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INVITED REVIEWS AND SYNTHESES

A practical guide to environmental association analysis in landscape genomics

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Abstract

Landscape genomics is an emerging research field that aims to identify the environmental factors that shape adaptive genetic variation and the gene variants that drive local adaptation. Its development has been facilitated by next-generation sequencing, which allows for screening thousands to millions of single nucleotide polymorphisms in many individuals and populations at reasonable costs. In parallel, data sets describing environmental factors have greatly improved and increasingly become publicly accessible. Accordingly, numerous analytical methods for environmental association studies have been developed. Environmental association analysis identifies genetic variants associated with particular environmental factors and has the potential to uncover adaptive patterns that are not discovered by traditional tests for the detection of outlier loci based on population genetic differentiation. We review methods for conducting environmental association analysis including categorical tests, logistic regressions, matrix correlations, general linear models and mixed effects models. We discuss the advantages and disadvantages of different approaches, provide a list of dedicated software packages and their specific properties, and stress the importance of incorporating neutral genetic structure in the analysis. We also touch on additional important aspects such as sampling design, environmental data preparation, pooled and reducedrepresentation sequencing, candidate-gene approaches, linearity of allele-environment associations and the combination of environmental association analyses with traditional outlier detection tests. We conclude by summarizing expected future directions in the field, such as the extension of statistical approaches, environmental association analysis for ecological gene annotation, and the need for replication and post hoc validation studies.

Keywords: adaptive genetic variation, ecological association, environmental correlation analysis, genetic–environment association, genotype–environment correlation, local adaptation, natural selection, neutral genetic structure, population genomics

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The emergence of landscape genomics

Changing environmental conditions force organisms to be phenotypically plastic, migrate or adapt to avoid extinction. Local adaptation (Williams 1966; Kawecki &

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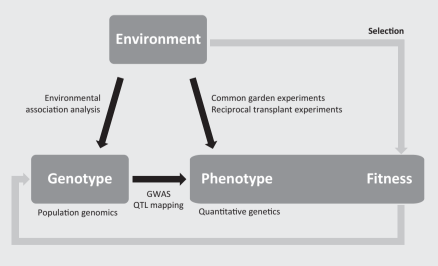
Ebert 2004; Savolainen *et al.* 2013) is the response to differential selective pressures among populations and habitats, acting on genetically controlled fitness differences among individuals. Hence, genes underlying heritable phenotypic variation are of great interest in evolution and ecology. To identify such genes, two types of approaches are currently used (Barrett & Hoekstra 2011). Top-down approaches, such as genomewide

association studies (GWAS, reviewed in Korte & Farlow 2013) and quantitative trait locus (QTL) mapping (reviewed in Stinchcombe & Hoekstra 2008), take advantage of phenotypic measurements and relate them to genotypic data (Box 1). Bottom-up approaches, such as population and landscape genomics, use genomic information to identify signatures of adaptive genetic variation and relate them to evolutionary processes and environmental variation. In population genomics, approaches based on identifying regions of high genetic differentiation among populations as compared to a neutral model are commonly used to detect positive selection (Luikart et al. 2003). Although they are frequently used, these outlier tests have drawbacks. First, in the case of positive selection, they are aimed at identifying loci that underwent selective sweeps of beneficial alleles. Adaptation to local conditions, however, can lead to subtle changes in allele frequencies that are hardly detected by outlier tests, for example in the case of polygenic additive effects (Pritchard & Di Rienzo 2010) or under high geneflow counteracting patterns of local adaptation (Kawecki & Ebert 2004). Second, outlier tests make the assumption that selection pressures differ among populations, but usually do not attempt to link to specific selection pressures that underlie adaptation. An approach that successfully integrates the environment, which is a major driving force behind natural selection, thus represents a valuable alternative to detect adaptive loci.

Some of the earliest examples of adaptation in natural populations come from observed concordances between phenotypic traits and environmental variation. Turesson (1922) was one of the first to consider the genotype as the relevant unit living in different habitats across the distribution of a species. Huxley (1938) reviewed several case studies of intraspecific variation in phenotypes across space. He coined the terms 'cline' to describe this phenomenon and 'ecocline' to describe the case where phenotypic variation is correlated with ecological factors. In recent years, with increasing

Box 1. Detecting signs of natural selection and genes involved in local adaptation

In the context of environmental, genetic, phenotypic and fitness variation, several approaches exist to uncover signs of natural selection and detect genes and environmental factors involved in local adaptation. The following simplified scheme presents some of these possibilities (modified from Sork *et al.* 2013). Boxes mark sources of variation that can be quantified, black arrows indicate the direction of the evolutionary process between cause and effect, and the grey arrow shows how selection acts on the different levels. Population genomics (reviewed in Hohenlohe *et al.* 2010b) and quantitative genetics (Stinchcombe & Hoekstra 2008) use genotypic and phenotypic information, respectively, alone to identify adaptive genetic variation. All other methods deal with the interaction of two of the different types of data. QTL (quantitative trait locus) mapping (Stinchcombe & Hoekstra 2008) and GWAS (genomewide association studies, Korte & Farlow 2013) are used to identify loci linked to specific phenotypes. Common garden and reciprocal transplant experiments (Savolainen *et al.* 2013) investigate the phenotypic and fitness differences of individuals originating from and living in different environments. Environmental association analysis (reviewed in this study) aims to correlate environment and genotypes. To our knowledge, only one methodological framework (Berg & Coop 2014) performs a joint analysis of all three aspects.



availability of genetic data from diverse species, a popular approach seeks to identify genetic variants strongly associated with specific environmental conditions (see Mitton et al. 1977; for one of the earliest examples). This approach, referred to as environmental association analysis (EAA; Boxes 1 and 2) and also called genetic-environment analysis (e.g. Lotterhos & Whitlock 2015), has the potential to uncover patterns induced by adaptive processes that are not detected by traditional population genomic approaches, or to complement and support results of these. EAA is at the core of landscape genomics, an emerging research field that integrates tools from landscape genetics and population genomics to identify the environmental factors that have shaped present-day (adaptive) genetic variation and the gene variants that drive local adaptation (Holderegger et al. 2010; Manel et al. 2010a; Manel & Holderegger 2013; Sork et al. 2013). In practice, EAA is often used in concert with other population genomic tools such as outlier analysis (e.g. Fischer et al. 2013). It is thus difficult to draw a distinct line between these two approaches. As with many other areas of molecular ecology, the emergence of landscape genomics has been strongly facilitated by next-generation sequencing (NGS), which allows screening thousands to millions of single nucleotide polymorphisms (SNPs) across the entire genomes of many individuals and populations at reasonable costs. The data sets describing environmental characteristics (e.g. spatially explicit data on abiotic factors such as topography, climate, bedrock type, but also biotic factors such as dominant species or vegetation types) have also greatly improved and increasingly become publicly accessible, owing to versatile remote sensing techniques and database harmonization, respectively.

Numerous statistical methods for environmental association studies have recently been developed. However, no single widely accepted statistical approach has yet emerged. Accordingly, researchers often find it difficult to navigate the many possible avenues for EAA provided by recent innovation. Here, we present a practical guide to EAA, both for the landscape genomics community as well as for those freshly entering this research field. This article complements earlier conceptual reviews on landscape genomics (Holderegger et al. 2010; Manel et al. 2010a; Schoville et al. 2012; Joost et al. 2013; Manel & Holderegger 2013; Bragg et al. in press) and comparisons of the statistical performance of selected methods (De Mita et al. 2013; Frichot et al. 2013; Jones et al. 2013; de Villemereuil et al. 2014; Lotterhos & Whitlock 2015) by focusing on the practical aspects of designing and analysing an environmental association study. First, we will introduce the basics of EAA by describing sampling designs and required data sets. Next, we present several

methods, focusing on their optimal application, also referring to dedicated software packages and their specific properties. Subsequently, we touch on limitations and extensions of EAA and conclude by describing future directions and possible improvements in the field of landscape genomics. This review concentrates on SNPs as genetic markers, because they are currently the marker of choice and because they can often be functionally annotated. However, several environmental association methods can also be used with other, less commonly used marker types such as expressed sequence tagderived simple sequence repeats (EST-SSR, e.g. Bradbury et al. 2013) or anonymous and dominant markers, such as amplified fragment length polymorphisms (AFLPs, e.g. Manel et al. 2012b).

Preparation of data

The basic goal of EAA is to test whether particular alleles or gene variants are significantly associated with any factor describing the environment in which they predominantly occur. For an environmental association study, two types of data are gathered, namely environmental factors and genetic polymorphisms, which should match in spatial resolution when establishing the sampling design. Processing environmental data includes data compilation (on-site measurement, data acquisition from existing sources), quality control, integration over time and/or space, and factor selection. Assessing genetic polymorphisms requires DNA extraction and sequencing or genotyping and is followed by bioinformatics, including quality control and data trimming. The two data components are then used in the actual EAA to assess evidence for allele-environment correlations. These steps are shown in Fig. 1 and detailed in the following sections.

Sampling design

When identifying sampling locations for an environmental association study, one intuitively thinks about sampling along environmental gradients. For instance, one could sample along a continental temperature or a local water salinity gradient. This design is appealing, but replication of gradients, also within evolutionary lineages, is important because multiple findings of the same candidate loci are a strong sign that they are true positives, and because replication reduces the confounding of population structure and covarying environmental factors. Usually, gradients of one particular environmental factor are the focus, but other environmental factors can be integrated into the analysis later. Another possibility is sampling in a categorical way, where researchers set up a 'quasi-experimental' design

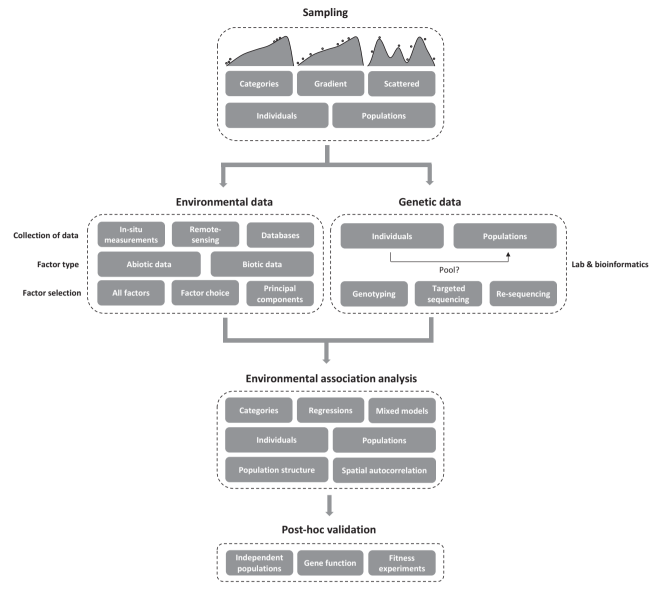


Fig. 1 A typical workflow in environmental association analysis (EAA). The three most important options per step are horizontally aligned. Genetic and environmental data are collected at the same sampling locations, processed separately and jointly analysed in EAA. The results can subsequently be validated with complementary approaches. All steps and options are described in detail in the manuscript.

with different 'treatments', for example low vs. high temperature or low vs. high salinity. Here, levels of a particular environmental factor are in focus. Categorical sampling seems attractive, but the range of subsequent adequate EAAs is limited, and one should clearly consider the number of replicates necessary for statistical significance testing. Researchers can also get a broad sample covering the entire environmental niche of a study species in a given area. Sampling locations would then be more widespread and scattered, or even randomly stratified (weighted random samples of representative subsets of sampling locations, Allaby 2009).

This scattered sampling design leaves a lot of flexibility as a variety of different environmental association methods and environmental factors can potentially be used (Table 1), but it comes with the drawback of (often) lacking replication and clear hypotheses to test. In a review on sampling strategies in landscape genomics, Manel *et al.* (2012a) suggest to use model-based stratification and simulations to establish sampling designs (if sufficient biological and environmental knowledge is available) instead of applying classical ecological sampling designs like random sampling. The authors recommend choosing the climatic or biological

 Table 1
 Overview of methods and software available for environmental association analysis in landscape genomics. Note that for some methods, other software or R packages are available

Method	Reference	Association type	Sampling design	Incorporation of neutral genetic structure	Incorporation of spatial autocorrelation	Individual/ population data	Mode for pooled data	Correction for sample size	Software/ R package
Categories		Categorical	Categorical	Possible	Possible	Both	Possible	Possible	Various statistical methods
Spatial analysis method (SAM)	Joost et al. (2007)	Logistic	Gradient/ scattered	Possible (in SAMBADA)	Possible (in SaMβada)	Individual	No	No	sam (Joost et al. 2008), samβada (Stucki et al. submitted)
Multiple logistic regression		Logistic	Gradient/ scattered	Possible	Possible	Individual	No	No	R (R Development Core Team 2011)
Generalized estimating equations (GEEs)	Carl & Kuhn (2007), Poncet et al. (2010)	Logistic	Gradient/ scattered	No	Yes	Individual	S O	No O	GEEPACK (Yan & Fine 2004)
Partial Mantel test	Smouse et al. (1986)	Linear/ rank- linear	Gradient/ scattered	Yes	Possible	Both	No	Š	& Urban 2007), VEGAN (Oksanen et al. 2013)
Multiple linear regression/ General linear models		Linear	Gradient/ scattered	Possible	Possible	Both	No.	No	R (R Development Core Team 2011), TASSEL (Bradbury et al. 2007)
Canonical correlation analysis (CCA)	Legendre & Legendre (2012)	Linear	Gradient/ scattered	Possible	Possible	Both	N o	No	VEGAN (Oksanen et al. 2013)
(Partial) redundancy analysis (RDA)	Legendre & Legendre (2012)	Linear	Gradient/ scattered	Possible	Possible	Both	No	No	VEGAN (Oksanen et al. 2013)

Table 1 Continued

(Bradbury et al. GINLAND (Guillot et al. 2013), LEA Villemereuil & BAYENV (Coop et al. 2010), Gaggiotti in press) SAYESCENV (de FMM (Frichot press)
EMMA (Kang
et al. 2008), (Günther & Coop 2013) Francois in 2007), LME4 (Bates et al. et al. 2014) (Frichot & Software/ R package BAYENV2 2014) TASSEL sample size Correction Yes Yes for Yes Νo Š Yes (in BAYENV2) Mode for pooled data Ν̈́ οŃ Š ρ Individual/ population Population Individual Both Both data Both autocorrelation Incorporation of spatial Yes Š Š Š Š Incorporation of neutral genetic structure Yes Yes Yes Yes Yes Gradient/ scattered Gradient/ scattered Gradient/ Sampling Gradient/ scattered Gradient/ scattered scattered design Differentiation-Association Linear/ linear based rank-Linear Linear Linear de Villemereuil & Gaggiotti Guillot *et al.* (2014) Frichot et al. Coop *et al.* (2010) (in press) Reference (2013)generalized Latent factor (SGLMM) models (LFMMs) methods F_{ST}-based models mixed model Spatial mixed Method linear mixed GWAS BAYENV

space over topographic or geographic space when developing a stratified sampling design. Finally, an interesting approach suggested by Lotterhos & Whitlock (2015) is to sample scattered and random pairs of closely situated populations that exhibit substantial differences in environmental conditions while being within geneflow distance. These authors showed, using simulated data, that this sampling design has increased power in detecting true positives compared to random or transect designs, especially in models with weak selection. The reason for this is that the paired design maximizes the differences in adaptive environment while it minimizes the differences in neutral genetic structure. Importantly, landscape genomic studies should be performed over an appropriate geographic scale, which depends on the ecology of the organism (reviewed in, e.g. Anderson et al. 2010; Manel et al. 2010a; Richardson et al. 2014). A major issue is the mobility, dispersal capacity and migration rate of the species under study: for example, the relevant scale for mobile animals may be quite different to the scale for stationary plants. Moreover, researchers should be aware of potential mismatches in time between genomic and environmental data; there might be a time lag between the process causing the genetic pattern and the observed genetic response to it (Anderson et al. 2010).

Sampling can either be performed on the individual or population level. In studies that include only a single individual per sampling location, laboratory costs (but not costs for field sampling) are decreased, as only a comparatively low number of individuals has to be processed. Individual sampling limits the range of EAAs to

individual-based approaches that can handle allele or locus genotype presences/absences or allele frequencies of 0/0.5/1 in the case of SNPs in a diploid species (Table 1 and Box 2, Figs C,E). In contrast, studies using population-based sampling can take advantage of population-based association approaches (Box 2, Figs A,B,D).

Environmental factors

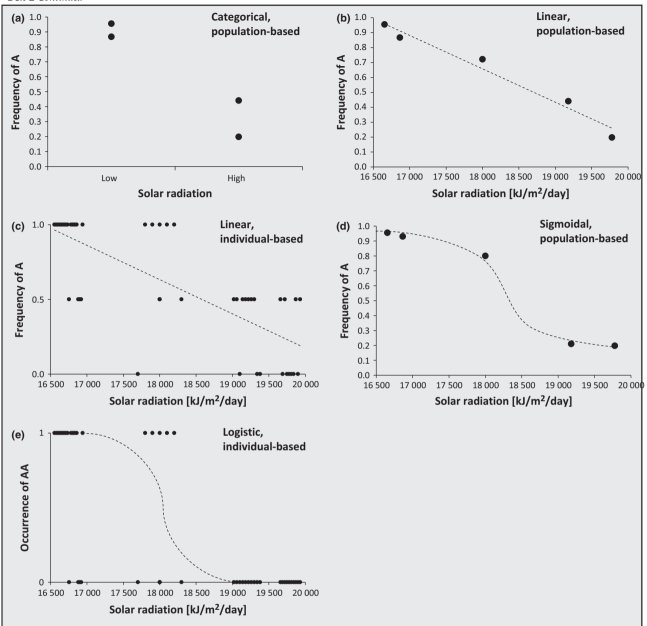
Sources of environmental information. As many abiotic and biotic factors are potentially acting as selective pressures, it is crucial to consider those factors that are most pertinent for the question asked and most likely to provide high explanatory power. Because this knowledge is usually missing a priori, environmental association studies are often rather explorative.

Abiotic data, in particular topo-climatic factors interpolated over large areas, are available from many public databases (Manel et al. 2010a; Thomassen et al. 2010). Limited to about 1-km² resolution is the ground-based interpolated WorldClim data (http://www.worldclim. org, Hijmans et al. 2005), where global climate layers for numerous factors, for recent as well as past and future periods, are freely available for analyses in a geographic information system (GIS) or in R (R Development Core Team 2011). Regional data sets based on ground-measured climate records, with higher resolution than the WorldClim data, are often available. Such climate data provide annual, seasonal, monthly or daily mean values as well as ranges and extremes. Increasingly, remote sensing supports data acquisition for large-scale environmental data, including elevation (http://glcf.umd.

Box 2. Response curves

The main goal of environmental association analysis (EAA) is to test whether a specific allele or locus genotype is associated with a specific environmental factor, while controlling for neutral genetic structure. However, depending on the genetic data available and the sampling design, different possibilities exist to detect such associations. Moreover, different response curves might be expected. This is illustrated by the following simplified examples. Imagine an adaptive SNP (locus X) with alleles A and G of a plant gene involved in response to light stimulus. In the simplest example, we sample four populations, two in each habitat with either low or high radiation. We genotype the locus in all individuals and calculate allele frequencies per population. If allele A at locus X is associated with low radiation, then we expect it to mainly occur in low-radiation populations, whereas allele G is mainly found in individuals of high-radiation populations (Fig. A). In the second case, we sampled five populations along a gradient in solar radiation. Here, an association (using linear regression) would look, for example, like in Fig. B, where the frequency of A in a population decreases when radiation increases. In a sampling design that includes scattered and geo-referenced single individuals from habitats with large differences in radiation, a significant association (using linear regression) should resemble Fig. C. Here, only three levels of allele frequencies (AA = 1, AG = 0.5 and GG = 0) are possible. If both alleles of locus X are mostly fixed for either high or low radiation, and intermediate frequencies are rare, we could expect a sigmoidal response of allele frequencies to the environmental gradient (Fig. D). Finally, in a logistic approach, one tests the association of the presence/absence of an allele or locus genotype, as, for example AA shown in Fig. E. Note that these simplified examples do not incorporate neutral genetic structure, which most of the more sophisticated environmental association methods do.

Box 2 Continued



In essence, most EAA methods assume linear responses. Nevertheless, in the case of a selective sweep caused by strong directional selection (Barrett & Schluter 2008) leading to the fixation of alleles in two extreme environments, one might rather expect a sigmoidal (Fig. D) than a linear response curve (Fig. B) of allele frequencies along the environmental gradient, because the two alleles lead to clear-cut fitness differences in response to the environment (Richter-Boix *et al.* 2011). None of the currently used methods for EAA is able to deal with such sweep-like response curves of population allele frequencies. Logistic regression best meets the assumptions underlying a sigmoidal pattern, but has other drawbacks (see Future directions). However, Prunier *et al.* (2012) conceptualized adaptation along an environmental gradient in the following way. Each allele essentially displays a bell-shaped frequency curve on a part of the whole environmental gradient. In other words, it provides strong adaptation to and thus highest fitness at a particular position of the gradient. The additive effects of the bell-shaped frequency curves of all alleles together enable the species to live along the whole environmental gradient. If such additive effects are occurring, we expect a linear rather than a sigmoidal response curve between allele frequencies and environmental factors.

edu), precipitation (http://trmm.gsfc.nasa.gov/data_ dir/data.html) and vegetation indices (e.g. http:// glcf.umd.edu, https://lpdaac.usgs.gov). The latter have been further developed through the use of light detection and ranging (LiDAR) technology, but such data are only available at regional (mostly national) scale so far. The primary limitations of current climate data sources are that they (i) often have a coarse spatial resolution, (ii) are integrated over a certain time period, (iii) represent spatial and temporal interpolations, and, consequently, (iv) ignore small-scale heterogeneity. Microscale conditions can therefore not be characterized in sufficient detail. Hence, researchers have started measuring data on site, for example by assessing soil conditions or using remote-sensing techniques (e.g. unmanned aerial vehicles, UAVs), but published examples are not available so far. While field-based measurements well reflect local site conditions in given years, they can fail in capturing long-term environmental conditions, which may often underlie adaptive response. An elegant, but restrictive way to make use of on-site measurements in EAA is to choose sampling locations where data recording has been performed over long periods.

For topographic data such as altitude, slope and aspect, detailed digital elevation or terrain models (DEMs/DTMs) are accessible at a worldwide scale and often at very high resolution (e.g. ASTER, http://asterweb.jpl.nasa.gov/gdem.asp). In this respect, new techniques, such as satellite- or plane-based LiDAR data assessment or UAVs help to improve the spatial resolution of DEMs to a few centimetres. From such high-resolution DEMs, microsite conditions may also be derived (Leempoel *et al.* in press). Furthermore, a wealth of other environmental data can possibly be considered, including geological factors, vegetation types, land cover, land use or species distributions, which might also serve as proxies for trophic interactions, prey availability or pathogen pressure (Gugerli *et al.* 2013).

Preparing environmental factors. A strategy that considers all environmental factors one-by-one provides a comprehensive analysis and reduces the risk of missing important loci and genes involved in local adaptation. However, using a large number of environmental factors rather than specific hypothesis tests increases the number of statistical tests, which needs to be considered in analyses to reduce the rate of false positives. In addition, many biotic and abiotic factors are highly correlated, for example altitude and temperature, or latitude and growing period. This leads to the testing of strongly interdependent models, increases variance in multivariate tests and makes estimates of the relative importance of different factors difficult. Including

highly correlated factors may thus lead to the wrong conclusions if an understanding of the environmental drivers of local adaptation is desired.

One way to avoid collinearity is to reduce the number of factors (for a review of methods, see, e.g. Dormann et al. 2013). For example, Fischer et al. (2013) removed highly correlated factors (Pearson's $|r| \ge 0.8$) based on a pairwise correlation matrix. Another possibility is to select or remove highly correlated factors based on their contribution to the first few axes of a principal component analysis (PCA), keeping only those factors with the highest contribution to each axis (e.g. Manel et al. 2010b; Zulliger et al. 2013). A further option is to control for multicollinearity with the variance inflation factor, by iteratively removing the most highly correlated factor until the highest factor is lower than a certain threshold. Nevertheless, when reducing the number of factors, the removed factors should still be taken into consideration when interpreting the results. Imagine a sampling design with several alpine plant populations at high altitude. Solar radiation is removed in the process of factor reduction, because it shows a positive correlation with temperature (which is often the case at high altitude; Körner & Riedl 2012). If a gene known to be involved in response to radiation is associated with temperature, one might easily misclassify the selective pressure.

PCA (or related multivariate statistics) offers another possibility to condense a large number of environmental factors. This approach creates new synthetic environmental factors, consisting of groups of variables (e.g. Eckert *et al.* 2010a; Mosca *et al.* 2012; Nosil *et al.* 2012). While this simplification to a few derived factors makes statistical analysis easy, it can make the biological interpretation of the results difficult, notably if several factors strongly influence principal components. It is therefore only recommended to use PCA loadings as environmental factors when their interpretation is straightforward. PCA may also be problematic if the data show high levels of spatial autocorrelation (Thomassen *et al.* 2010).

Genomic data

Historically, after the use of isozymes (e.g. Mitton et al. 1977), dominant AFLPs were the marker of choice for EAA (Manel et al. 2010a), because they allowed testing up to hundreds of loci with a relatively simple and inexpensive laboratory protocol. The sequence of an AFLP marker and its flanking region, however, is commonly unknown unless additional sequencing efforts are made (e.g. Buehler et al. 2013; Zulliger et al. 2013). These anonymous markers have largely been replaced by SNPs, which are abundant across the entire genome, can easily be standardized

among laboratories, and whose flanking sequences can be directly queried in public databases (Morin et al. 2004). In the coming years, whole-genome sequencing of all individuals will eventually become the standard in EAA and enable the association of millions of SNPs of known location and function. So far, we are aware of only one published environmental association study (Yoder et al. 2014) that used individually sequenced whole genomes. If such deep sequencing is not possible due to large sample sizes and genomes, researchers aiming for environmental association studies can reduce costs mainly by sequencing pooled samples (Pool-Seq) or by targeting a fraction of the genome (e.g. candidate-gene approach or genome complexity reduction). We detail these three options in the following sections.

Pool-Seq (reviewed in Schlötterer et al. 2014) is a cost-effective method of NGS, because the DNAs of several individuals are equimolarly pooled before sequencing (Futschik & Schlötterer 2010). This approach can lead to accurate SNP allele frequency estimates (reviewed in Rellstab et al. 2013) and population genomic parameters (Futschik & Schlötterer 2010; Schlötterer et al. 2014). As a drawback, individual multilocus genotypes and information on heterozygosity are inaccessible. As many environmental association approaches can handle population allele frequencies (Table 1), the use of whole-genome Pool-Seq is an attractive option, but only BAYENV2 (Günther & Coop 2013) yet accounts for the variance introduced by variation in sequencing coverage in Pool-Seq. Nevertheless, whole-genome Pool-Seq data have only rarely been used in EAA so far (but see Turner et al. 2010; Fabian et al. 2012; Fischer et al. 2013).

In a candidate-gene approach, genes or loci are characterized which have already been identified or known to potentially play an important role in local adaptation, or which are involved in a biological process related to the tested environmental factors. This is an especially appealing strategy for study species for which only limited genomic information is available. Information about biological processes can be retrieved, for example from homologous genes of species for which gene ontology (GO) databases exist (Primmer et al. 2013). For SNP genotyping, there are various high-throughput methods on the market (e.g. real-time PCR, KASP, Infinium, GoldenGate, pyrosequencing). Some genotyping technologies can also be used to accurately determine allele frequencies of population pools, for example Infinium (e.g. Bourret et al. 2013) or pyrosequencing (e.g. Gruber et al. 2002; Rellstab et al. 2011). To sequence genes or gene regions, targeted amplicon sequencing of individual or pooled samples using one of the NGS platforms is an attractive option (e.g. Homolka et al. 2012; Ho et al. 2014).

An alternative strategy to lower costs is complexity reduction of the genome. In exome capture (Bamshad et al. 2011), only the part of the genome is sequenced which hybridizes to probes covering exons. This approach requires at least partial knowledge about the transcriptome. In restriction-site associated DNA sequencing (RAD-Seq) and its variants (Puritz et al. 2014), the complexity of the genome is reduced using restriction enzymes, and the flanking regions of restriction sites are sequenced by NGS (Davey et al. 2011). This approach has successfully been applied to pooled population samples (Emerson et al. 2010). However, RAD-Seq identifies substantially fewer polymorphisms, from a few thousand to tens of thousands (e.g. Emerson et al. 2010; Hohenlohe et al. 2010a), as compared to millions of SNPs when using whole-genome Pool-Seq (e.g. Turner et al. 2010; Fabian et al. 2012; Fischer et al. 2013).

Incorporating neutral genetic structure

EAAs need to consider various types of autocorrelation, which arise from the mere historical relationships of individuals across the sites where they live. Consider two locations, where several individuals are sampled. The samples from the same location share a similar environment, which in turn is likely to differ from the other location. Likewise, individuals from one location tend to be more closely related to each other than to individuals from the second location. This concept can be expanded to any spatial scale and applies to both individual- and population-based sampling. If EAAs do not consider such dependencies, the identified associations might just be the consequence of the spatial arrangement and demographic history of the individuals or populations, and not a signature of local adaptation. It is therefore important to correct for neutral genetic structure in EAA. Alternatively (or additionally), some studies and methods (Table 1) include pure spatial autocorrelation in their approaches. Because spatial autocorrelation can serve as a proxy for neutral genetic structure, given isolation-by-distance patterns, a joint incorporation of both parameters (genetic and spatial structure) in such a situation is actually overly conservative. As spatio-environmental relationships are well covered in a recent review (Thomassen et al. 2010), we touch this issue only briefly and focus on how to deal with neutral genetic struc-

To account for the spatial signal in the data, one may just incorporate one or more spatial factors in regressionbased models. A simple approach integrates either the

geographic coordinates of, or the pairwise Euclidean distances between sampling locations into analysis (e.g. Guillot et al. 2014; Stucki et al. submitted). In a more elaborate strategy, Manel et al. (2010b) included Moran's eigenvector maps (MEMs, based on coordinates of the sampling locations, Borcard & Legendre 2002; Dray et al. 2006). MEMs represent environmental variation not specifically included in the model as well as pure spatial signals. Using generalized estimating equations (GEEs), Poncet et al. (2010) considered spatial autocorrelation of sampled individuals within populations. This concept assumes that individuals sampled within the same location share respective properties (habitat, kinship), whereas individuals sampled at any other site do not.

Neutral population genetic structure is defined as allele frequency differences among populations that have arisen due to neutral processes such as genetic drift, gene flow and mutation. The patterns of differences in allele frequencies among populations are the background against which loci contributing to local adaptation — a non-neutral process — are assessed in EAA. Neutral processes affect all loci across a genome, whereas non-neutral processes affect only a subset of loci. Corrections for neutral genetic structure are important in EAA, because neutral genetic structure can mimic patterns expected under non-neutral processes (Excoffier & Ray 2008; Excoffier et al. 2009; but see Vilhjalmsson & Nordborg 2013). For example, post-Pleistocene expansion by a species from a southern refugium may create clines of allele frequencies at neutral loci that are correlated with latitude, and any environmental factor related to latitude, resulting in false positives in EAA (but see Frichot et al. 2015). For instance, in Picea sitchensis along the western coast of North America (Holliday et al. 2010), demography created clines in allele frequencies that confounded tests of neutrality. Controlling for neutral genetic structure reduces the concern about this kind of false positives, because associations among SNPs and environmental factors are assessed after removing the confounding effects of neutral genetic structure (Sillanpää 2011).

Ideally, the subset of neutral markers used to estimate neutral genetic structure is known a priori. However, given that it is generally not possible to know which markers are neutral, a decision about how to best represent neutral genetic structure must be made. First, one can generate a large number of markers across the genome, and all these markers are used to estimate neutral genetic structure (e.g. Eckert et al. 2010a,b). This approach implicitly assumes that the number of loci affected by non-neutral processes in the data set is so small that their effects on global estimates of neutral genetic structure are negligible. Second, two sets of molecular markers can be created, where one set is

used to estimate and control for neutral genetic structure and the other (often including all available markers) is used in EAA (e.g. Bourret et al. 2013). Typically, control markers are from sites in the genome thought to be neutral, such as nonoutliers, synonymous sites (coding for the same amino acid), or noncoding regions. They should be carefully matched against the focal loci with respect to heterozygosity, sample size, minor allele frequency, ascertainment scheme and location in the genome (e.g. in regions with similar levels of background selection, see Berg & Coop 2014; Tiffin & Ross-Ibarra 2014). Consequently, nuclear microsatellites are not the best choice for estimating neutral genetic structure in an EAA using SNPs, as they have very different properties (e.g. mutation rate, allelic diversity) than SNPs.

Traditional methods for estimating neutral genetic structure rely on estimating global or pairwise fixation indices among populations (see, e.g. Holsinger & Weir 2009). In EAA performed at the level of population allele frequencies, not only pairwise fixation indices (e.g. Fischer et al. 2013), but also population-specific fixation indices (sensu Foll & Gaggiotti 2006) can be used to control for neutral genetic structure. Another choice with which to describe population genetic structure in EAA is the estimation of kinship. Numerous estimators of kinship exist (Weir et al. 2006), which can yield substantially different results. Kinship is calculated in a pairwise fashion for all individuals in the data set and is used in subsequent analyses. Note, however, that association approaches using a kinship matrix were developed for GWAS of mostly inbred lines of model organisms. In natural populations, neutral genetic structure might substantially differ from these cases, eventually having unpredictable consequences on the kinship estimator. The use of kinship as an estimator for neutral genetic structure may therefore be inappropriate and remains to be tested. Other popular methods, at the level of individual samples, include matrix factorization methods (e.g. PCA, Patterson et al. 2006) and clustering algorithms like STRUCTURE (Pritchard et al. 2000). Matrix factorization methods produce scores for each individual on each synthetic component, which are used to control for neutral genetic structure in downstream analyses. In contrast, model-based clustering methods result in a Q-matrix, which describes the fraction of each individual's genome attributable to one of the inferred clusters, which is then used to control for neutral genetic structure in EAA.

Analysis of data

In the following, we introduce and discuss the most important and popular methods for EAA (for an overview see Table 1 and Box 2), divided into five broadly defined categories. We recommend applying several environmental association approaches to compare results. This selection is not complete, there are further but less commonly applied methods described in the literature (see, e.g. Jones *et al.* 2013).

Testing categorical factors

Landscape genomics in its simplest form compares allele frequencies of individuals or populations from different types of environments (Box 2A), for example northern vs. southern or high- vs. low-altitude populations. In statistical terms, the different types of environment are introduced as categorical variables in parametric or nonparametric tests. Typically, a neutral genetic model is not implemented (but see, e.g. Foll et al. 2014), and all other environmental factors than the one defining the sampling design are ignored. The most prominent example for such an analysis comes from Turner et al. (2010), who performed Pool-Seq on four populations of Arabidopsis lyrata; two populations originated from serpentine and two from granitic soils. Across eight million SNPs, the authors detected several loci indicative of serpentine soil adaptation, because alleles at these loci were differentiated between soil types and were located in genes with functions associated with conditions characteristic of each soil type.

Logistic regressions

Logistic regressions test whether an environmental factor affects the presence or absence of an allele or single-locus genotype. Although mostly used for dominant markers such as AFLPs, which provide binomial information, logistic regression can also be applied to codominant markers such as SNPs. It is then necessary to prepare the data set in a format that describes the absence and presence of every allele or locus genotype. Because logistic regression can only take two states into account (the presence/absence of an allele or locus genotype), there is no clear way to deal with three or more genotypic states that occur in loci with heterozygous individuals. In this case, an EAA requires multiple analyses, two when using alleles and three when using single-locus genotypes in the case of a bi-allelic SNP. Sampling individuals from diverse habitats or along environmental gradients is ideally suited for this type of analysis.

The spatial analysis method (SAM; Joost *et al.* 2007) was the first implementation of logistic regression in EAA. This approach ignored neutral genetic structure, possibly leading to high false-positive rates under various demographic scenarios (De Mita *et al.* 2013; Frichot *et al.* 2013). Despite this, SAM has been intensively used

in studies of local adaptation. For example, Quintela et al. (2014) combined SAM with the outlier locus detection approach BAYESCAN (Foll & Gaggiotti 2008) to identify AFLP markers and mitochondrial haplotypes associated with water temperature in the freshwater gastropod Radix balthica. Similarly, Nielsen et al. (2009) identified seven outlier SNPs that were related to temperature and/or salinity at spawning grounds of Atlantic cod (Gadus morhua).

Recently, an extended version of SAM, SAMBADA (Stucki et al. submitted; available on arXiv) was developed to overcome some of the limitations of SAM. The software now includes the possibility of multivariate analyses testing, enabling the introduction of neutral genetic structure as an additional factor. SAMBADA can further quantify the level of spatial autocorrelation of genotypes. According to tests performed by the authors, the software is substantially faster than BAYENV2 and LFMM with the univariate model (i.e. not including neutral genetic structure) and faster than BAYENV2 with a bivariate model. SAMBADA comes with a module that can split and remerge large data files. Hence, analyses can be run on different processors in parallel, potentially enabling genomewide analyses. Multiple logistic regressions to test several factors simultaneously including neutral genetic structure can also be performed in R using the generalized linear model function, as shown by Grivet et al. (2011) in a candidate-gene approach in two Mediterranean pine species. An alternative logistic approach is formalized in generalized estimating equations (GEEs, Carl & Kuhn 2007), an extension of generalized linear models with a logit-link and binomial error distribution that considers spatial autocorrelation within populations. It is an individual-based method best suited for sampling designs including many locations from a broad range of environmental conditions, and with a low number of samples per population. According to simulations, GEEs suffer from high falsepositive rates under various demographic scenarios (De Mita et al. 2013).

Matrix correlations

In matrix correlations, one aims to test for correlation between matrices that express distances or dissimilarities between sampling units. A simple Mantel test estimates the strength of correlation (linear or rank linear) between two distance matrices (Mantel 1967) and computes a *P*-value for the correlation coefficient in a permutation procedure. As an extension, the partial Mantel test checks if there is a correlation between two distance matrices given a third matrix (Smouse *et al.* 1986). In EAA, partial Mantel tests can be used with individual or population data. The first matrix includes pairwise

genetic distances or differentiation among individuals or populations at particular loci, the second matrix consists of environmental distances between sampling locations, and the third matrix can be used to control for genetic structure with neutral pairwise genetic distances. Hancock et al. (2011a) performed rank-linear partial Mantel tests using genomewide SNP data from Eurasian accessions of Arabidopsis thaliana, controlling for neutral genetic structure using a kinship matrix based on genomewide genetic variation. They found an enrichment of likely functional variants and could use the results to predict relative fitness in a common garden experiment. Fischer et al. (2013) used linear partial Mantel tests in their study of natural populations of Arabidopsis halleri, with pairwise whole-genome F_{ST} values of over 2 million SNPs as a measure of neutral genetic structure, to identify candidate SNPs for adaptation to five environmental factors.

The (partial) Mantel test has several nice features. For example, it can deal with distances and does not rely on any parametric assumptions. However, Mantel tests have been criticized (e.g. Oden & Sokal 1992; Guillot & Rousset 2013; but see Legendre & Fortin 2010). Guillot & Rousset (2013) showed that, if there is spatial autocorrelation in the two matrices, Mantel tests result in P-values that are not well calibrated, because the permutation procedure fails to produce a valid null hypothesis. One possible solution to overcome this problem is to ignore P-values and concentrate on effect sizes instead (i.e. the correlation coefficient r) when identifying top associations between loci and environmental factors. For example, Fischer et al. (2013) used the 99% quantile of 100 000 simulated r-values as a threshold for relevant environmental associations. Another solution is the use of the nonparametric extension of BAYENV2, which provides a robust alternative approach to (rank based) partial Mantel tests in cases where parametric assumptions are not met.

General linear models

General linear models are statistical models in which a response variable is modelled as a linear function of some set of explanatory variables. These models can account for neutral genetic structure and include statistical methods largely familiar to biologists.

Multiple linear regressions and univariate general linear models. Multiple linear regressions test linear effects of several environmental factors on population allele frequencies and thus enable including neutral genetic structure. For example, several studies (Manel *et al.* 2012b; Zulliger *et al.* 2013) investigated adaptive genetic variation for diverse alpine plant species and used

multiple linear regressions including multiple environmental factors and MEMs to account for the effects of spatial structure and/or unobserved environmental variation. Both studies (Manel *et al.* 2012b; Zulliger *et al.* 2013) found that temperature and precipitation are the driving factors behind local adaptation in alpine plant species.

Some environmental association studies (e.g. Bradbury et al. 2013) have taken advantage of general linear models previously used in GWAS, in which the genotype is the explanatory variable and a phenotypic trait measure the response variable, while controlling for neutral genetic structure with a covariate, for example with the elements of the Q-matrix of STRUCTURE (Pritchard et al. 2000). In EAA, however, environment instead of phenotype is used as response variable. As the environment experienced by an organism is not caused by its genotype, this might seem conceptually counterintuitive. It is assumed, however, that environmental factors that are strongly correlated with heritable traits can replace them in statistical models. An example is illustrated by Eckert et al. (2009), who showed that a linear association between bud flush and mean annual temperature for Douglas fir (Pseudotsuga menziesii) can be described through an association of a SNP affecting bud flush with mean annual temperature. Such general linear models are implemented, for example in the software TASSEL (Bradbury et al. 2007) or can be performed using standard linear modelling in R.

Canonical correlations and multivariate linear regressions. The general linear model framework can be extended to models with multivariate response variables to account for the polygenic architecture of adaptive traits. The most popular method is canonical correlation analysis (CCA), which finds the linear combinations of two sets of variables - multiple loci and multiple environmental factors - that are maximally correlated (Legendre & Legendre 2012). The results are orthogonal sets of canonical variables that can be tested for significance. The loadings by loci and environmental factors indicate which loci respond to which environmental factors. However, users should be aware that strong patterns of multicollinearity could skew the results. Moreover, as CCA does not allow missing data, global deletion of samples or imputation of missing values is often required. Along this line, Mosca et al. (2012) used CCA to show how geographic factors shape the population genetic structure, based on several hundred SNPs, of four subalpine conifer tree species in the European Alps.

A useful approach to test hypotheses about specific environmental factors is redundancy analysis (RDA,

Legendre & Legendre 2012). It allows for building and testing models of varying complexity, including those that condition results based on neutral genetic structure or spatial effects, referred to as partial RDA (pRDA). Significance of the model, each synthetic orthogonal axis and each explanatory variable can be tested using a permutation-based analysis of variance (Legendre & Legendre 2012). Lasky et al. (2012) used pRDA to assess correlations between multivariate climate and multivariate genetic variation in A. thaliana while controlling for spatial effects and identified putatively adaptive SNPs by looking at the contribution of each SNP to the first RDA axis. Using large sets of SNP loci, populations and environmental factors, Bourret et al. (2013) identified temperature and geological factors as drivers of local adaptation in Atlantic salmon (Salmo salar) with RDA. Many of the putatively adaptive genes showed growthrelated functions.

Mixed effects models

The use of mixed effects models is powerful in EAA because they provide a unified statistical framework for controlling for the effects of neutral genetic structure. Here, allele frequencies of individuals or populations are treated as response variables, environmental factors are used as fixed factors, whereas neutral genetic structure is incorporated as a random factor. Approaches differ in how significance is tested, how neutral genetic structure is incorporated, and which type of genotype–environment association (linear/rank-linear/logistic) is assumed.

BAYENV. Coop et al. (2010) developed a Bayesian approach, BAYENV, to assess evidence for correlations between loci and environmental factors. For a given genetic variant, BAYENV tests whether a model that includes an environmental factor has an improved fit to the data compared to a null model that includes only neutral genetic structure, which is represented by a covariance matrix of estimated allele frequencies. BAYENV delivers Bayes factors for each locus-variable combination. One should note, however, that these factors may not be directly compared across environmental variables because of variable-specific value ranges. An advantage of BAYENV is that it allows for the incorporation of uncertainty of allele frequencies that arises from differences in sample sizes. It is not applicable to individual and scattered sampling designs. More recently, Günther & Coop (2013) published BAYENV2, which can be robustly applied to data from Pool-Seq and includes the option of nonparametric tests (Spearman rank correlation). Using Spearman rank correlation showed low detection power in two scenarios simulated by Lotterhos & Whitlock (2015). In cases where the data diverge from assumptions of linearity, however, the relative power of nonparametric tests should increase. Coop et al. (2010) emphasized that the fit of the null model may be imperfect, presumably due to complexities in demography that are not captured by the covariance matrix. Therefore, they suggested to additionally examine other evidence that the approach identifies true signals of selection, such as enrichment of likely functional variants (e.g. nonsynonymous substitutions) in the distribution tails of the resulting Bayes factors. A recent study by Blair et al. (2014) showed that the run-to-run variation of BAYENV (version 1) can be large. These authors thus advise to average Bayes factors among multiple runs to produce more stable and reliable results.

BAYENV was the first method specifically developed for EAA that controlled for neutral genetic structure. As a result, it has been used in several large-scale studies of candidate genes and for genomic data sets. Hancock et al. (2008) applied an early version of this approach to candidate loci for energy metabolism genotyped in a worldwide set of human populations. Subsequently, they used BAYENV with a human genomic data set to identify correlations using both continuous and categorical environmental factors (Hancock et al. 2010, 2011b). The studies identified enrichment of nonsynonymous SNPs, variants associated with disease traits and ecologically relevant sets of genes among the loci correlated with environmental factors. BAYENV has also been applied to studies of local adaptation in candidate genes in tree species, first by Eckert et al. (2010a) in loblolly pine (Pinus taeda) and later in different spruce (Picea) species (Chen et al. 2012; Prunier et al. 2012).

Using simulations, BAYENV was shown to detect a relatively low rate of false positives (De Mita *et al.* 2013) and to perform best under scenarios with weak hierarchical genetic structure (de Villemereuil *et al.* 2014). However, BAYENV is slow because it is computationally very intensive (De Mita *et al.* 2013; Stucki *et al.* submitted) and therefore less suited for analyses of a large number of genetic polymorphisms. A related method is GINLAND (Guillot *et al.* 2014), a spatial generalized mixed model (SGLMM) which uses a Markov chain Monte Carlo (MCMC)-free approach with shorter computing time. GINLAND also considers pure spatial autocorrelation based on a geographical distance matrix. To our knowledge, GINLAND has not yet been used in any empirical study.

Latent factor mixed models (LFMMs). In LFMMs (Frichot et al. 2013), neutral genetic structure is introduced as a random factor with the so-called latent factors, which

are similar to principal components and calculated from all available markers. The advantage of this linear approach is that the effects of environmental factors and neutral genetic structure on allele frequencies are simultaneously estimated. Moreover, computing time is reasonably fast, making LFMM attractive for EAA with whole genomes or subsets of large random batches of SNPs in parallel. This approach surpasses the need for specifically formalizing neutral genetic structure, and it works without knowledge about which loci are putatively neutral, which is often not available in advance. LFMM computes Z-scores and P-values to quantify the strength of associations and which are also informative when compared among environmental factors. Before starting the final analysis, the number of latent factors (K) has to be chosen, either by an analysis of histograms of test P-values for different K-values (i.e. it should look similar to a uniform distribution), by performing a Tracy-Widom test on the eigenvalues of a PCA on the genetic data, or using programs such as STRUCTURE (Pritchard et al. 2000) to determine plausible values for K. As the stochastic algorithm of LFMM (MCMC) does not provide exact results, Frichot et al. (2013) recommend to perform multiple runs, use the median of the resulting Z-scores and adjust their P-values as described in the software manual. The software LFMM comes with two different interfaces, a graphical user interface and a command-line version. Only the latter can handle population allele frequencies. LFMM is therefore suited for both population based and scattered, individual-based sampling designs.

Frichot et al. (2013) found that LFMM has low rates of false positives and negatives and that it performs slightly better than BAYENV in detecting weak selection. de Villemereuil et al. (2014) showed that LFMM provides the best compromise between detection power and error rates in situations with complex hierarchical neutral genetic structure and polygenic selection. Finally, Lotterhos & Whitlock (2015) showed that LFMM is quite robust to a variety of sampling designs and underlying demographic models. LFMM has been used in several recent empirical studies. For example, Zueva et al. (2014) investigated pathogen- and environment-driven selection in populations of Atlantic salmon. They identified around 900 of the 4631 tested SNPs to be associated with one of the five environmental factors considered, including parasite-induced mortality as a measure for pathogen-driven selection. De Kort et al. (2014) found strong associations between temperature and 15 outlier SNPs in black alder (Alnus glutinosa) and showed, with additional evidence from a common garden experiment, that temperature is the main driver of local adaptation in this drought-sensitive tree species.

GWAS mixed models. Mixed models have been a standard approach for some time for the discovery of genotype-phenotype associations (Korte & Farlow 2013). As in the general linear models described above, environmental association studies have taken advantage of computationally efficient GWAS methods by replacing the response variable phenotype by environment. Kang et al. (2008) developed an efficient mixed-model association (EMMA) method that includes a simple identity-bystate allele sharing kinship matrix to control for neutral genetic background. EMMA was used to associate the RegMap panel SNPs (Horton et al. 2012) in A. thaliana to cold- and moisture-related climatic factors (Lasky et al. 2014). Genes with genetically variable expression responses to abiotic stress were enriched by SNPs strongly associated with climate. It is important to note that EMMA is optimized to test associations of only one allele with climate. Allowing heterozygous genotypes of outbred individuals is possible, but complex and computationally intensive (Kang et al. 2008). Moreover, the use of a kinship matrix to describe neutral genetic structure of populations may be inappropriate. Similarly, a linear mixed-model method is implemented in the software TASSEL (Bradbury et al. 2007). For example, Yoder et al. (2014) tested for associations of nearly 2 million SNPs to three climatic factors in 202 inbred accessions of barrel clover (Medicago truncatula). They identified more than 20 genes that were associated with climate and have a function in response to abiotic factors and pathogens in homologs of A. thaliana. GWAS mixed models are designed for individual rather than population sampling, making them best suited for analyses with samples continuously distributed across a study region.

Limitations and extensions of environmental association analysis

The main hurdle for EEAs (and notably also of population genomic approaches, De Mita et al. 2013; Lotterhos & Whitlock 2014) is that they might result in high rates of false positives (De Mita et al. 2013; Lotterhos & Whitlock 2014; de Villemereuil et al. 2014; Frichot et al. 2015), which are significant associations that are actually not casual. The main reason is that geographic and demographic processes can lead to patterns that mimic those observed as a consequence of selection. In fact, de Villemereuil et al. (2014) found high rates of false discovery in some scenarios with complex, hierarchical structure and polygenic selection. Fortunately, applying analyses that control for neutral genetic structure can mitigate this problem. De Mita et al. (2013) simulated different demographic, selective and mating type scenarios and found false-positive rates of up to 40% (logistic regression) and 50% (GEE) for approaches not specifically correcting for neutral genetic structure, but only 20% for BAYENV, which corrects for structure. Depending on the combination of approach and scenario, power and error rates differed greatly in this study. Similarly, Frichot et al. (2013) reported low false-positive rates (0-7%) for methods that correct for neutral genetic structure. Unfortunately, some demographic scenarios may be particularly challenging for EAA. For example, scenarios in which the range expansion of a species creates a cline in allele frequencies along an environmental gradient (Keller et al. 2009; Novembre & Di Rienzo 2009) or in which individuals/populations are under strong isolation by distance (Lotterhos & Whitlock 2015) are hard to deal with in EAA (but see, Frichot et al. 2015). False positives can also arise due to the failure to account for multiple testing, which is needed when a large number of loci and environmental factors are included in the analysis. We strongly recommend to control for false-discovery rate (FDR) using the algorithms described by Benjamini & Hochberg (1995) and Storey & Tibshirani (2003). FDR (unlike, e.g. classical Bonferroni correction) does not depend on the number of tests and aims to accurately estimate the proportion of false discoveries among positive findings. A third cause of false positives is that it can be difficult to distinguish between correlated environmental selective pressures. More specifically, observed correlations with a specific environmental factor can be due to adaptation to covarying factors that were not included in the analyses or excluded in the process of factor reduction. In these cases, it is the association, not the locus, that represents a false positive. In other words, the detected locus might actually play a role in local adaptation, but is linked to a different factor. For example, the presence of an allele may be correlated with high temperature, but is actually involved in defence against pathogens whose development, survival and transmission is sensitive to temperature (Harvell et al. 2002). Moreover, correlations among loci (i.e. linkage disequilibrium between an adaptive locus and other variants) can result in a spurious signal of correlation at linked variants (hitchhiking, Strasburg et al. 2012). Finally, false positives can also derive from coincidental outlier values of environmental factors and allele frequencies. A simple way to deal with these cases is to avoid populations with extreme environmental values already in the sampling design, or to use rank-based, nonparametric statistics such as BAYENV2 or rank-linear partial Mantel tests. In any case, landscape genomic studies should carefully consider the issue of false positives, keeping in mind that applying stricter thresholds to possibly account for this issue will result in lower power to detect true positives and will inflate the rate of false negatives.

As for most biological studies, the results of EAAs are restricted to the sampled populations and environmental conditions. Therefore, several studies (e.g. Poncet et al. 2010; Prunier et al. 2012; Buehler et al. 2013) have considered geographical subsets that were analysed separately to detect more general patterns. Overlap among identified loci of adaptive relevance of such population subsets is, however, often minimal. For example, Poncet et al. (2010) found 61 and 21 climate-related AFLP loci in populations of the alpine rockcress (Arabis alpina) from the French and Swiss Alps, respectively. Only four of these loci were found in both regions. This result implies the presence of false positives (in the case of the SNPs that were only identified in one region) or to geographically restricted patterns of adaptation.

Combined approaches and downstream analyses

Given the issues discussed in the preceding section, it is desirable to combine EAA with other approaches in order to reduce the rate of false positives and to assess the relevance of findings. In this section, we list a selection of such integrative approaches (for more ideas, see, e.g. Pardo-Diaz *et al.* 2015) and exemplify them with respective empirical studies.

Combination with tests for outlier locus detection. Instead of opposing EAA and outlier detection methods, one could combine them to obtain more information from the data. For example, one could first perform an outlier test using, for example BAYESCAN (Foll & Gaggiotti 2008), FDIST and derivates (Beaumont & Nichols 1996; Beaumont & Balding 2004), FLK (Bonhomme et al. 2010), or ARLEQUIN (Excoffier & Lischer 2010) and use only the resulting outlier loci in subsequent EAA. For example, Fischer et al. (2013) used POPOOLATION (Kofler et al. 2011) to select the most extremely differentiated SNPs of A. halleri and subsequently correlated the resulting outlier loci to topo-climatic factors using partial Mantel tests. Selection processes that lead to small shifts in allele frequencies, however, are not likely to be detected with this strategy, and the overlap among different methods can be small (de Villemereuil et al. 2014). Alternatively, one could perform multiple analyses in parallel using the entire set of loci, and then discuss the results by comparing the two lists of putatively adaptive loci (e.g. Quintela et al. 2014). Finally, in EAAs using a categorical sampling design, one could perform outlier tests among groups of individuals that are defined by the environment (e.g. Buehler et al. 2013; Roda et al. 2013), while appropriately dealing with neutral genetic structure. Buehler et al. (2013) used DFDIST (Beaumont & Balding 2004) in A. alpina to identify one

outlier AFLP marker that exhibited particularly high genetic differentiation among three contrasting habitat types. Foll *et al.* (2014) recently presented a flexible hierarchical extension of the BAYESCAN approach (Foll & Gaggiotti 2008), which allows for the simultaneous analysis of populations living in different environments in several distinct regions. It includes a convergent (parallel) evolution model that directly identifies candidate loci in replicated pairs of populations instead of using intersecting sets of candidate loci.

Gene function and gene ontology analyses. Recent technological and scientific advances have not only resulted in the availability of reference genomes for numerous species, but also led to the establishment of public databases where annotated genes are described in detail. For several model species, large parts of their genomes are now annotated, although not with the same level of reliability (Primmer et al. 2013). Most studies on evolutionary and molecular ecology, however, focus on nonmodel species. While draft genomes for nonmodel species are emerging (Ekblom & Galindo 2011), they still often lack annotation (Primmer et al. 2013). Fortunately, in most cases, annotation from related model organisms can be transferred to less well-studied species by identifying homologous sequences, assuming that they have the same function in both model and study species.

Gene ontology (GO) databases describe the biological process, molecular function and cellular component of a gene in a standardized, species-neutral vocabulary (Primmer et al. 2013). They therefore enable linking EAA with gene function. Many EAA studies rely on GO databases in one or the other way, not only in the planning phase (e.g. for selecting candidate loci), but also for downstream analyses. In most cases, researchers try to verify the biological function of a gene post hoc. In the best case, gene function appears reasonable in the context of the associated environmental factor (e.g. Eckert et al. 2009). This inference increases evidence that a given association is not purely coincidental. An additional option for EAA are GO enrichment tests (e.g. Fischer et al. 2013), which examine whether certain gene functions are over- or under-represented in a set of genes (e.g. those associated with an environmental variable).

Nonsynonymous vs. synonymous substitutions. Not all nucleotide substitutions lead to changes in the encoded amino acid. Usually, the third nucleotide of a codon is silent (synonymous, i.e. the derived codon codes for the same amino acid) and therefore thought to be selectively neutral. Annotation of investigated polymorphisms can therefore be applied to interpret the results

obtained from EAA. This is only feasible if a reference genome of the investigated or a closely related species is available. The occurrence of nonsynonymous (amino acid changing) SNPs, especially if it also concerns SNPs significantly related to environmental factors, can increase evidence for relevance in adaptation. If many substitutions are present, one can calculate the ratio of nonsynonymous to synonymous variants within the distribution tail of the EAA and compare this to the ratio in nonsignificant loci. For example, Hancock *et al.* (2011a) looked at the top 1% of SNPs associated with climate in *A. thaliana* and found an enrichment of nonsynonymous compared to synonymous and nongenic substitutions.

Post hoc validation in independent data sets. Replicated patterns of local adaptation can derive from the spread of an adaptive allele to multiple geographic locations or by repeated and parallel adaptation (discussed, e.g. in Schmidt et al. 2008; Nosil et al. 2009; Prunier et al. 2012; Buehler et al. 2014; Tiffin & Ross-Ibarra 2014). However, studies using an independent data set to test the generality of adaptive loci are rare. Buehler et al. (2014), using 30 independent populations of A. alpina, did not find the same association of an AFLP outlier locus as identified previously (Buehler et al. 2013). In contrast, 15 previously identified AFLP loci of the gastropod Littorina saxatilis exhibiting signs of selection were distributed in the same clinal manner on two independent shores along the Atlantic coast in England (Grahame et al. 2006). Although such a validation step represents a useful addition to EAA, successful validation in an independent data set is not necessarily expected. This is because locus-specific selection is crucially dependent on the local genomic context and local environmental conditions, and genotype-by-environment interactions may modulate selection patterns in an unpredictable way (Schmidt et al. 2008), leading to geographically restricted local adaptation. However, finding recurrent patters in independent data sets greatly improves evidence for the generality of adaptive patterns detected.

Experimental validation. Direct proof that a genetic variant actually leads to a fitness advantage in a local environment can only be obtained experimentally (Barrett & Hoekstra 2011; Savolainen et al. 2013). Compelling support for EAA (or GWAS) findings is to employ a common garden experiment, in which genotyped individuals coming from different habitats share the same natural or manipulated environment(s) and are measured for fitness-related phenotypic traits (e.g. Fournier-Level et al. 2011; Hancock et al. 2011a; De Kort et al. 2014; Yoder et al. 2014). To this end, Hancock et al. (2011a) identified climate associations in A. thaliana

accessions from across Eurasia and found that the identified SNPs could be used to predict rank fitness in a common garden. Conversely, Fournier-Level et al. (2011) grew hundreds of inbred A. thaliana lines derived from natural populations across their native distribution and planted them in four European field sites (common gardens) that spanned the species' native range. Alleles that were associated with higher fitness in particular common gardens were more frequent in the respective environment the plant originated from. In theory, only in reciprocal transplant experiments, it is possible to test whether the fitness of 'home' populations is actually higher than that of 'away' populations (Kawecki & Ebert 2004). Although reciprocal transplant experiments have been carried out repeatedly in the past (e.g. see Savolainen et al. 2013), they have mostly been conducted at the phenotypic level and have rarely taken advantage of genomic information. In the context of EAA, reciprocal transplant experiments are the perfect addition to check for fitness advantages of given alleles associated with particular environments. We are not aware of a study that has validated identified associations with this often laborious approach. While transplant and common garden experiments with genetic variants might be feasible in the case of processes of monogenic adaptation, they could be challenging for polygenic adaptation. One should also bear in mind that the potentially different genetic backgrounds of populations included in experiments can interfere with the detection of the adaptive signal (Holderegger et al. 2008). Finally, it should be noted that even if a fitness advantage is not detected in the above-described experiments, it does not mean it does not exist, as the results and interpretation of the experiment is bound to the experimental conditions (site conditions, duration, age of individuals, interactions with pathogens, etc.) and the measure of fitness.

Future directions

Options for further developing the field of EAA are manifold and involve theoretical, methodological and statistical issues, some of which we highlight in this section. One of the major challenges in EAA is the computational bottleneck resulting from increasingly large data sets, made possible by decreasing sequencing costs. Effort therefore will likely be put into developing faster algorithms that adequately control for neutral genetic structure.

Another issue to be improved in EAA is the lack of methods that can deal with nonlinear responses at the level of population allele frequencies. In a scenario that does not assume gradual changes in allele frequencies along an environmental gradient, but in contrast rather corresponds to a sweep model where beneficial alleles are mostly fixed, one would expect extreme allele frequencies. Accordingly, only a narrow range of environmental conditions should exhibit intermediate allele frequencies (Box 2D). Therefore, we suggest that effort should be put into the development of alternative population-based models. SAM and SAMBADA, which use logistic regression with a sigmoid-like response curve, deal with individual-based presence/ absence data and not allele frequencies per population. Hence, they are not applicable to Pool-Seq. In addition, the interpretation of loci with heterozygous individuals is difficult. One possibility could be sigmoidal curve fitting with the challenge of incorporating neutral genetic structure. A promising development in this respect is integrated in BAYESCENV (de Villemereuil & Gaggiotti in press), which extends the outlier detection software BAYESCAN (Foll & Gaggiotti 2008) by implementing an additional model that includes information about the environment. It is based on genetic and environmental distance and can also detect patterns of allele frequency that are not linearly dependent on environmental factors. Along the same line, although one of the main arguments for using EAA is the detection of polygenic adaptation, most methods presented here only test single-locus effects. There is a clear lack of approaches that test associations with multivariate response variables (Sork et al. 2013), with the exception of multivariate analyses like CCA and RDA described above.

Analogs to the GO databases, which mostly stem from a cellular perspective, have recently been advocated based on an ecological association ontology that includes findings from ecological and evolutionary studies (Landry & Aubin-Horth 2007; Pavey *et al.* 2012). This so-called ecological gene annotation would complement existing GOs, introduce a vocabulary used by evolutionary ecologists, present links to ecological and evolutionary studies and decrease the proportion of genes with unknown function. Landscape genomics could nicely contribute to, and also greatly profit from such an additional source of information (for discussion, see also Primmer *et al.* 2013).

Finally, there is need for more post hoc validation regarding function and adaptive generality of the alleles, loci or genomic regions identified in EAA. Many studies, including most of those described in this review, perform EAA, present a list of interesting loci, compare it to GO databases and stop there, that is half way to the goal of identifying those genes that are functionally involved in local adaptation of natural populations. Instead, studies should go further and test their findings using, for example independent populations, knockout mutants, common garden and reciprocal

transplant experiments. The effort of such follow-ups should, however, not be underestimated.

Conclusions

The recent advances in sequencing technologies have opened the door for analyses of the genetic basis underlying local adaptation. Landscape genomics is an emerging research area, focused on understanding the role of the environment in genetic adaptation and identifying putatively adaptive loci. EAAs are the main tools of this research field. In this review, we described several strategies to test for associations between genotypes and environment. We strongly recommend controlling for neutral genetic structure in the analyses, carefully applying and comparing complementary statistical approaches, using a large number of populations and/or individuals, and looking at as large as possible fraction of the genome. It is clear that for many researchers, especially for those with limited financial budgets or those working on nonmodel organisms, some of these goals are hard to reach. Luckily, sophisticated and increasingly inexpensive genomic technologies are emerging. It is important to note that landscape genomics, however, has yet several relevant issues to solve. Besides the potential for further extending current statistical models for EAA, the main concern is the lack of post hoc testing of fitness advantages of putatively beneficial genetic variants in specific environments. After the gold rush in finding loci that are associated with important environmental drivers, it is now time for the field to take the next step and show that promising candidates are linked to fitness-related variation and thus relevant for adaptation.

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