Giving eyespots a shiner: Pharmacologic manipulation of the Io moth wing pattern [version 2; referees: 2 approved]

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Abstract

Our knowledge of wing pattern formation in Lepidoptera has advanced significantly in recent years due to the careful examination of several groups of butterflies. The eyespot is a prominent feature of Lepidoptera wing pattern, especially in the family Saturniidae. The present study examined how sulfated polysaccharides affected the wing pattern formation of the Io moth, *Automeris io* (Saturniidae). Prepupae and pupae of this species were subjected to injections of heparin and cold shock. While the cold shock had little to no effect on wing pattern, the aberrations resulting from heparin injections were moderate to profound and depended on the dose and the stage at which injection was made. The changes consisted of expansion of the black ring around the dorsal hindwing eyespots and distortion of discal spots on both dorsal and ventral sides of forewings, suggesting a possible link between genetic controls of these elements. Several different types of scales form the normal color pattern of *Automeris io*, and heparin-induced changes correspond to changes in shape of scales. The resulting aberrations are dubbed ‘Black Eye’ and ‘Comet Eye.’ Other known aberrations of *Automeris io* eyespots are summarized, illustrated, and named.

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Author roles: Sourakov A: Conceptualization, Data Curation, Formal Analysis, Investigation, Methodology, Resources, Visualization, Writing – Original Draft Preparation, Writing – Review & Editing

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Amendments from Version 1

Following the reviews of Version 1, I have made significant changes to this paper, which are listed below in order of importance:

1. Additional specimens: The sample size of the individuals transformed by heparin injections increased from two to seven due to the emergence of one additional specimen and dissections of dead pupae.
2. Pupal staging: I described in as great detail as possible how the timing of injections was evaluated using time-lapse photography.
3. Injection dose: I described precisely how heparin solution was produced for the injections.
4. Eyespot structure: I examined the composition of the eyespots, which are formed by several different types of scales. I also examined transformed and aberrant individuals and showed that changes in wing pattern involved not only changes in color, but also changes in the types of scales involved.
5. Discussion: I discussed the observed results in light of current knowledge about the effect of heparin on gene function. However, this part of the manuscript remains very concise to preserve the Research Note nature of this publication. I made use of reviewers’ comments by referring the readers to them, as they appear side-by-side with the manuscript and contain many thoughtful comments and important references.
6. As a result of the above changes, the number of figures has doubled, including figures submitted as supplementary material, showing time-lapse photos of pupal development and microscopic study of wing scales involved in the formation of the eyespots.

See referee reports

Introduction

While our understanding of the mechanisms involved in butterfly wing pattern development has been increasing exponentially in the recent two decades, the work has been largely limited to butterflies such as Junonia, Heliconius, Papilio and Bicyclus. Thanks to these ‘model’ genera, we now understand homologies among wing pattern elements and the adaptive radiation that led to the kaleidoscope of intriguing ‘designs’ found among ca. 160,000 Lepidoptera species (Martin & Reed, 2010).

Natural and artificially generated aberrations serve as windows into the developmental mechanisms and evolutionary history of animals. In addition to many naturally occurring melanic aberrations and some melanic recessive phenotypes that can be obtained and/or maintained through inbreeding, the dark markings of Lepidoptera wings can sometimes be amplified by the timely application of a colder regime to the immature stages (e.g., Sourakov, 2015 and references within). Serfas & Carroll (2005) first demonstrated that injections of heparin into the early pupal stage can simulate cold shock and alter wing patterns in similar ways. Martin & Reed (2014) utilized heparin injections to understand genetic controls and homologies among separate wing pattern elements.

Eyespots are characteristic of many Lepidoptera, and considerable advances have been made towards understanding their evolution (Monteiro et al., 2006). In Automeris io, a species whose name, if anglicized (‘Eye’‘Oh!’), invokes associations with its pair of magnificent dorsal hindwing eyespots that are exposed when the moth (otherwise cryptic) is threatened. Several recessive mutations causing deformations of the black ring surrounding the dark blue eyespot with a white center have been obtained through inbreeding, conducted first by Thomas Manley (1978, 1990) and, more recently, myself (Table 1 below). However, the most dramatic aberration, which involves the melanization of almost the entire hindwing, was found in an A. io male collected in 1966. It was noticed only recently while the MGCL Saturniidae collection was being re-curated (Cowell, 2012).

A previous study by Sourakov (2015), in which the development of the Bella moth was manipulated by temperature change, suggested that the black wing pattern elements of the hindwings may be forming during the prepupal stage. Hence, in the present study, heparin injections were done to both prepupal and pupal stages, and the cold shock was delivered to the former. The results of these injections, while not replicas of known genetic aberrations, are quite dramatic. They are illustrated along with the slight aberrations possibly resulting from cold-shock and heritable aberrations, both those heretofore described and those previously unrecorded.

Structure of Automeris io eyespots (See Supplementary File for figures)

Close examination of both wings in normal individuals reveals not only color but also structural differences between different color pattern elements, supporting similar findings in Nymphalid butterflies (Iwata & Otaki, 2016). As pointed out by Martin (2017) and Marcus (2017) and is confirmed here (Supplementary Figure S1) the white center of the small ventral forewing eyespot (DI element) is located on the discal cell’s crossvein between M₁ and M₃. Similarly, the dorsal hindwing eyespot center is also tied to the discal cell’s crossvein between M₁ and M₃ (Supplementary Figure S2). Sibling moths can frequently be recognized by the size and shape of the eyespot, the width of each of its elements, and its position on the wing. As the examination of several specimens with differently positioned eyespots suggests, the position of the eyespot on the wing, which in some broods is shifted distally towards the outer black band of the wing, is determined by the wing venation (Supplementary Figure S2 & Supplementary Figure S3).

Microscopic examination makes it obvious that the white and blue center of the dorsal hindwing eyespot is inset deeper within the wing plane compared to the eyespot’s black ring, which is level with the surrounding yellow areas. Removing successive layers of scales using Scotch tape revealed that the ground layer of all three of the above elements is formed by similarly shaped, wide, short cover scales (Supplementary Figure S3). In that first layer of scales covering the wing membrane, the white (fluorescent in UV light), flat, short, and wide scales, aggregate mostly along the discal cell’s crossvein that runs through the middle of the eyespot. The center of the elongated white spot that these scales form most likely
Table 1. Aberrations of dorsal hindwing eyespots found in Automeris io.

| Aberration name | Description | Author, details |
|-----------------|-------------|----------------|
| “Black eye”     | Figure 1A and Figure 2D – expansion of black eyespot ring, so that the area between eyespot and outer black band entirely or almost entirely black | Heparin-induced, present study |
| “Broken eye”    | Figures 3B,C – vertical streaks of black medially of the eyespot | Manley, heritable aberration |
| “Teardrop”      | Figure 3D – eyespot shape modified, with an appendix extending towards wing base | Manley, heritable aberration |
| “Caecus”        | Figure 3A and perhaps Figure 4C – eyespot completely disappears, masked by black pigment | Wild, collected by J.L. Boughner; Heparin-induced, present study |
| “Comet eye”     | Figure 1C, Figure 2A – black ring around eyespot with smudges extending towards wing base | Heparin-induced, present study |
| “Barley eye”    | Figure 1E and Figure 3F – black ring uneven, bulging or protruding locally | Cold shock-induced, present study; Obtained through inbreeding by Sourakov |
| “Winking eye”   | Figure 3E – blue circles forming eyespots are of uneven size on left and right wings due to uneven expansion of black ring | Obtained through inbreeding by Sourakov |

corresponds to the developmental focal point, akin to that of Junonia coenia, as described by Nijhout (1980).

The dark-blue similarly shaped, iridescent scales intermingling with additional white scales described above form the blue-and-white area of the eyespot. Their non-iridescent-black and translucent-yellow counterparts also underlie the black ring around the eyespot and the surrounding yellow area, respectively (Supplementary Figure S3). In the black ring, however, these short and flat ground scales are hidden by another layer of long and flat scales that are ca. 1.5 times longer than the first type and half as wide. These types of scales are absent in the blue-and-white eyespot center (Supplementary Figure S4). The color of the blue eyespot area can vary depending on the proportion of white scales that is intermixed with the blue ones.

While the scale type that dominates the surface layer of the black ring is also numerous in the surrounding yellow areas, it is not readily visible as it is covered by thin, long, bristle-like scales that are twice as long and only a quarter as wide. As a result, the surface of hindwing outside of the eyespot appears hairy rather than scaly. In the black ring, the bristle-like scales (colored black) are also present, but they are few compared to the rest of the hindwing.

Methods

Representatives of five broods of Automeris io from local stock (over 300 caterpillars) were reared on sugarberry (Celtis laevigata) in Gainesville, Florida, in the fall of 2016, resulting in 130 pupae. Caterpillars were maintained in large clear cellophane bags under a natural light regime in an unheated room with windows. Automeris io caterpillars take 2–3 months to develop undergoing 6 (males) or 7 (females) instars. The previously recorded time-lapse photography of the pupation process (Sourakov & Schlachta, 2016; Sourakov & Schlachta, 2017) allowed for the estimation of age of prepupae and pupae. While the caterpillars were pupating in November, when temperatures fluctuated daily between 10 and 25°C, the pupation process was monitored visually through a slit in a cocoon. It takes ca. 7 days from cocoon spinning to pupation and about 8 hours for a pupa to change color from green to dark brown. Hence, the approximate age of a pupa can be judged relatively easily up to 8 hours after pupation (Supplementary Figure S5).

Ten pupae of different stages, but not older than 8 hours, were injected using a 10µl syringe and 10µl (1 drop) of heparin solution. The solution was obtained by purchasing 50 mg of heparin sodium salt from porcine, manufactured by MP Biomedicals, Inc. and adding 0.1 ml of distilled water. Additionally, two late-stage prepupae were injected within a day of pupation with 5µl and 10µl of heparin solution. Injections of pupae were conducted under the wing through the soft cuticle separating thorax from abdomen. The two prepupae were injected in the side approximately at the mid-length on the 7th day after cocoons were spun and within a day of pupation. Upon injections, prepupae and pupae were placed back into their cocoons and in individual plastic bags with paper towel available for the moth to perch on upon emergence. Additionally, 12 prepupae were subjected to cold shock in the refrigerator (7°C) for 24 hours. During diapause, all pupae were kept under ambient light conditions, with temperatures fluctuating between ca. 10 and 30°C, until most of the untreated pupae (75%) and some of the injected ones emerged during May-August 2017.

Results and discussion

While control pupae and most of the individuals cold-shocked as prepupae had an emergence rate of 75%, and showed little or no deviation from the expected wing pattern, most of the pupae that were injected by heparin did not emerge. The three individuals
that emerged (two males from injected prepupae and one female from a pupa injected ca. 5 hours after pupation) exhibited a substantial variation from the norm (Figure 1 and Figure 2). Additionally, upon close examination and dissection of cocoons and pupae from the rest of the experimental group, five more males were recovered, four of which showed significant deviation from the norm.

A transformed female is illustrated in Figure 1A. Injection may have damaged the right hindwing, so it did not spread properly (Figure 1A.ii), but the left side was structurally intact and strikingly different, with the hindwing almost entirely black due to the expansion of the black ring around the eyespot. The control sibling female is illustrated in Figure 1B for comparison. Also, in Figures 1C and 1E, the slight changes in the black ring around the eyespot exhibited by two females cold-shocked as prepupae are illustrated next to control siblings.

Two aberrant males, whose prepupae were injected with 5µl and 10µl of heparin within a day of pupation are illustrated in Figure 2. The one that received a smaller dose was only slightly aberrant in its dorsal hindwing eyespots (Figure 2A). There, the black rings have smudges extending towards the wing base akin to comet tails, hence the aberration is nicknamed “Comet Eye,” following the tradition started by Manley (1978, 1990), who gave genetic aberrations of Automeris io names, such as “Broken eye,” (Figures 3B and 3C) and “Teardrop” (Figure 3D).

The male from prepupa that was injected with 10µl of heparin was transformed much more drastically (Figure 2D), with symmetrical changes in forewing DI elements on both dorsal and ventral surface akin to these of the aberrant female in Figure 1A. The black rings around the hindwing eyespots underwent similarly broad expansions that are not symmetrical in left and right hindwing. In the aberrant male, it is more apparent that, while the ring around the eyespot expanded, the outer black band of hindwing underwent little or no change. A name “Black Eye” is proposed for the eyespot aberrations shown in Figure 1A and Figure 2D. The ventral surface of the wing in “Black Eye” shows considerable expansion and diffusion of the small and compact black ring of control specimens around the small white ventral eyespot that corresponds to the DI element of the dorsal forewing surface (Figures 1A.ii vs. 1B.ii and Figures 2D.ii vs. 2B.ii). In the “Comet
Eye,” the changes in DI element are only noticeable on the ventral surface (Figure 2A.ii), and it is very likely that the differences in the degree of transformation between the two aberrant males in Figure 2 can be explained by the difference in the dose of heparin they received. Serfas & Carroll (2005) observed no asymmetry in the action of heparin even when it was injected only in one forewing, explaining this observation by suggesting that it influences “the secretion of cold shock hormone by a structure located near the body midline, rather than acting on receptor function within the wing.” While the “Comet Eye” aberrant and the forewing changes in “Black Eye” support this hypothesis, the asymmetry of hindwing pattern changes in male “Black Eye” in Figure 2D, as well as some additional observations provided below, suggest otherwise.

Heparin injections are known to enhance wingless gene signaling and have been previously shown to modify forewing DI elements across Lepidoptera (Martin & Reed, 2014). Özsü et al. (2017) recently showed that wingless is a regulator of eyespot color patterns in Bicyclus anynana butterflies. It is also quite possible, based on the present study, that wingless is involved in controlling the black ring around A. io eyespot (see reviews of Version 1 of this paper by Martin (2017) and Marcus (2017)). Martin & Reed (2010) proposed that DI elements are found in both forewing and hindwing and that the hindwing ones should “be considered as serial homologs of their forewing counterparts.” Perhaps the black ring of the hindwing dorsal eyespot is homologous with the black ring element of the forewing ventral eyespot as they are positioned very similarly in relation to their respective wing venation (Supplementary Figure S1 & Supplementary Figure S2), and both react similarly to heparin injections. It is interesting to note that despite the dramatic changes to wing pattern, the ventral hindwing surface remained identical in experimental and control individuals (Figures 1A.ii vs. 1B.ii; Figures 2D.ii vs. 2E.ii).

Iwata & Otaki (2016), showed correlation between scale size and color in various wing pattern elements of nymphalid butterflies. The close examination of normal dorsal hindwing eyespot reveals that there are at least four different types of scales that form the color pattern in this area: the eyespot center is formed by short, flat scales that are white under normal light and fluorescent under UV; in the blue area of the eyespot, these scales intermix with similarly shaped iridescent blue scales (Figure 2F.iii and Supplementary Figures). These two types of scales form the eyespot proper and are surrounded by a ring with a top layer of longer, flat, non-iridescent black scales. Beyond the black ring lies yellow or pink wing areas the surface of which are covered with bristle-like scales. In the eyespot transformed by heparin, the expansion of black ring is associated with the expansion of the number of the
black-ring type surface scales into the yellow areas of the wing (Figure 2D.iii). Some of these scales appear separately from the black ring, which can be explained by the intracellular communication between cells within the developing wing membrane via epithelial feet that may be mediating cell rearrangement (Nardi & Magee-Adams, 1986). Marcus (2017) suggests that observed change may be happening on the level of color pattern determination processes that involves cell-cell signaling by signal transduction processes.

Among the additional five specimens (all males) from dissected dead pupae that were all injected 10µl of heparin solution, the one that was given an injection as a green pupa at about 3 hours after pupation showed no transformation, while four that were injected as brown pupae at ca. five hours after pupation underwent strong transformation that is quite striking even though the wings are not expanded (e.g., Figures 4B,C vs. control specimen in Figure 4A). One of the transformed individuals, shows a higher level of change in the ventral forewing (Figure 4C.ii) and dorsal hindwing (Figure 4C.iii). Unlike all other A. io modified by heparin injections, the hindwing eyespot is not visible, yielding its place to black scales underlined with scales that appear translucent and produce light diffraction. The dorsal hindwing margin colors and elements are absent in this individual. This level of transformation is comparable to the “Caecus” aberration found in the wild-collected specimen (Figure 3A). The close examination of the eyespot area in this latter wild specimen (Figure 3A.i) suggests that it consists of short, flat, iridescent-black-blue scales that differ from long, black scales of the surrounding wing area. In other words, the “Caecus” aberration is the result of the disappearance of the fluorescent white scales and the expansion of the black-ring type scales all the way to the normally yellow hindwing margin.

Such differences in the shape of the scales forming different wing pattern elements is not restricted to the dorsal hindwing. When examining the ventral DI elements of non-expanded forewings more closely in control and transformed moths, shown in Figures 4A.ii and C.ii, it is easy to note that they, too, are formed by differently shaped, flatter, and shorter scales than the surrounding yellow areas that are formed by longer, bristle-shaped scales (Figures 4A.iiiz and 4C.iiiz). This also supports the idea that ventral forewing eyespot and dorsal hindwing eyespot, different as they are to the human eye, are homologous organs.

It must be noted that the transformations associated with heparin injections may be more profound than changes in the color and shape of individual scales. In at least one of the four transformed moths (the one whose dorsal hindwing resembles “Caecus” aberration and is illustrated in Figure 4C), the forewings are not symmetrically modified, with little pattern appearing on the right pale forewing (Figure 5A). Also, during dissections of the dead pupae injected with heparin, the underlying surface of the forewing was revealed (Figure 5B) and the black pattern (Figure 5B.i) and change in fluoresce properties (Figure 5B.ii) as compared to the control (Figure 5C) suggest that transformation of scales is accompanied by the changes in the underlying wing membrane. Laboratories involved in research on Lepidoptera wing and scale embryology (e.g., Dinwiddie et al., 2014) may find it an interesting avenue of research to pursue.
The wing pattern research has now entered the phase when functions of individual genes are being rapidly revealed (e.g., Marcus, 2005; Monteiro et al., 2006; Zhang et al., 2017). It is hoped that the present publication, while documenting unique aberrations in a single species, will be useful in the future work directed at understanding wing pattern evolution and development, in general, and will prompt additional experiments that will clarify the observations presented here.

Competing interests
No competing interests were disclosed.

Grant information
The author(s) declared that no grants were involved in supporting this work.

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**Supplementary material**

**Supplementary File.** Figures S1–S5.

**Figure S1.** Forewing venation in *Automeris io* and position of ventral eyespot in relation to venation. A. Whole forewing with pigmentation cleared; B. Ventral eyespot. (i) Intact eyespot, photo in LED light; (ii) Scales partially removed to expose venation, photo in UV + incandescent light. Wing venation nomenclature after Heppner, 1998. Photos by Andrei Sourakov

**Figure S2.** Hindwing venation in *Automeris io* and position of dorsal eyespot in relation to venation. A. Whole forewing with pigmentation cleared; B. Dorsal eyespot in UV + incandescent light. (i) Intact eyespot; (ii) Scales partially removed to expose venation. Wing venation nomenclature after Heppner, 1998. Photos by Andrei Sourakov

**Figure S3.** Scales involved in formation of dorsal hindwing eyespot in *Automeris io*. A. Dorsal eyespot with some of the scales removed. B. Intact eyespot. Photographed in LED light (i) Whole eyespot; (ii) Close-up of eyespot center; (iii) Close-up of black-ring/yellow field border. (1) white center corresponding to underlying vein, (2) blue part of eyespot, (3) black ring, (4) surrounding yellow field. Photos by Andrei Sourakov

**Figure S4.** Three types of scales removed with scotch tape from the dorsal hindwing eyespot of *Automeris io*. A. First layer of scales. B. Second layer of scales. Photographed in LED light. C. Third layer of scales. Photographed in UV + incandescent light. (1) white center corresponding to underlying vein, (2) blue part of eyespot, (3) black ring, (4) surrounding yellow field. Photos by Andrei Sourakov

**Figure S5.** A. Staging of pupae: time-lapse photographs of a pupa of *Automeris io* reflect the time since pupation. B–D. Two seven-day-old prepupae and (D) a representative ca. five-hour-old pupa that were injected with heparin and resulted into transformed moths. (i) and (ii) shedding of larval skin… (iv) 2 hours and 3 minutes after pupation, etc. Photos by Andrei Sourakov & Steven Schlachta.

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Open Peer Review

Current Referee Status:  

Version 2

Referee Report 04 October 2017

doi:10.5256/f1000research.13785.r26373

Arnaud Martin
Department of Biological Sciences, The George Washington University, Washington, DC, USA

The author went out his way to improve the first version, and added valuable observational data on replicated experiments. I feel the data is preliminary but useful, and carefully discussed.

Competing Interests: No competing interests were disclosed.

Referee Expertise: Developmental genetics, Lepidoptera

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Referee Report 02 October 2017

doi:10.5256/f1000research.13785.r26372

Jeffrey M. Marcus
Department of Biological Sciences, University of Manitoba, Winnipeg, MB, Canada

I congratulate the author for making revisions to create a much improved paper. Thank you for the opportunity to review this work.

Competing Interests: No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 1

Referee Report 17 August 2017

doi:10.5256/f1000research.13271.r24774
Jeffrey M. Marcus
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This manuscript describes a very promising approach to expanding the experimental study of color pattern development beyond the select group of model butterfly species that have received the bulk of the attention thus far. The Saturniid moth Automeris io was used for these experimental studies. Also described are A. io specimens derived from breeding experiments and wild-caught specimens with aberrant phenotypes.

While I enjoyed reading the manuscript, there are a number of changes that I would like to suggest to the author:

Clarification of methods:
There are a number of methodological details that should be clarified.
1. “sugarberry”. Please give scientific name. Is this Celtis laevigata?
2. Staging of pupae. As much detail as possible should be given here about when and how the injections were done relative to the pupal molt. How many hours at what temperature? Was the cuticle sclerotized yet? Where on the experimental specimens’ bodies did the injections take place? Under what conditions were the animals allowed to recover from their injections? What was the manufacturer and purity of the heparin used? For coldshock, how many hours is “overnight”?
3. Control injections: what was the rationale for using manitol? Many experiments pair heparin with chondroitin sulfate B (keratan sulfate) as a negative control because it is structurally similar to heparin, but lacks the biological activity associated with heparin.
4. Overwintering conditions: the author reports that the experimental animals were kept at ambient conditions until emergence the following spring. Were they kept indoors or outdoors? If indoors what was the typical temperature and lighting conditions during this period? If outdoors, try to provide descriptors of climatic conditions during the appropriate period using National Weather Service or other data.

Clarification of results:
Additional clarification of the results is also required.
1. The author reports that most of the injected individuals did not emerge. How many failed to emerge? Were there any differences in eclosion rate between the heparin injected individuals and the chondroitin sulfate B injected individuals?
2. What was the emergence rate of unmanipulated individuals? Was it different from the emergence rate of cold-shocked individuals?

Reinterpretation of Results and Elaboration of Discussion:
1. The author should keep in mind that Lepidopteran color patterns can be altered by manipulation of the developmental processes responsible for determining color patterns as well as by manipulation of the developmental processes responsible for color pattern differentiation. Determination processes might involve cell-cell signaling by signal transduction processes, while differentiation of color patterns involves the expression and regulation of biosynthetic pathways responsible for pigment synthesis. Manipulations such as cold-shock might have effects on both kinds of developmental processes, if they are taking place at the time of the manipulation. Additional
references to prior work that examines the effects of coldshock (Nijhout 1984; Serfas and Carroll 2005; Mahdi et al. 2010; Dhungel and Otaki 2013) might be warranted.

2. Everything that we know about the action of heparin suggests that it interacts with signal transduction pathways such as wingless/wnt (Binari et al. 1997), and there is no evidence that it interacts with melanin pigment biosynthesis. Discussing the experimental results of heparin injection from this study in reference to what is known about wingless/wnt signaling and its effects on color pattern determination in insects (Carroll et al. 1994; Monteiro et al. 2006; Martin and Reed 2010; Werner et al. 2010) would be highly desirable. I also urge the author to extend the discussion further and to connect his work with prior work on Saturniid moth color pattern development including both classic cautery studies (Henke 1933; Henke 1944) and studies of gene expression (Monteiro et al. 2006). Interestingly, the discal eyespots of Automeris io are positioned on top of crossveins, structures that also employ wingless/wnt signaling during their development (Conley et al. 1997; Marcus 2001). Relating eyespot development to underlying crosvein development in Saturniid moths, perhaps through modulation of the wingless/wnt pathway would be a very interesting research trajectory to pursue. Understanding how discal eyespots in some Saturniid moths are both similar to and different from border eyespots in butterflies would be a very interesting avenue for further studies of the evolution and development of Lepidopteran color patterns.

I would like to encourage the author to consider these points in his revisions and also to continue his experimental explorations in future work.

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**Is the work clearly and accurately presented and does it cite the current literature?**
Partly

**Is the study design appropriate and is the work technically sound?**
Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**
No

**If applicable, is the statistical analysis and its interpretation appropriate?**
Not applicable

**Are all the source data underlying the results available to ensure full reproducibility?**
Yes

**Are the conclusions drawn adequately supported by the results?**
Partly

**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Referee Report 04 August 2017

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Department of Biological Sciences, The George Washington University, Washington, DC, USA

This manuscript presents some exciting pattern aberrations observed in the eyespots of the Io Moth, including specimens obtained from drug injections, a wild-caught individual, and previously undescribed collection specimens.

I enjoyed reading the manuscript and thank the author for publishing this. The reported phenotypes provide valuable information on the formation of eyespot rings in this particularly spectacular species.

I noticed the following minor issues.
What was the rationale for injecting mannitol?

Serfas and Caroll (2005) -> Carroll with two r (typo)

Methods: "Ten randomly selected pupae were injected"
Please provide more details if possible on the staging. Were they tender and relatively fresh? Is there a way to determine the maximum age they were at, or a time estimate based on their cocoon spinning?
5mg of heparin is a large dose, so it may have killed the younger pupae, while the aberrant female came out by chance by being at a more resilient stage (or was accidentally injected with much less compound)

Methods, please confirm the injected concentration of the injected heparin (apparently 0.5mg/uL), and provide the exact origin of the compounds (sodium salt? molecular weight? manufacturer?)

Figure 1 Legend: the abbreviation for DI BR and EIII are missing.

The author is wrong calling the outer black line “EIII”. Schwanwitsch 1956 and Henke 1936 (in 3 other Saturniids) suggest this is MI (central symmetry system outer band). I am personally inclined to say that while it may not be M1 in A. io … it is certainly not EIII

“A less aberrant male, whose prepupa was injected with 5mg of heparin a day before pupation, also emerged”
I believe the author meant “A less aberrant male, whose prepupa was injected with \textbf{2.5mg} of heparin \textbf{ONE} day before pupation, also emerged”

“Heparin injection must have enhanced or prolonged the process of expansion of black pigment once it formed in the black ring around the blue scales.”

There is a way to deepen the discussion here, and I will try to explain briefly (feel free to use this suggestion).

The A. io Discal Spots (DI = Discalis I element of the Schwanwitsch pattern homology system) are stereotypical patterns that are always overlapping with the discal crossvein. Martin and Reed (2010) suggests that these spots always express the wingless morphogen in Lepidoptera, and Martin and Reed (2014) also shows that DI elements are responsive to heparin treatment in nymphalids. Interestingly, heparin is known to enhance wingless signaling in Drosophila: Baeg et al. (2001), Binari et al (1997) and Greco et al (2001).

Thus, the eyespot ring expansions observed upon heparin injection here suggest the exciting possibility that wingless, or perhaps other heparin-sensitive morphogens, are deployed during pre-pupal and pupal development to pattern different aspects of the discal ocelli structure.

To be honest, the current insight about melanization is a little too phenomenological, because melanin pigment synthesis happens much later in development and there is no known interaction between heparin and melanin biosynthesis...
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Is the work clearly and accurately presented and does it cite the current literature?
Yes

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Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Partly

If applicable, is the statistical analysis and its interpretation appropriate?
Not applicable

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

*Competing Interests*: No competing interests were disclosed.

*Referee Expertise*: Developmental Genetics, Lepidoptera

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.