we focus our interest on the remarkable difference in the appearance of the cuticular surface between males and females, which is well known for most *Poecilus* species (Fig. 1).

Certainly in Central European populations of *P. lepidus*, all males exhibit a bright elytral surface, all females a matt one, due to the virtually smooth surface in the males and micro-sculptured, knobbly one in females. At first glance, this phenomenon appears to result from secondary sexual character inheritance. However, specimens of *P. lepidus* from two other geographic regions differ from this pattern, females display the same bright elytral surfaces as males in the Italian Apennine Mountains (P. Brandmayr, pers. comm.) and throughout Bulgaria (B. Gueorguiev, pers. comm.). Crossbreeding experiments with German and Bulgarian specimens resulted in a dominance of matt over bright elytra in the classical Mendelian F₃ : 1 ratio, but only in females. All the males showed a bright elytral surface. These findings throw doubt on the interpretation given above, i.e. that the effect is a secondary sexual character.
Because classical modes of inheritance do not give a sound interpretation of the data without additional assumptions, we discuss two forms of epigenetic mechanisms that might be responsible for the elytral patterns observed between males and females of this particular beetle species: (1) silencing of a gene by genomic imprinting; and (2) haplo-insufficient dominance of one allele over its counterpart. Lastly, we interpret the observed pattern by sex-limited inheritance that works when an allele (allele 1) of a gene S is present but does not work with another allele at the same locus (allele 2).

This paper focuses on three questions: (1) What is the difference in surface structure between bright and matt elytral cuticles? (2) How are these characters inherited? and (3) Is there any adaptive value apparent for these differences?

**MATERIAL AND METHODS**

**Material**
Specimens derived from Germany (Lueneburg Heather, leg. J. Eggers, T. Assmann) and Bulgaria (Batchevo, north of Razlog, Pirin-Macedonia, leg. H. Turin).

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Fig. 1. Elytral surface of *Poecilus lepidus*. Bright colour impression of a male, and matt of a female from a German population. Elytron breadth = 2.4 mm.

Fig. 2. Scanning electron micrographs of *Poecilus lepidus* elytral cuticle, dorsal view. Parental (A, C) and F2 (B, D) representative specimens; females display both forms in F2: knobbly structured matt (B) and smooth (D) cuticle.
Methods

Scanning electron microscopy (SEM) micrographs

These were produced using a ZEISS Auriga FIB/SEM (Focused Ion Beam & Scanning microscope) at the Institute for Micro Sensors, Actuators and Systems (IMSAS), Department of Physics and Electrical Engineering, University of Bremen, Germany. Probes were not sputtered.

Crossbreeding experiments

*Poecilus lepidus* has one generation per year. Initially, beetles were reared for many years in order to study the life cycle of this species and later on to analyse the genetics of colour morph production. Some male specimens from Bulgaria were used in the crossing experiments performed in the present study.

P generation

Ten violet virgin females of the “Lüneburger Heide” strain (mat elytra) were kept without males until they reached sexual maturity. At this point, five green and sexually mature Bulgarian males were transferred to the females for a few days each (for further details of breeding conditions and crossbreeding experiments, see Paarmann et al., 2008).

F1 generation

Here, beetles were separated according to their colour in order to extend our former colour experiments with the same material, i.e. violet (n = 27), green (n = 25) and red (n = 17) morphs were placed into three containers. Their offspring resulted in the F1 morph numbers as shown in Table 4. Matt and bright cuticle status of females and males was recorded for each container and each generation, respectively.

Reflectance spectra were measured with a diode array spectrometer (MSC 621 II with CLH 600, ZEISS Optical Sensor Systems, Jena, Germany). The diameter of the measuring spot was about one millimetre across. Measurements were taken every four nanometres. PTFE (polytetrafluoroethylene) was used as a white standard. The reflectance curves were calculated and managed using the computer program Aspect Plus (ZEISS Optical Sensor Systems, Jena).

Photographs were taken using a Canon EOS 550D mounted on a LEITZ Photo-Macroscope. The probes were illuminated using a LED ring light (Mueller Optronik MK-S64T). Multiple pictures were combined with the programme Helicon Focus.

RESULTS

Cuticle surface structure of elytra

The surface structure of the elytra in *P. lepidus* is virtually smooth in males and sculptured in females. Under the assumption of Mendelian inheritance, character states should split in the F2 generation into a different number of morphs. Therefore, we tested different F2 offspring under the scanning electron microscope for intermediate states. Two morphs resulted: a smooth and a micro-sculptured one, conforming to the matt and bright forms (Fig. 2). The cell width in both males and females was 10–15 µm.

![Fig. 3. Scanning electron micrographs of *Poecilus lepidus* elytral cuticle, 64° view to the surface at the breaking edge of the scissor cut. Female (A) and male cuticle (B). A part of the multiple layers producing the structural colour are resolved in the cutting section above the arrows (B).](image)

The impression of the microsculpture depends on the viewing angle (normal view: convex, picture turned by 180°: concave); therefore, the knobbly structure was proved by looking on the probes breaking edge at an angle of 64° (Fig. 3).

Crossbreeding experiments

The first crossbreeding experiment was performed with German females and Bulgarian males. All the males displayed bright elytra, all the females matt ones in the F1 generation. In the F2, a split occurred, but only in females; the matt form was dominant over the bright form in a ratio near a 3 : 1 ratio. All the males showed a bright elytral surface (Table 1). More details and an interpretation are given in Table 4.

<table>
<thead>
<tr>
<th>P: GER matt × BUL bright</th>
<th>P: GER matt × BUL bright</th>
<th>P: BUL bright × BUL bright</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1: matt bright</td>
<td>F1: bright</td>
<td>F1: bright</td>
</tr>
<tr>
<td>F2: matt &amp; bright</td>
<td>F2: bright</td>
<td>F2: bright</td>
</tr>
</tbody>
</table>

Reflection measurements

The reflectance of the two structural morphs differs distinctly. The maximum reflectance in the male was 28.9%
pressions as well as that differing in wavelength due to the mixture of the colour of the plain bottom area of the de-

Schultz & rankin, 1985). Thus, the colour reflectance were also found in many other species (e.g. Schultz, 1983; Geo-

drical ridges on an undulating surface, additionally cov-

ered by larger tubercles (Stegmann, 1930). Such structures are due to a multilayer system (Mossakowski, 1979). These
center of about 8–10 μm (Mossakowski, 1979). These
ticidae). The cuticle surface of matt C. campestris tiger beetles of the genus Cicindela.

2008) and display a bright reflection only if the cuticle sur-

face structure of elytra (Fig. 2).

The colours displayed by the cuticular surface of P. lepi-
dus are due to a multilayer system (Mossakowski et al., 2008) and display a bright reflection only if the cuticle sur-

face is more or less smooth (Fig. 2). This condition was

found in all male specimens examined, regardless of their geographical and genetic origin. Two morphs are present in the females: a bright and a matt one. The latter differs by a sculptured surface structure of the elytra (Fig. 2).

C. campestris L. is covered by depressions with a di-

ameter of about 8–10 μm (Mossakowski, 1979). These
grooves occur on an undulating surface, additionally cov-

ered by larger tubercles (Stegmann, 1930). Such structures were also found in many other species (e.g. Schultz, 1983; Schultz & Rankin, 1985). Thus, the colour reflectance is a mixture of the colour of the plain bottom area of the depres-

sions as well as that differing in wavelength due to the
different angles of the walls (Seago et al., 2008). These authors referred to such depressions as “dimples”.

The cuticle surface of matt P. lepidus is sculptured but the effect of the uneven structure of these humps causes the same effect as the dimples in Cicindela spp.

DISCUSSION

Cuticle surface structure of elytra

The colours displayed by the cuticular surface of P. lepi-
dus are due to a multilayer system (Mossakowski et al., 2008) and display a bright reflection only if the cuticle surface is more or less smooth (Fig. 2). This condition was found in all male specimens examined, regardless of their geographical and genetic origin. Two morphs are present in the females: a bright and a matt one. The latter differs by a sculptured surface structure of the elytra (Fig. 2).

In beetles, a similar but inverse situation was found in tiger beetles of the genus Cicindela. The cuticle surface of C. campestris L. is covered by depressions with a diameter of about 8–10 μm (Mossakowski, 1979). These grooves occur on an undulating surface, additionally covered by larger tubercles (Stegmann, 1930). Such structures were also found in many other species (e.g. Schultz, 1983; Schultz & Rankin, 1985). Thus, the colour reflectance is a mixture of the colour of the plain bottom area of the depressions as well as that differing in wavelength due to the

Geographic differences. See discussion.

Parent-of-origin

Genomic imprinting

Haplo-insufficient dominance

Sex-limited

Table 2. Modes of inheritance tested to interpret the crossbreeding results for Poecilus lepidus.

<table>
<thead>
<tr>
<th>Model</th>
<th>Counter-argument</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mendelian</td>
<td>No uniformity in F₁; split in F₂ depends on gender.</td>
</tr>
<tr>
<td>Sex chromosome linked</td>
<td>Males do not display different phenotypes; 3 : 1 in F₂ females.</td>
</tr>
<tr>
<td>Attached X chromosome</td>
<td>Needs only 2 sex-linked phenotypes, because the other two are not viable. Our breeding success is &gt;90%.</td>
</tr>
<tr>
<td>Maternal</td>
<td>Does not conform because of 3 : 1 split in F₂.</td>
</tr>
<tr>
<td>Parent-of-origin</td>
<td>Does not conform because of 3 : 1 split in F₂.</td>
</tr>
<tr>
<td>Genomic imprinting</td>
<td>Needs additional assumptions. See discussion.</td>
</tr>
<tr>
<td>Haplo-insufficient dominance</td>
<td>Needs additional assumptions. See discussion.</td>
</tr>
<tr>
<td>Sex-limited</td>
<td>Geographic differences. See discussion.</td>
</tr>
</tbody>
</table>

F₁, F₂ refer to crossbreeding German × Bulgarian Poecilus lepidus. *Based on females with two X chromosomes fused at the centromere and usually a Y chromosome. The phenotypic effect of an allele depends on whether it is inherited from the mother or the father. 604
tions and always produce the matt form of elytra. If the first gene is autosomal, only one of the two alleles at the locus will be silenced, whilst the other should produce matt males. Such males have so far not been found to occur in nature nor also in our experiments. This problem may be resolved if it is supposed that the gene governing the elytral traits is on the sex chromosome. D. patens has an XX/XO sex determining chromosomal system (Nettmann, 1986; Serrano & Galian, 1998). Hence, the offspring should show a ratio of 1 : 1 in females in the F1 generation (matt : bright), instead of our experimental result of 3 : 1. As a consequence, we cannot present a sound interpretation based solely on genomic imprinting.

**Haplo-insufficient dominance**

This form of incomplete dominance exists if the expression of one copy of a gene at a particular locus is insufficient to produce the coded protein. It could work under the assumption that the two alleles produce a different amount of units in terms of their gene products and a threshold must be overcome to induce the next step in protein production.

When we specify that allele S1 produces 12 and S2 produces 8 units in terms of their gene products, and that females will increase their temperature more quickly than matt females. In consequence, it may be hypothesised that females will increase their temperature more quickly and to a higher value than males under identical solar radiation. Because the difference is lower than 10% (Table 5), the temperature effect on the elytra will be small.

Several effects of bright versus matt cuticular surface may play a role in adaptation, such as camouflage, temperature modulation, or adhesion during copula.

**Sex-limited inheritance**

Sex-limited inheritance fits the data when the differences in females from northern and southern populations is interpreted as being due to a dimorphism that is under the control of an autosomal gene. If a trait has a sex-limited expression this phenomenon will normally occur within a population. The consequence is that selection can only affect the sex with the sex-limited trait; in the other sex the gene is hidden to selection. Such a scenario may well have significant impacts on selection (Reinhold, 1999). In contrast, the matt and the bright females of *P. lepidus* occur in different geographical regions. We have a sex-limited trait in northern populations (bright males; matt females), uniformity in southern populations, and a dimorphism in females only in our crossbreeding experiments between German and Bulgarian specimens (Table 4). All attempts to interpret the data lead to an autosomal gene. Such a conclusion is concordant with the statement that autosomal genes are involved in traits limited to the homogametic sex, in the case of *P. lepidus*, the female sex (Reinhold, 1999).

**Adaptation**

Several effects of bright versus matt cuticular surface may play a role in adaptation, such as camouflage, temperature modulation, or adhesion during copula. The reflectance measurements show the expected differences. Bright males of the same colour reflect more light than matt females. In consequence, it may be hypothesised that females will increase their temperature more quickly and to a higher value than males under identical solar radiation. Because the difference is lower than 10% (Table 5), the temperature effect on the elytra will be small.

**Table 3.** Interpretation of crossbreeding German × Bulgarian *Poecilus lepidus* under the assumption of haplo-insufficient dominance inheritance. A sound explanation is only possible when the gene is assumed to be on an autosome and one of the alleles in males, S1 or S2, is suppressed or inherited sex-limited (bold-marked allele). Numbers in the right column indicate protein units encoded by the gene (see Discussion).

<table>
<thead>
<tr>
<th>Female</th>
<th>Male</th>
<th>F1</th>
<th>F2</th>
</tr>
</thead>
<tbody>
<tr>
<td>♀ S1S1</td>
<td>♂ S2S2</td>
<td>S1S1 matt</td>
<td>S1S1, 2 S1S2 matt (50)</td>
</tr>
<tr>
<td>F2: ♀ S1S1 matt</td>
<td>♂ S2S2</td>
<td>S1S1 bright (40)</td>
<td></td>
</tr>
<tr>
<td>F2: ♀ S1S1 matt</td>
<td>♂ S2S2</td>
<td>S1S1, 2 S1S2 bright (50)</td>
<td></td>
</tr>
<tr>
<td>S2S2 bright (12)</td>
<td>S2S2 bright (Σ 83)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Table 4.** Interpretation of crossbreeding experiments with German (GER) and Bulgarian (BUL) *Poecilus lepidus* compared with the assumed situation in the natural populations. Numbers of offspring are indicated in brackets. In F2, matt is dominant to bright; phenotype matt is suppressed in males.

<table>
<thead>
<tr>
<th>GER ♀ × GER ♂</th>
<th>GER ♀ × BUL ♂</th>
<th>BUL ♀ × BUL ♂</th>
</tr>
</thead>
<tbody>
<tr>
<td>P: S1S1 matt × S1S1 bright</td>
<td>P: S1S1 matt × S2S2 bright</td>
<td>P: S2S2 bright × S1S1 bright</td>
</tr>
<tr>
<td>F1: S1S1 matt × S1S1 bright</td>
<td>F1: S1S2 bright (30)</td>
<td>F1: S2S2 bright × S1S1 bright</td>
</tr>
<tr>
<td>F1: S1S1, 2 S1S2 matt</td>
<td>S1S2 bright (40)</td>
<td>F1: S2S2 bright × S1S1 bright</td>
</tr>
<tr>
<td>S2S2 bright (12)</td>
<td>S2S2 bright (Σ 83)</td>
<td>F1: S2S2 bright × S1S1 bright</td>
</tr>
</tbody>
</table>
this morph, the southernmost edge of the species range on the Balkan Peninsula.

In general, the females of the genus Poecilus show matt elytral surfaces, but P. lepidus is the only species with such a geographical variability in this particular feature. Schultz & Hadley (1987) failed to find any significant temperature differences between isolated elytra of specimens of the same Cicindela species with either black or coloured elytra (compare also Hadley et al., 1992). Only specimens with white cuticles displayed a difference. All these cincindelids have a surface structure comparable to the dimples described above and their inner structure is the same, a multilayer system. But white in Cicindela depends on a quite different internal structure (Schultz, 1983). In P. lepidus, the surface structures differ between smooth in bright and sculptured in matt cuticles. If the matt structure of the cuticle has a mimetic effect or has an effect on copulation, the question arises “Why are such clear geographic differences found?”

CONCLUSION

Structure

We found two structural morphs for the elytral surface, which produce the different impressions of brightness to the human eye.

Inheritance

We cannot interpret the data on the basis of genomic imprinting alone, because every new approach produced new obstacles. An interpretation by insufficient haplo-dominance need two assumptions: the hypothesised gene must be on an autosome and one of the alleles must be silenced in males. The gene cannot be located on the X chromosome because under this assumption, the F2 offspring ratio in females should be 1:1.

An interpretation by sex-limited inheritance fits the situation in northern populations (bright males, matt females). The uniform southern most populations must have different alleles; therefore, the observed pattern may be interpreted by sex-limited inheritance that depends on the presence or absence of a particular allele.

Adaptation

The measurements gave a remarkable difference between matt and bright cuticle reflectance. As a consequence of this difference, a matt cuticle of the same colour under identical radiation input may warm up to a higher temperature, due to structural differences of the cuticle surface, which may in turn have adaptive survival advantages to insects living in cooler climes.


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