Exploring Gloger's ecogeographic rule

Why organisms are darker in wetter & warmer environments



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Gloger's rule is an ecogeographic trend wherein organisms possess darker pigmentation in warmer, more humid environments (Gloger 1833; Gaston et al. 2008; Lev-Yadun 2015; Roulin & Randin 2015; Bishop et al. 2016). Gloger (1833) first described this pattern in birds, and consequently the rule is largely treated as applying to endothermic animals (Bishop et al. 2016). However, evidence mounts for the pattern's applicability to ectotherms (e.g. Rapoport 1969; Bishop et al. 2016), and plants (e.g. Lev-Yadun 2007; Koski & Ashman 2015). Though, there may be different processes in different taxa that result in this pattern of organismal pigmentation across climatic gradients.

Herein, I discuss the treatment of ecogeographic rules in ecological and evolutionary science, and within this context discuss the degree to which Gloger's rule has merit as a genuine pattern in nature. I discuss insights into possible mechanisms that generate the Gloger's rule pattern, and how these vary between taxa, and across levels of biological organisation. I argue that there is sufficient evidence to support Gloger's rule as a useful and true generalisation.

Ecogeographic & biogeographic "rules"

The theory vs empiricism of "rules" ...

It is important to consider what is mean by the term "rule" in biology. There are many of biogeographic and ecogeographic patterns that are referred to as rules, e.g. Bergmann's rule, Allen's rule, etc. Gloger's rule, the focus here, is also often grouped with these.

Lomolino et al. (2006) outline the study of observed patterns or trends of organismal traits across geographical space with ecogeographic rules (i.e. observations). They highlight examples of *patterns* at broad ecological scales as consequence of *processes* at a range of scales. Indeed, the patterns themselves are also manifest at a variety of scales and levels of organisation. As such, Lomolino et al. (2006) notes, underlying causal mechanisms behind these patterns are difficult to ascertain.

Lomolino et al. (2006) also talk about theory vs empirical evidence, and varying scales, and also the *evolutionary* side to *ecog*eographic rules.

Olalla-Tárraga (2011) discusses approaches to studying rules too, with focus on Bergmann's rule. Olalla-Tárraga (2011) advocates a pluralist approach, wherein the manifestations of ecogeographical rules at multiple scales and levels of biological organisation are considered. He also

outlines that "laws" and "rules" in ecology, and in science generally, need not *always* contain mechanistic statements, and need not be without exception, as they are "correlative generalisations". Though, mechanistic understandings of the processes that generate patterns is desirable, and indeed often the aim of research, this does not imply that correlative ecogeographic rules are not useful intrinsically. Not least, these rules are often the starting point of research. Thus, these rules represent interesting observed patterns that can motivate research, *and* useful generalisations that can be employed in other work.

Ecogeographic rules operate across various levels of biological organisation. Gaston et al. (2008) differentiate between *intra*specific, *inter*specific, and *assemblage* patterns in organismal traits across environmental gradients. Bishop et al. (2016), for example, describe community trait averages in order to describe the extent of Gloger's rulein ants.

Millien et al. (2006) discuss ecotypic (i.e. intraspecific, i.e. between populations) variation in terms of ecogeographic rules, too.

Booth (1990): Ontogenetic level too!

Gloger's rule—evidence & examples

Origins of the rule ...

As stated above, Gloger's rule stems from his observations of bird plumage darkness across climatic moisture gradients (and, by the simplest extension, pigementation in endothermic animals too). Roulin & Randin (2015) describe the logic of Gloger's rule lucidly:

The rule is based on the assumption that melanin pigments and/or dark coloration confer selective advantages in warm and humid regions.

They investigate animal fitness in warm, humid environments as conferred via alternative phenotypic adaptations (viz. in owls). The implication of *fitness* in a given habitat allows one to see the *evolutionary* aspect to Gloger's *ecogeographic* rule (a sentiment Lomolino et al. (2006) highlights).

Studies concerning the rule, and indeed most patterns in nature (Lomolino et al. 2006), must ideally consider the historical, evolutionary context to organismal form—not just local deterministic mechanisms.

Though, it must be noted, the extent to which historical and deterministic processes govern organismal pigmentation largely depends on the scale of investigation. Ontogenetic changes in pigementation are a-historical. Intraspecific patterns may represent micro-evolutionary fitness benefits to certain genotypes. Interspecific and assemblage patterns, however, would be indicative of more long-term, persistent selection pressures—e.g. ant species assemblages follow Gloger's rule in humid environments, implying that there may be long-standing, consistent strength to Gloger's rule in ants. In other words, enough evolutionary time must pass

[...].

Conversely, with respect to interspecific and community ecogeographic patterns, environmental filtering could conceivably produce trait patterns over shorter, ecological timescales. In this case, if the traits in taxa that follow an ecogeographic rule (e.g. Gloger's rule) do confer a fitness benefit as the rule would predict, they would not be adaptations (sensu Baum, Mayr, [...]), having evolved elsewhere.

Distinguishing between whether organismal pigmentation follows the Gloger's rule pattern (if and when it does) is governed by deterministic (i.e. environmental filtering) or historical (i.e. adaptation) process would be insightful. Research concerning this distinction would further allow one to ascertain the degree to which Gloger's rule is a consequence of multiple processes (at multiple spatio-temporal scales), and the number and relative strength of processes that partake therein. In such a case that these patterns have, say, historical drivers in one system at one scale, and deterministic drivers in another, this would emphasise the complexity of ecogeographic patterns. It would also mirror the notion Olalla-Tárraga (2011) puts forward, wherein correlative generalisations, such as Gloger's rule, need not encapsulate one, if any, mechanisms. Though, it is unlikely that there is no mechanisms behind Gloger's rule patterns. Instead, it is interesting and insightful that multiple mechanisms, combined or in isolation, govern these patterns.

Manifestations of Gloger's rules across various scales and levels of organisation [...] intrasp. Gloger's rule manifestations (= "ecotypic", sensu Millien et al. (2006)) svs intersp./community level manifestations (Lev-Yadun 2015)

Evidence & examples

Gloger vs anti-Gloger patterns (sensu Lev-Yadun 2015), e.g "negative Gloger's rule" in invertebrates, e.g. Collembolates (Rapoport 1969)

Does the pattern exist? Is it observed in the first place? Dissect the observations at the intrasp., intersp., and assemblage levels

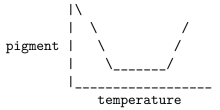
Kamilar & Bradley (2011) primate *inter*specific coat colour follows Gloger's rule -> little mechanistic work in this paper though! They speculate about 1) background matching, 2) anti-bacterial stuff, and 3) maybe thermoregulation (#FutureStudies). Unlikely to be UV, because primates live in trees! (not even in the upper-canopy, where UV is strong—primates do not spend tonnes of time up there)

VanderWerf (2012) looked at bird body size, bill size, and plumage in populations two closesly related birds across environmental gradients in Hawai'i. With regards to plumage, bird individuals (N.B. this is basically intraspecific-scale work!), Gloger's rule was supported, and with "smoothly clinal distributions" [sic] (as a function of thew few dispersal barries and steep environmental gradients.

Bishop et al. (2016) ant *assemblage* work (ECTOTHER-MAL):

- organismal darkenss as a modal pattern
- at low temperatures for thermoregulatory needs

- at higher temperatures for UV-B protection
- · also darkness incr with smaller body size
- these patterns are evident at the assemblage level



(confounded w/ rainfall, sexual selection)

Plants vs endothermic animals vs ectothermic animals; UV-B protection (Gloger's rule) vs thermal hypothesis vs dessication hypothesis (Pinkert et al. 2016); Plants foliarly (sensu Lev-Yadun 2007) vs florally (sensu Koski & Ashman 2015)

Animal vs plant e.g. -> differences in meaning and interpretation. Also consider Dominy & Lucas (2004) food colour and primates—could THIS connect to an environmental pattern in plant colour?

Mechanisms behind the pattern

Burtt & Ichida (2004) found that dark pluamge is resistant to bacterial degradation (\sim pigments), a common problem in humid climes; methods: measure intrasp. Δ colour vs bacterial activity.

cf Koski & Ashman (2015) -> UV role (in plants)

Tate et al. (2016) found persistent colour polymorphism within a populations of a species -> equivalent fitness of the morphs in heterogeneous habitats; results: darker species forage/hunt better in darker habitats (~ hiding in the ambient background). Since (Ruan says) darker habs are wetter (ish), this relates to Gloger's rule.

- Connect evidence above to mechanisms described in their respective papers (if applicable)
- & mechanisms from other papers concerning pigment and environment (e.g. Tate et al. 2016)
- Dissect the observations at the intrasp., intersp., and assemblage levels MECHANISTICALLY
- Animal vs plant e.g. -> differences in meaning and interpretation for MECHANISM

Concluding remarks

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Millien et al. (2006) traits vary with geography, but also with global cliamte change!

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