Using connectivity to identify climatic drivers of local adaptation

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Abstract

Despite being able to conclusively demonstrate local adaptation, we are still often unable to objectively determine the climatic drivers of local adaptation. Given the rapid rate of global change, understanding the climatic drivers of local adaptation is vital. Not only will this tell us which climate axes matter most to population fitness, but such knowledge is critical to inform management strategies such as translocation and targeted gene flow. While simple assessments of geographic trait variation are useful, geographic variation (and its associations with environment) may represent plastic, rather than evolved, differences. Additionally, the vast number of trait–environment combinations makes it difficult to determine which aspects of the environment populations adapt to. Here we argue that by incorporating a measure of landscape connectivity as a proxy for gene flow, we can differentiate between trait–environment relationships underpinned by genetic differences versus those that reflect phenotypic plasticity. By doing so, we can rapidly shorten the list of trait–environment combinations that may be of adaptive significance. We demonstrate how this reasoning can be applied using data on geographic trait variation in a lizard species from Australia’s Wet Tropics rainforest. Our analysis reveals an overwhelming signal of local adaptation for the traits and environmental variables we investigated. Our analysis also allows us to rank environmental variables by the degree to which they appear to be driving local adaptation. Although encouraging, methodological issues remain: we point to these issue in the hope that the community can rapidly hone the methods we sketch here. The promise is a rapid and general approach to identifying the environmental drivers of local adaptation.
Significance Statement

Despite 150 years of evolutionary research, we still do not have a good method for determining the climatic drivers of local adaptation. Having such a method is critical if we are to understand and mitigate the impact of climate change. Here we exploit the fact that local adaptation is eroded by gene flow to develop a new method for identifying the environmental drivers of local adaptation. Our method will allow workers around the world, for the first time (and often using existing datasets), to determine the climatic drivers of local adaptation in their system. As such, the method has powerful implications not only for pure evolutionary research, but also for management and conservation.
Introduction

It is only recently that we have begun to appreciate the speed with which evolution can happen; not only over relatively short timespans (e.g., 1, 2-4), but also at small spatial scales (5). Rapid local adaptation has been recorded in response to a wide suite of environmental drivers, including invasive species, and pollution (6). We expect climate to also be a major driver of local adaptation (e.g., 7, 8), and understanding the way in which species respond to climate is of increasing importance because anthropogenic climate change is proceeding at such a rate that there are concerns that many species will be unable to evolve rapidly enough to avoid extinction (9, 10).

Evolution typically optimizes phenotypes, but the optimum will vary through both time and space (11, 12), in turn leading to populations ('demes') that have, on average, higher fitness in their home environment than an immigrant would: local adaptation. While adaptive optima for traits almost always vary geographically, it does not follow that all geographic trait variation is due to local adaptation. Geographic trait variation can arise due to other factors, such as phenotypic plasticity (including developmental plasticity and maternal effects), neutral clines, and environmental factors (such as geographic variation in fitness-reducing parasites). These factors can give the appearance of local adaptation (10, 11), complicating our identification of climate-relevant adaptive variation.

To circumvent these issues, evolutionary biologists use experimental approaches to demonstrate local adaptation (12, 13). Experiments designed to detect local adaptation typically utilise one of two techniques: 1) reciprocal transplants, which are done in situ, and are considered the gold standard for demonstrating local adaptation; or 2)
common garden experiments, which are usually done in the lab where it is easier to control each environmental variable (12). Both of these techniques can be difficult, for reasons of time, expense, logistics, or ethics. This difficulty increases as the number of separate demes and environmental variables to be tested increases and as the generation time of the organism increases (12). Additionally, although reciprocal transplants will detect signs of local adaptation, they are not necessarily suited to identifying the environmental drivers of that local adaptation (14). This is because in situ reciprocal transplants necessarily encompass all the environmental variables that differ between the transplant locations. Lab-based common garden approaches may, in principle, be more suited to identifying environmental drivers (because the environment may be under a degree of control), but in practice it often remains impossible to identify the environmental drivers of trait variation seen in the wild. Thus, the best experimental tools we have for studying local adaptation are demanding in terms of time and cost, and are unsuitable for assigning environmental drivers (such as climate variables) to adaptive variation. If we are looking for climate-driven local adaptation, this is a problem: we want to know which climate variable or variables are the main drivers of adaptation, and we urgently need this information for many species.

By definition, local adaptation has a genetic basis and is consequently weakened by high levels of gene flow (11, 15, 16). Demes with excessive inward gene flow are therefore likely to be less optimally adapted, causing an observable mismatch between optimal and actual phenotypes. Some examples of this are birds dispersing and producing clutch sizes that are not optimised for the habitat quality in which they are now nesting (17), larval salamander colouration not matching streambed colouration
due to high levels of gene flow from nearby but predator-free streams (18), and stick
insects in smaller habitat patches having non-cryptic colouration when the
surrounding patches are larger and environmentally dissimilar (19). These
observations of "migrant load" suggest an alternative technique for identifying and
assessing local adaptation. First, we look across populations for relationships between
the environment (e.g., mean annual temperature) and traits (e.g., morphology,
physiology). By themselves, these relationships are not sufficient evidence of local
adaptation — they could also be caused by phenotypic plasticity. Second, knowing
that local adaptation is hindered by gene flow, we can look at whether gene flow
diminishes the environmental effect. With some caveats (discussed below), in cases
where data on gene flow are absent (which is often the case), habitat connectivity can
be used as a substitute for gene flow. Trait–environment relationships that are strong,
but which are also weakened by connectivity, are indicative of trait–environment
relationships that have a genetic basis. In a statistical model, this idea would be
represented as follows:

\[ \text{Trait}_i = A + B_{\text{env}} \times \text{Env} + B_{\text{conn}} \times \text{Conn} + B_{\text{int}} \times \text{Env} \times \text{Conn} + \text{error}_i \]

Where:

\[ \text{trait}_i = \text{trait value for individual } i \]

\[ A = \text{intercept} \]

\[ B_{\text{env}} = \text{coefficient of the environmental variable} \]

\[ \text{Env} = \text{environmental variable (e.g., annual mean temperature) at the} \]

\[ \text{individual’s site} \]

\[ B_{\text{conn}} = \text{coefficient of connectivity} \]

\[ \text{Conn} = \text{connectivity at the individual’s site} \]

\[ B_{\text{int}} = \text{coefficient of the interaction between Env and Conn} \]
\[
error_i = \text{deviation between expected value and trait value of individual } i
\]

Which, with slight rearrangement, can also be expressed as:

\[
\text{Trait}_i = A + (B_{\text{env}} + B_{\text{int}} \times \text{Conn}) \times \text{Env} + B_{\text{conn}} \times \text{Conn} + error_i
\]

showing that the slope of the relationship between the trait and the environment now depends on the connectivity value. When the signs of \(B_{\text{env}}\) and \(B_{\text{int}}\) are in opposition, then we have a situation in which the relationship between the trait and the environment diminishes as connectivity increases.

If we now collect data on a large number of trait–environment relationships, and their interaction with connectivity, we can imagine several possible patterns emerging. These possibilities are depicted in Figure 1. Each panel represents a possible relationship between trait–environment coefficients (along the x-axis) and the interaction between environment and connectivity (y-axis). Panel A shows a set of trait–environment relationships that vary in strength, but that are not influenced by connectivity (i.e., no environment–connectivity interaction). This pattern is indicative of a system in which trait–environment relationships are predominantly driven by plastic responses of traits to their environment (i.e., traits always match the local environment, regardless of the level of inward gene flow). Panel B shows a system in which trait–environment relationships are eroded by connectivity: increased connectivity diminishes the relationship between the environment and the trait. In this situation, the interaction between the environmental variable and connectivity is negative when the environmental coefficient is positive (i.e., greater connectivity causes the environmental coefficient to decrease towards zero; bottom-right quadrant), and positive when the environmental coefficient is negative (i.e., greater connectivity causes the environmental coefficient to increase towards zero; top-left quadrant).
This is the pattern we would expect if there is a genetic basis to the trait–environment relationship, such as is exhibited by local adaptation. Panel C shows the situation where the effect of the environment tends to be enhanced by connectivity. This pattern might arise in organisms that are highly mobile and can actively move to their ideal environment, thus avoiding the selective pressures that would lead to local adaptation.

Understanding how species respond to specific aspects of their environment is vital if we are to have any hope of halting the current rapid loss of biodiversity. Climate change is undoubtedly one of the biggest threats to global biodiversity (20, 21), and conservation biologists are looking to a variety of techniques to assess and help mitigate the impacts of climate change on vulnerable species (22-24). One technique that is likely to see increasing use is targeted, or assisted, gene flow [TGF; for review, see (22, 25)]. This technique involves the spatial redistribution of long-standing adaptations, and acts to increase genetic diversity in recipient populations, thereby bolstering capacity for evolutionary adaptation (10, 22, 24, 25). When applying TGF to help species adapt to climate change, we need to find an existing location that matches the future climate at our recipient site, and then translocate animals from that source location. It is a simple idea, but climate is multidimensional and species will not be adapting equally to each climate axis: is a difference of 0.5°C in mean temperature more important than a difference of 100mm in annual rainfall? The answer depends upon which aspects of climate (hereafter “climatic axes”) have the strongest influence on fitness.
Here we explore the idea of using connectivity to infer local adaptation. To do this we develop a case study of a lizard species from northern Australia. We use this system to examine the relationship, across sites, between traits and climatic variables. We assess how habitat connectivity affects these relationships and use the interaction between the environmental variable and connectivity to rank trait–environment combinations. In doing so, we reveal a set of trait–environment relationships that appears to be dominated by local adaptation.

**Methods**

**Study species and site selection**

The Rainforest Sunskink (*Lampropholis coggeri*) is a small (snout–vent length up to 45 mm), diurnal scincid lizard restricted to the rainforests of the Wet Tropics region of northeastern Australia (26). The rainforests of this region cover a wide range of environmental conditions, spanning significant elevation (0–1600 m ASL), precipitation (annual mean precipitation of 1432–8934 mm, not including input from cloud stripping), and temperature (annual mean temperature of 16.3–25.8°C) gradients. This heliothermic skink is active year-round, often seen basking in patches of sunlight on the rainforest floor. Lizards were captured by hand from sites that were selected to maximize the environmental heterogeneity sampled (Fig. 2).

Morphological measurements were obtained from 532 skinks from 32 sites. Physiological measurements were obtained from a smaller subset of these lizards: 259 skinks from 12 sites. At each site, 8–20 skinks were caught per collecting trip. Following capture, skinks were transported to James Cook University (JCU) in...
Townsville for trait measurement. All procedures involving lizards were approved by the JCU animal ethics committee (projects A1976 and A1726).

Physiological trials

Physiological trials commenced within seven days of skinks being collected from the field; skinks being used only for morphology were measured and released back at their point of capture within seven days. The following measures were taken from each skink (n = 259) during laboratory trials: critical thermal minimum (CTmin), critical thermal maximum (CTmax), thermal-performance breadth for sprinting (breadth80), maximum sprint speed (Rmax), temperature at which sprint speed is optimized (Topt), active body temperature as measured in a thermal gradient (Tactive), and desiccation rate (des) (see Table S1 for further details). Details of trait measurement procedures are detailed elsewhere (see 27, 28).
Morphological measurements

The following measurements were taken from each skink (n = 532) using digital calipers: head width (HeadW); head length (HeadL); interlimb length (ILimbL); hindlimb length (HindLL). Left and right measurements were averaged to obtain one measurement for that trait. We also recorded snout–vent length (SVL), total length, and mass (see Table S1 for further details). All measurements were taken by one person (SLM) to minimize observer bias. All morphological variables were log-transformed prior to regression analyses.

Climatic variables, and connectivity

Because our study aimed to assess adaptation to local climate, various temperature and precipitation variables were extracted for each site (see Table S2 for details). We considered both means and extremes. It is important to consider climatic extremes, because temperature extremes may be increasing faster than mean temperatures (29), and selection may often occur during extreme weather events (30). Many environmental variables are highly correlated (27), so only the less-derived variables were used in analyses, specifically: annual mean precipitation (AMP); seasonality of precipitation (Pcov); precipitation of the driest quarter (Pdry); annual mean temperature (AMT); coefficient of variation of temperature (Tcov); average minimum daily temperature (Tmin); average maximum daily temperature (Tmax); average variance of daily maximum temperature (TmaxVar); and average variance of daily Tmin (TminVar).
Our connectivity index was designed to capture the flux of individuals through a location and is detailed in (31). Briefly, it is a measure of habitat suitability for our focal skink species, averaged over space using a species-specific estimate of dispersal potential. This approach is reasonable for any species exhibiting diffusive dispersal, and similar techniques (though different spatial-weighting functions) can be used for species exhibiting non-diffusive dispersal. As our species is an obligate rainforest-dweller, grid cells in the landscape that are rainforest and that are surrounded by rainforest have high connectivity indices, while grid cells of rainforest surrounded by non-rainforest matrix have low indices. See Table S2 for further details on all variables, and Figure S1 for correlations between all variables.

Analysis

Our analysis aimed to assess: 1) the relationship, across sites, between each trait and each environmental variable; and 2) how connectivity affected each of these relationships (i.e., the interaction between connectivity and environment). To allow comparison of coefficients across variables, and to make interaction effect-sizes meaningful, all trait and environmental variables were standardized so they had a mean of 0 and a standard deviation of 1. Linear models were fitted for each pair of environment–trait variables, with all models including the effect of lizard body size and sex, as well as the interaction between environment and connectivity:

$$\text{trait}_i = A + B_{svl} \times \text{SVL}_i + B_{sex} \times \text{Sex}_i + B_{env} \times \text{Env} + B_{conn} \times \text{Conn} + B_{int} \times \text{Env} \times \text{Conn} + \text{error}_i$$

Where:

$$\text{trait}_i = \text{standardized trait value of interest for lizard } i$$

$$A = \text{intercept}$$

$$B_{svl} = \text{coefficient of SVL}$$
SVL = lizard snout–vent length, to control for effect of body size

$B_{sex}$ = coefficient of Sex

Sex = lizard sex (this species is sexually dimorphic in some morphological traits)

$B_{env}$ = coefficient of environmental variable

Env = environmental variable (e.g., annual mean temperature) at the lizard’s site

$B_{conn}$ = coefficient of connectivity

Conn = connectivity index at the lizard’s site

$B_{int}$ = coefficient of interaction between Env and Conn

$error_i$ = deviation between expected value and trait value of lizard $i$
A score for ranking the strength of local adaptation ($L$) was then calculated as:

$$L = -B_{env} \times B_{int}$$

If the signs of the two coefficients ($B_{env}$ and $B_{int}$) are opposite (which indicates an trait–environmental relationship that is diminished by increasing connectivity, i.e., evidence for local adaptation), $L$ will be positive. If the signs are the same (which indicates an environmental effect being enhanced by increased connectivity, a situation not consistent with local adaptation), $L$ will be negative. Thus, higher numbers on this scale equate to stronger evidence for local adaptation in that environment–trait pair. This score can, in theory, range from $-\infty$ to $+\infty$. Once many environment–trait combinations have been assessed, the coefficients for all pairs can be plotted (see Fig. 1). As described in the Introduction, in a system dominated by local adaptation, we expect to see a negative relationship between $B_{env}$ and $B_{int}$ (Fig. 1B). All analyses were conducted in R v3.2 (32).

**Results**

There was substantial variation in the effect of environment ($B_{env}$) and its interaction with connectivity ($B_{int}$) across climate and trait variables, with $B_{env}$ ranging from -1.8 to 1.61, and $B_{int}$ ranging from -0.73 to 0.78 (Fig. 3). Despite this variation, a clear pattern is evident, with most points in Figure 3 appearing in the top-left or bottom-right quadrants: the quadrants in which the two coefficients have opposing signs, and where we would expect points to fall if trait–environment relationships have a genetic basis. Across these trait–environment combinations there is a distinct negative linear trend (slope= -0.36, p < 0.001). It is especially noteworthy that the trait–environment pairs with the largest coefficients are in the two quadrants indicative of local adaptation.
Overall, physiological traits showed substantially stronger environmental effects (i.e., larger values of $B_{env}$) than did morphological traits, with the largest environmental effects being exhibited by CTmin (AMP: -1.80; Tmax: 1.35; Pdry: -1.55) and CTmax (Pdry: 1.61; AMP: 1.21). Physiological traits also showed stronger interactions between environmental effects and connectivity, again with CTmin and CTmax showing the largest interactions. These trends are apparent when we examine our index of local adaptation, $L$. Figure 4 shows a heatmap of all trait–environment pairs, ranked via reciprocal averaging according to the strength of their local adaptation index. The trait–environment pairs that show the strongest signature of local adaptation appear at the top-left in red. There is a rough divide, with most of the physiological traits on the left and most of the morphological traits on the right. The exceptions are the physiological traits $T_{opt}$ and $R_{max}$, which appear at the far right of the figure.

The two environmental variables that produced the strongest effects (topmost rows in Fig 4) were both precipitation related: annual mean precipitation (AMP) and precipitation of the driest quarter (Pdry). In our system, AMP and Pdry are both highly correlated with connectivity (see Fig. S1). This is expected, because our connectivity index is largely a measure of where rainforest is, and the distribution of rainforest in our study region is driven to a large degree by rainfall.

**Discussion**

Understanding relationships between traits and the environment will help us plan management strategies, such as targeted gene flow (TGF), that can mitigate the
impact of climate change on vulnerable species. Numerous studies have looked for (and found) trait–environment relationships (e.g., 18, 19, 33, 34-36), but the interpretation of these associations is plagued with uncertainty: are they associations due to local adaptation, neutral clines, habitat choice, or plasticity? By acknowledging that gene flow undermines adaptation, we can incorporate connectivity (a proxy for the flux of genes) into our analysis, and in doing so, separate those relationships due to fixed genetic differences, from those due to plasticity or habitat choice.

Local adaptation

In the trait–environment combinations we assessed, physiological traits typically showed a substantially stronger effect of environment ($B_{en}$) than did morphological traits, with the largest environmental effects shown in CTmax and CTmin (Figs. 3 & 4). Physiological traits also generally showed stronger environment–connectivity interactions ($B_{en}$), again with CTmin and CTmax showing the largest interactions. Overall, physiological traits generally showed stronger evidence of local adaptation than did morphological traits. This result is intuitive: we would expect an ectotherm’s physiological traits to be under strong selection from climate (37-39), but the fitness link between morphology and climate is much less clear. Had we also included some environmental variables that had a clearer bearing on morphology, we might have detected stronger trait–environment relationships for morphology. For example, skinks that occur in rockier habitats show various morphological adaptations to that environment (40). Including a measure of rockiness in our set of environmental variables might have allowed us to detect a signal of local adaptation for limb length. Here, however, our focus is on climatic aspects of the environment.
Of the environmental variables used, our analysis suggests that precipitation is a very strong driver of local adaptation, even in thermal traits that might not seem obviously related to precipitation (e.g., CTmin, CTmax). Although this may seem a surprising result, precipitation has been shown to directly affect growth rate, body temperature, activity patterns, and thermoregulatory opportunities in lizards (38, 41-45). Wetter areas also have higher thermal inertia (and so lower cyclical thermal fluctuations (46)), and changed environmental variance in temperature potentially has a strong influence on thermal limits (47). Additionally, Bonebrake and Mastrandrea (48) found that changes in precipitation can significantly affect modeled fitness and performance curves. Finally, comparative analyses also suggest that precipitation can influence thermal traits in many species (38). Thus, although the mechanisms linking precipitation to thermal limits are diffuse and poorly resolved, they do exist, and our analyses suggest that precipitation is a strong driver of local adaptation at thermal physiological traits.

Our analysis also suggests that temperature is an important driver of local adaptation in this system, but that extremes of temperature (encapsulated in minimum and maximum temperatures) are at least as strongly associated with local adaptation as is mean temperature. Again, this result is intuitive (natural selection from climate is likely stronger during extreme events than during normal daily temperatures) and agrees with results of empirical studies (38). Finally, the environmental variables with the weakest signals of local adaptation are Tcov (temperature seasonality), TminVar, and TmaxVar (variance of minimum and maximum daily temperatures, respectively). These variables represent predictable environmental variation occurring within an individual’s lifespan and so are variables to which we might expect individuals to
develop plastic responses, rather than fixed differences; local adaptation to these variables would likely be reflected in reaction norms, rather than point values for traits. (49-51).

System-wide signal of local adaptation
The clear negative linear trend displayed in Fig. 3 is precisely what we would expect in a set of trait–environment combinations dominated by local adaptation. Migrant load (the negative effect of the immigration of less-locally adapted individuals) scales positively with immigration as well as with the strength of selection [see equation 5 in Polechová, Barton and Marion (52)]. The reason for this is that, when the strength of selection is moderately high, the environment will have a large effect on relevant traits, and therefore any immigrants coming from differing environments will be particularly maladapted and will therefore have a large and negative impact on the local phenotype. Thus, we expect trait–environment combinations with strong local adaptation to show strong effects of connectivity on the trait–environment relationship (52).

By setting up a statistical model in which the trait–environment relationship is altered by connectivity, we have allowed the possibility that the trait–environment relationship could be reversed as connectivity increases. Such an outcome is absurd from a theoretical perspective. In practice, however, our interaction coefficients were typically estimated to be around 0.36 times as strong as the main effect of environment. In this situation, reversal would only happen when connectivity values were more than 2.7 standard deviations beyond the mean (a situation that is exceedingly rare). Thus, encouragingly, our system wide analysis consistently
provides parameter estimates that are theoretically sensible, despite there being no
constraint within the model for them to be so.

We used long-term climatic averages and found strong evidence that local adaptation
dominates over plasticity in our trait–environment set. If we had included different
environmental variables, such as the conditions the lizards had recently encountered,
signals of plasticity may have been more apparent. Clearly environmental variables
that are similar across generations should lead to local adaptation, while
environmental variables that fluctuate within generations should have a strong
influence on phenotypic plasticity.

Phenotypic plasticity
The importance of accounting for phenotypic plasticity is, however, exemplified in
our dataset by the relatively strong effect of precipitation of the driest quarter (Pdry)
on the temperature at which maximum sprint speed is achieved (Topt) and on
maximum sprint speed (Rmax) itself. On their own, these strong trait–environment
relationships might be interpreted as evidence for local adaptation. Our analysis,
however, suggests that the environmental effect is largely independent of
connectivity, implying that variation in these traits is due to plasticity rather than
genetic differentiation. Other work (27) has shown little temporal variation in Topt
(within generations) despite clear geographic variation and this, together with our
results, suggests that this trait undergoes developmental plasticity, but is fixed in adult
lizards. In principle, this non-effect of connectivity could also arise due to selection
that is so strong that it maintains local adaptation despite high levels of gene flow
[i.e., immigrants are selected against so strongly that they do not contribute to the
recipient population [11]. The trait–environment relationships for Topt and Rmax
are, however, weaker than those for some other traits (e.g., CTmax and CTmin) that
show clear effects of connectivity, so extremely strong selection seems an unlikely
explanation for the pattern we see here.

The generally weak evidence for plasticity in our dataset should not be considered
weak evidence for plasticity in these traits. Indeed many of the physiological traits we
use (e.g., CTmax) are notoriously plastic, responding reversibly on timescales ranging
from hours to months [53, 54]. That we do not see signals of plasticity in these traits
here reflects our choice of environmental variables: long-term climatic variables,
rather than short-term weather variables (such as the temperature in the week before
an animal was collected). We chose these long-term variables precisely because we
are interested in unearthing patterns of local adaptation, rather than patterns due to
rapid, reversible plasticity.

Caveats and challenges

Our intent here has been to point out the additional inference that can be drawn from
data on geographic trait variation if we account for the effect of gene flow on trait
differentiation. The idea that local adaptation is eroded by gene flow offers a novel
way to identify the environmental drivers of local adaptation. Such a capacity is of
fundamental interest, and is also sorely needed if we are to effectively manage the
impacts of climate change. The methods we use here are, however, embryonic, and in
the following we point out caveats and challenges for future work.

Gene flow and connectivity
Our approach requires a measure of gene flow across a landscape. Here we have used environmental connectivity as a proxy for gene flow. We chose connectivity because it can be calculated relatively easily for many species by using broad scale habitat mapping datasets [e.g., vegetation mapping from DERM (55)]. Of course, these measures of connectivity should be calculated at a scale relevant to the scale of dispersal of the species in question [as ours was, using dispersal rate data for *Lampropholis coggeri* from Singhal and Moritz (56)]. While connectivity measures will often correlate with gene flow, e.g., (57)], a measure of gene flow, rather than the flow of individuals, would be preferable. Such measures are increasingly becoming available with the rise of landscape genomics tools (e.g., 58), but may still be cost-prohibitive in many cases.

While there may be better measures of gene flow, our inference might also be improved by taking into account landscape heterogeneity in the environment. Gene flow, per se, does not erode local adaptation. Rather it is an influx of maladapted genes that erodes local adaptation. Thus, a better index of this “migrant load” may well be one in which connectivity is multiplied by a measure of environmental heterogeneity, where connectivity and heterogeneity are calculated over the same spatial scale (e.g., 31). An index such as this should, in principle be a better measure of migrant load than our simple measure of connectivity. The cost, however, is that this index would need to be calculated in a standardized manner for every environmental variable under consideration.

Clearly connectivity is an imperfect measure of migrant load. By using it, we implicitly assume that all migrants are equally maladapted and have equal fitness in
the recipient population. Nonetheless, connectivity should scale positively with
migrant load, and our analysis using simple connectivity generated a coherent and
intuitively sensible result. This is encouraging, suggesting that, in the absence of
precise estimates of migrant load, a readily calculable connectivity metric may suffice
to elucidate broad patterns.

Linear trait–environment relationships, and covariation with connectivity

Our method assumed that traits have a linear relationship to the environment (at least
at the environmental scale across which we are looking). In many instances, this will
be a reasonable null assumption: it seems unlikely, for example, that a trait such as
desiccation resistance would be high in dry environments, low in moderately wet
environments, and then high again in very wet environments. The assumption bears
particular mention, however, in the situation where the connectivity index is strongly
correlated with one or more of the other environmental variables being used. In our
system, for example, AMP and Pdry are correlated with connectivity (Fig. S1). Where
the environment–connectivity correlation is very strong, the interaction term in our
model (Conn×Env) could be interpreted as a quadratic term for environment (i.e.,
Env²). In these cases, it is possible that a strong connectivity interaction is, in fact,
pointing to a non-linear trait–environment relationship. Thus, for environmental
variables that correlate with connectivity (and there will always be some), careful
consideration needs to be given to the possibility of a quadratic fitness function
between trait and environment. In our case, it remains possible, for example, that the
strong influence of precipitation on local adaptation in our system is spurious, and
instead reflects non-linear relationships between optimal trait values and precipitation.
We can, however, think of no obvious reason why thermal limits should respond
quadratically to precipitation, nor why desiccation rates and other physiological traits should also do so. Thus, in our case, we are inclined to accept the importance of this environmental variable in driving local adaptation in our system.

Covariation between explanatory variables

As in any multiple regression analysis, our capacity to make precise coefficient estimates diminishes if there is substantial covariation between our explanatory variables. If a sampling regime is being designed *de novo*, care should be taken to sample sites in such a way that covariation between environmental variables (including connectivity) is avoided as far as possible. Such an aim can be achieved by, for example, strategically exploiting latitudinal and altitudinal gradients.

Multivariate traits and environments

Here we examined one trait–environment combination at a time. Doing so may potentially miss relationships that only appear in multivariate analyses. For example, if two environmental variables are negatively correlated but both have a positive effect on a trait, it is possible that these countergradients can obscure the univariate relationship. Similar problems are encountered when examining response to selection over time (59) and, with our approach, may lead us to underestimate the number of important environmental drivers of local adaptation. Analysis incorporating multiple environmental predictors is possible, but such a model will rapidly become saturated with parameters. To minimize the problem of countergradients, again, care should be taken to sample environmental spaces in such a way as to minimize correlations between environmental variables.
An additional analytical challenge is to treat traits as multivariate. Here we have treated each measured trait as independent. In reality, however, traits covary and this covariance can have both genetic and environmental origins (60). As a corollary, selection acts on the multivariate trait, and causes populations to move in multivariate trait space (61). Consequently, local adaptation perhaps should be measured in a multivariate trait space rather than on a univariate basis. Such an aim, however, requires considerable theoretical development and may well require substantially more data. For now, however, we should be aware that we are collapsing our trait space, and each of our measured traits is not independent. For example, in our system there is a strong correlation between CTmin and CTmax, thus we should be aware that these two traits should not get equal weighting when we use our traits to rank environmental variables by their importance to local adaptation.

**Neutral clines**

Finally, our approach should allow us to identify when geographic variation is a result of genetic variation. That is, it can weed out relationships that are driven by plasticity or habitat choice. Covariation between genotype and environment will often be the result of local adaptation, but can also arise for non-adaptive reasons, the most obvious being trait clines caused by the historical spread of population (62). In principle, and again, with careful attention to sample design (i.e., a sample design which minimizes the covariation between space and environment), it should be possible to separate spatial from environmental patterns.

**Conclusion**
There is increasing urgency to identify populations that will act as suitable sources for targeted gene flow efforts in the face of climate change. To identify these populations, we need to know which traits influence sensitivity to climate and are locally adapted. Traditional approaches to unearthing local adaptation (reciprocal transplants and common garden experiments) are time consuming, and often cannot attribute adaptation to any particular environmental driver. Local adaptation is, however, undermined by gene flow, and we should be able to use this fact to sort patterns of local adaptation from patterns with other causes. Here we have demonstrated this approach: using connectivity as a proxy for gene flow, and looking for its effect on trait–environment relationships. Our analysis, using a species of lizard from Australia’s Wet Tropics rainforest, suggests the approach has merit: the results we achieve are coherent and suggest local adaptation is the overwhelming signal in the set of trait–environment relationships tested. As well as implying a strong role for local adaptation, we have effectively ranked environmental drivers of local adaptation, finding evidence that precipitation and temperature are important environmental variables with regard to local adaptation in our system. Our analysis also suggests that some traits exhibit strong plastic responses to the environment, particularly in response to precipitation of the driest quarter and the seasonality of temperature and precipitation. These specific results will likely apply to other species that are phylogenetically or ecologically similar to our focal species, but the method has the potential to apply much more broadly. Analytical and sampling challenges remain, however, and we point to avenues whereby the method can be improved. Given the potential of this method to provide evidence of local adaptation, and to provide rapid ranking of the climatic drivers of local adaptation, assessment of the method in a broader array of systems is warranted.
Acknowledgments

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The authors are extremely grateful for her patience and attention to detail while undertaking this task.
References


Figure Captions

Figure 1. Graphs showing the concepts illustrated by plotting a set of trait–environment coefficients (e.g., the coefficient from a linear model examining the effect of annual mean site temperature on the sprint performance of organisms from that site: \( B_{\text{env}} \), x-axis) and the corresponding environment–connectivity interaction coefficients (\( B_{\text{int}} \), y-axis). Broad grey line represents the approximate area in which these points would fall. A) Phenotypic plasticity is suggested when trait–environment relationships are strong, but are not influenced by connectivity. B) Local adaptation is suggested when increasing connectivity diminishes the relationship between the environment and the trait. C) The effect of the environment is enhanced by connectivity. This latter pattern might arise in organisms that are highly mobile and can actively move to their ideal environment, thus avoiding the selective pressures that would lead to local adaptation.

Figure 2. Map of the southern Australian Wet Tropics bioregion, showing the distribution of rainforest in green and the sampling locations as black dots.

Figure 3. Scatterplot showing the results of 99 linear models run to assess the relationship between each trait–environment pair, and the environment–connectivity interaction. Trait–environment coefficients (\( B_{\text{env}} \)) are on the x-axis, and environment–connectivity interaction coefficients (\( B_{\text{int}} \)) are on the y-axis. Local adaptation is suggested when these two parameters are opposite in sign: in trait–environment pairs in which a strong environmental effect is eroded by increasing connectivity.

Figure 4. Heatmap showing the relative rankings of climate variables (rows) and morphological and physiological traits (columns). The matrix has been sorted (by reciprocal averaging) and coloured according to the strength of local adaptation, with higher values coloured red and being sorted to the top/left. See Tables S1 and S2 for explanations of the trait and environmental variables used.

\[ L = \text{local adaptation index: } -B_{\text{env}} \times B_{\text{int}} \]
Figures

A) Plasticity

B) Local adaptation

C) Ideal free distribution

Figure 1
Figure 3
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**Figure 4**