Common garden experiments to study local adaptation need to account for population structure

Pierre de Villemereuil\textsuperscript{1}, Oscar E. Gaggiotti\textsuperscript{2}, and Jérôme Goudet\textsuperscript{3}

\textsuperscript{1}Institut de Systématique, Évolution, Biodiversité (ISYEB), École Pratique des Hautes Études / PSL, MNHN, CNRS, Sorbonne Université, Université des Antilles, Paris, France
\textsuperscript{2}Scottish Oceans Institute, University of St Andrews, Fife, KY16 8LB, United Kingdom
\textsuperscript{3}Department of Ecology and Evolution & Swiss Institute of Bioinformatics, University of Lausanne, 1015 Switzerland

Abstract

1. Common garden experiments are valuable to study adaptive phenomenon and adaptive potential, in that they allow to study local adaptation without the confounding effect of phenotypic plasticity. The $Q_{ST} - F_{ST}$ comparison framework, comparing genetic differentiation at the phenotypic and molecular level, is the usual way to test and measure whether local adaptation influences phenotypic divergence between populations.

2. Here, we highlight that the assumptions behind the expected equality $Q_{ST} = F_{ST}$ under neutrality correspond to a very simple model of population genetics. While the equality might, on average, be robust to violation of such assumptions, more complex population structure can generate strong evolutionary noise.

Synthesis We highlight recent methodological developments aimed at overcoming this issue and at providing a more general framework to detect local adaptation, using less restrictive assumptions. We invite empiricists to look into these methods and theorists to continue developing even more general methods.

Keywords: Local adaptation, common garden, $Q_{ST} - F_{ST}$ comparison, population structure, phenotypic divergence, population genetics, quantitative genetics.

Introduction

There are three main possible responses from organisms subjected to climate change (Parmesan \textsuperscript{2006}; Aitken, Yeaman, Holliday, Wang, and Curtis-McLane \textsuperscript{2008}): they can (i) disperse and change their range limits, matching the new geographic repartition of their ecological niche, (ii) quickly acclimate to the new climatic conditions through (possibly transgenerational) phenotypic plasticity or (iii) in the longer run, evolve genetically to match the new climatic conditions through an adaptive process. Predicting the likeliness of these three outcomes (or any combination thereof) is challenging and requires both reliable models and detailed information.

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about the focal species. Probably one of the most difficult tasks is to assess the extent to which phenotypic plasticity or genetic adaptation allow a species to cope with the climatic challenge. This is, in part, because phenotypic plasticity (including epigenetic changes) and adaptive evolution cannot be disentangled from simple observations in the natural environment, and also because they greatly differ in their characteristics to respond to climate change (phenotypic plasticity having a much shorter response time).

Fortunately, “common gardens”, experimental settings in which offspring from different populations are raised under the same (sets of) environmental conditions, allow to circumvent this difficulty and to assess the amount of adaptive genetic variation that exists among populations (Kawecki and Ebert 2004; Savolainen, Lascoux, and Merila 2013; de Villemereuil, Gaggiotti, Mouterde, and Till-Bottraud 2016). A particular challenge in the analysis of such experiment is to distinguish between neutral and adaptive genetic variation among populations. Indeed, even in absence of selection, and as a result of the combined effect of drift, mutation and migration, populations are expected to diverge from a phenotypic point of view, just as they do from a genetic one (Lande 1992). As the fixation index ($F_{ST}$) measures the genetic divergence between populations from a molecular point of view, the phenotypic divergence can be measured through a parameter named $Q_{ST}$ (Spitze 1993). Both can be defined as the ratio of the between-population genetic variance over the total genetic variance (Lande 1992; Spitze 1993):

$$Q_{ST} = \frac{V_B}{V_B + 2V_A},$$

where $V_A$ is the within-population additive genetic variance and $V_B$ is the between population genetic variance.

Moreover, in the absence of selection (i.e. for a purely neutral trait), it is expected that $F_{ST} = Q_{ST}$ (Whitlock 2008). Thus, hypothesis testing based on a null hypothesis of neutrality, as first suggested by Spitze (1993) has generated a lot of literature focused on the $Q_{ST} - F_{ST}$ comparison (reviewed in Leinonen, O’Hara, Cano, and Merilä 2008; Leinonen, McCairns, O’Hara, and Merilä 2013). In these studies, a phenotypic trait with $Q_{ST}$ significantly larger than the $F_{ST}$ estimated from neutral markers is considered as being under local adaptation, while a $Q_{ST}$ significantly smaller than $F_{ST}$ is taken as a sign of balancing selection (Spitze 1993; Leinonen et al. 2013). This framework of $Q_{ST} - F_{ST}$ comparison has been heavily criticised for being subjected to many issues and limitations. For example, $Q_{ST}$ is notoriously difficult to estimate and its (often large) uncertainty should be carefully accounted for in the comparison (O’Hara and Merilä 2005); it can also be influenced by the effect of dominance and inbreeding (Goudet and Büchi 2006; Goudet and Martin 2006; Santure and Wang 2008) and mutation rates might not be comparable between the Quantitative Trait Loci (QTLs) and the neutral markers used to compute $F_{ST}$ (Edelaar, Burraco, and Gomez-Mestre 2011; Edelaar and Björklund 2011). Besides offering criticisms, these studies also provide strategies to overcome some of the limitations, and because such a framework is so crucial to the study of local adaptation, it is still very popular in the literature (Leinonen...
Here, we will focus on an issue that has been been less extensively discussed and, in our experience, not always considered by empiricists using $Q_{ST}$: the influence of population structure on the $Q_{ST} - F_{ST}$ comparison. We explain the assumptions behind the “classical” computation of $Q_{ST}$, show a little example of how population structure can affect the test for local adaptation and explore some alternatives to perform statistical tests excluding the neutral hypothesis of divergence between populations with less restrictive assumptions.

**The typical population model**

$Q_{ST}$ is typically estimated from common garden phenotypic measurements $Y$, in which each individual $i$ belongs to a distinct natural population of origin $p$. The average population effects $a_p$ are considered as being of genetic origin and their variance $V_B$ is assimilated to the between-population genetic variance. While Spitze (1993) used an ANOVA to compute the very first empirical $Q_{ST}$, it is nowadays common (Leinonen et al. 2013) to use a mixed model to compute $V_B$ and the within-population additive genetic variance $V_A$, using:

$$Y_i = \mu + u_i + a_p(i) + e_i,$$

(2)

where $\mu$ is the model intercept, $u_i$ is the individual-level genetic random effect, $a_p(i)$ is the population-level genetic random effect (of the population $p(i)$ the individual $i$ belongs to) and $e_i$ is the residual. The variance associated with the $a_p$ effects is $V_B$ and the variance associated with $u_i$ is $V_A$. The assumptions about the population genetics model lie in how the structure of the random effects are specified. Generally, in order to estimate the $a_p$ effect in the model, one simply uses a “simple” random effect based on the population ID, which results in the assumption that the $a_p$’s are independent and identically distributed. In a more formal way:

$$a_p \sim N(0, IV_B),$$

(3)

where $I$ is the identity matrix and $N$ is the normal distribution. This identity matrix is akin to assuming populations are structured according to an island model, where all equally-sized populations receive and send the same number of migrants from a common pool (see Box 1A). Note that this model is also the one generally used to derive $F_{ST}$ estimates (Weir and Cockerham 1984, but see Gaggiotti and Foll 2010; Karhunen and Ovaskainen 2012; Weir and Goudet 2017 for other approaches). In practice, this model is quite robust to slight deviations from its assumptions, as shown by decades of experience from population geneticists using $F_{ST}$ (Holsinger and Weir 2009).

Theoretically, the equality $F_{ST} = Q_{ST}$ under neutrality should be valid, on average, for population structures.
Box 1: Comparing island and stepping-stone models

To illustrate the effect of population structure on the \( F_{ST} - Q_{ST} \) contrast, we compare the results of simulations of the island model (A), where all populations exchange the same number of migrants with all others, and the one dimensional stepping stone model (B), where each population exchanges migrants with its nearest neighbours only. The data were generated with the ms program (Hudson 2002), assuming a genome of 20 chromosomes each 100'000 base pair long. Migration was adjusted to obtain an overall \( F_{ST} \) around 0.2 for the two population structures. In both sets of simulations, the number of populations is 20, and 500 individuals per population are sampled.

Figure C shows the results. For each scenario (IslM: Island model, SS1D: Stepping-Stone), 500 neutral, purely additive and genetically determined traits are simulated with either 10, 100 or 1000 (top to bottom row) causal loci and effect sizes drawn from a normal distribution. The red horizontal line on each panel is the genomic \( F_{ST} \), the violin plots labelled \( F'_{ST} \) correspond to \( F_{ST} \) estimated from causal loci, and the violin plots labelled \( Q_{ST} \) show the distribution of \( Q_{ST} \) for each scenario. Dots are the observations. The blue points show the mean value of the observations. For both models, \( F'_{ST} \) is essentially unbiased, with more variation when the number of loci encoding the trait is small, as expected. For \( Q_{ST} \), while more variable that \( F'_{ST} \), the estimation for the island model are centered around their expected value, with an empirical 95% percentile interval [0.1, 0.27]. For the one-dimensional stepping stone model, despite the mean being close to its expected value, a majority of \( Q_{ST} \) estimates are smaller than the expected value, with a mode around 0.1 and the range of variation is extremely wide, from 0 to 0.7, and an empirical 95% percentile interval [0.05, 0.44]. Such a variation due to very large evolutionary stochasticity for \( Q_{ST} \) (note that statistical sampling errors have been minimized here, since \( V_A \) is the phenotypic variance and \( V_B \) is estimated from 20 populations, so using the statistical framework in Equation 2 would most likely aggravate the issue) makes carrying a test for the null hypothesis \( Q_{ST} = F_{ST} \) prone to a high level of type I errors, if not accounting for such effect of population structure. Indeed, for the island model, 5% of the simulated \( Q_{ST} \) are significant at the 5% nominal level when using the modified Lewontin-Krakauer test proposed by Whitlock and Guillaume (2009), while 30% are significant for the one dimensional stepping-stone model.
other than the island model (Whitlock 1999). We show in Box 1 this equality to hold for the island model and, on average indeed, for the one-dimensional stepping-stone model. However, for this last model, a majority of traits gave $Q_{ST}$ values (much) lower than $F_{ST}$, and some gave $Q_{ST}$ values much higher than $F_{ST}$ (Box 1C). We argue that the test $Q_{ST} = F_{ST}$ is not reliable in general because of the much larger evolutionary stochasticity associated with $Q_{ST}$, which inflates type I errors. Hence implementation of the test should take into account such evolutionary variance. This is particularly important when the population structure deviates from a simple island model because the increase in evolutionary stochasticity is much more pronounced for $Q_{ST}$ than for $F_{ST}$.

**Toward a solution: more general strategies**

It is difficult to come up with a universal model of neutral evolution, that can account for any sort of population structure. However, there have been some methodological developments that aim at testing local adaptation in a common garden setting with more general assumptions about population structure. Here, we wish to highlight two of them.

Ovaskainen, Karhunen, Zheng, Arias, and Merilä (2011) method is not based on a direct $Q_{ST} - F_{ST}$ comparison. Instead, it is based on a theoretically motivated neutral model of phenotypic divergence that allows for differences in migration and drift among populations, as well as preferential migration between populations. To achieve this, the identity matrix $I$ is replaced by a between-population relatedness matrix (here noted $B$) in Equation 3:

$$a_p \sim N(0, BV_B),$$

(4)

To estimate this matrix from neutral marker data the same authors propose an extension of the F-model (Gaggiotti and Foll 2010) that also allows the simultaneous estimation of the other parameters associated with Equation 2. Because $B$ is a matrix, and not just a single number as is $F_{ST}$, this framework offers both a more accurate description of the population structure and more power to detect deviation from neutrality. It does so by alleviating the issues affecting the direct comparison of $Q_{ST}$ with $F_{ST}$ and provides a statistical test (coined “S-test”) measuring the deviation of the population means themselves from the neutral expectation. A more recent implementation of the method in R (driftsel, Karhunen, Merilä, Leinonen, Cano, and Ovaskainen 2013) first estimates the matrix using neutral markers and an admixture F-model (Karhunen and Ovaskainen 2012), and then incorporates quantitative trait data to estimate all remaining parameters and further refine the estimate of the matrix. Finally, a new alternative method to estimate the $B$ matrix (which can then be used by driftsel) is provided by the unified approach to characterise population structure and individual relatedness and inbreeding recently put forward by Weir and Goudet (2017). Using this model of neutral evolution ap-
Applicable to both molecular and quantitative trait data also allows for the further addition of environmental information, and as a result, perform more powerful statistical tests using habitat information (Karhunen, Ovaskainen, Herczeg, and Merilä 2014) or single environmental values (de Villemereuil, Mouterde, Gaggiotti, and Till-Bottraud 2018), while accounting for the (potentially confounding) effect of genetic drift and population structure.

Sometimes, the population structure is so complex that even defining populations can be very difficult and, in the end, a fairly subjective process (Waples and Gaggiotti 2006). Martins, Caye, Luu, Blum, and François (2016) have shown that, in such cases, an equivalent to $F_{ST}$ can be derived even in absence of delimited populations, using either the genomic proportion of an individual assigned to a given cluster (the $Q$ matrix of the Structure program) or the individual scores along the different axes obtained from a Principal Component Analysis of the genotypes. This later approach was used by Josephs, Berg, Ross-Ibarra, and Coop (2019) to define an equivalent to $Q_{ST}$ (coined $Q_X$) in absence of explicitly defined populations. However, to circumvent the need for a direct comparison to $F_{ST}$, Josephs et al. (2019) suggest testing for a phenotypic excess of variance ($F$-test) along some of the Principal Component axes retained for the analysis. The difficulty in the analysis lies in identifying the set of first axes defined to be “among” populations (the other lower axes being considered “within”). Josephs et al. (2019) offer various ways of defining this limit between among and within populations, ranging from using an arbitrary threshold to using the Tracy-Widom test.

**Conclusion**

Since it was first proposed by Spitze (1993), the $Q_{ST} - F_{ST}$ comparison framework has been an invaluable tool to investigate the prevalence and characteristics of local adaptation. Nonetheless, we encourage empiricists to consider the alternative strategies discussed here to better account for various effects of population structure.

As these new strategies themselves have their own limitations, we also urge theorists to continue developing new methods to study the phenotypic impact of local adaptation in common garden, while accounting for population structure and evolution stochasticity as accurately as possible.

**Authors contribution**

PdV led the writing of the manuscript with critical contributions from OEG and JG. JG performed the simulation analysis, with analysis from all authors. All authors agreed to the publication.
Data Availability

We provide the code for replicating the analysis in Supplementary Information.

References


