





Constrained ordination method as a tool for performing genome scan and environmental genomic studies

Eric Bazin,*,1 Michael G. B. Blum,2

¹LECA, Université de Grenoble

 $^2{\rm TIMC},$ Université de Grenoble

*Corresponding author: E-mail: eric.bazin@univ-grenoble-alpes.fr

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Abstract

Key words:

1 Introduction

Performing genome scan in order to detect 21 genomic region of interest is a common task 22 in population genomic area (Foll and Gaggiotti, 23 2008; Frichot et al., 2013; Luu et al., 2016; Vatsiou 24 et al., 2015). Some methods aim at detecting genes 25 that has suffered from a loss of genetic diversity 26 and increase of linkage disequilibrium following 27 the appearance of a beneficial allele and its spread 28 by the mean of selective sweep. Others aim at 29 picking up alleles with strong correlation with 30 some environnemantal variable (e.g. Temperature, 31 drought) with the idea that these alleles may 32 confer a selective advantage to the individuals 33

involved in local adaptation process. These region should have an increased differentiation between

population because different alleles tend to be

(Coop et al., 2010; Frichot et al., 2013). Finally,

other methods aim at detecting genomic region

between population is excepted under the hypothesis of geographical isolation. Therefore, this region can be detected by quantifying the level of differentiation using some statistics and detecting the regions with unexpectedly high values. A common statistic and very easily comprehensible in population genetic is Fst. Many methods use this parameter as a basis in many different implementation of genome scans (Bazin et al., 2010; de Villemereuil and Gaggiotti, 2015; Foll and Gaggiotti, 2008). These are model based method where parameters such as Fst are usually inferred using likelihood or Bayesian methods. This mean that users must have some a priori on their parameter value and the best model that fits their data in order to expect the best from their analysis. However, it is often difficult to get a satisfactory a priori picture of the demographic and population structure of the

beneficial in each environment. Differentiation

40 species one is interested in Indeed many species © The Author 2013. Published by Oxford University Press on behalf of the Society for Molecular Biology and Evolution. All rights reserved. For permissions, please email: journals.permissions@oup.com









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are not clearly structured in different populations 73 longitudinal) but they are not necessary linked to but more or less show a pattern of isolation 74 an environmental variable such as Temperature, by distance without clear geographical barrier 75 drought, diet habit, etc. Therefore, when this to gene flow. One solution would be to use 76 information exists, it has to be a posteriori used more complex model that better reflect reality 77 as a mean of interpretation but are not involved but these are difficult to implement in Bayesian 78 in the inference process. It should be recalled framework. Additionally, these latter methods are 79 that natural selection is the result of a complex very time consuming and the increase of both 80 set of environmental pressures and that it most model complexity and the amount of data to 81 often acts on several characters simultaneously analyze in terms of the number of individuals and 82 and that these characters are encoded by several loci makes them more and more difficult to use. 83 genes which generally have weak effects. In A new path has opened recently with the use 84 order to extract the maximum of all available of multivariate methods. The idea is to capture 85 information, it seems therefore necessary to use the whole genome geographic structure using an 86 approaches that are able to compile all kind of ordination method such as ACP. Following this 87 variable (e.g. alleles, phenotypic measurement, analysis, outliers loci are detected if they have 88 biotic and abiotic variables). One natural way extremely high correlation with one or more 89 to overcome this limitation would be to use ordination axis (Duforet-Frebourg et al., 2014; 90 more sophisticated ordination method than ACP Luu et al., 2016). These are very efficient methods 91 like methods. Constrained ordination methods and simulations have shown that while they 92 (i.e. Redundancy Analysis, RDA, Canonical are very fast, they show similar efficiency than 93 Correspondence Analyis, CCA) are well-known classical Bayesian method and sometimes perform 94 set of approaches in Ecology for instance to better when the simulated demographic model 95 explain the species distribution pattern by drift from the model implemented in bayesian 56 the mean of environmental data. They have method, usually the island model. For instance 97 specifically been designed in order to deal with Luu et al. (2016) have shown their method 98 biological complexity. In the population genomic to be better when population are structured 99 era, it seems that data amount, complexity in hierarchical set or in isolation by distance 100 and heterogeneity is often a limitation to the pattern. Nevertheless, one conundrum of such 101 use of inference methods based on classical approaches is the difficulty to interpret ordination 102 population genetic models. Although they are axis in term of ecological meanings. These are 103 more difficult to interpret, such approaches would

usually tight to geographical axis (latitudinal or 104 be complementary to the model based method









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because of their long-term use in ecology and their 137 results than PCA-based method. Second, thanks efficiency on complex and large datasets. These 138 to these simulations, we show that RDA can method have sometimes been used in population 139 indeed help to identify important environmental genomic studies, not as a genome scan but in 140 order to quantify multilocus adaptation to an 141 environmental gradient (De Kort et al., 2014; 142 Steane et al., 2014; ?; ?). These studies whereby 143 relationships between environmental data and 144 large multilocus data is explored are becoming 145 more and more popular and are often coined 146 as Ecological Genomics or Landscape Genomics 147 studies. However the concept of using constrained 148 ordination methods to analyse genomic data has never been tested on simulated datasets. This 150 paper aims at filling this gap. First, we show 151 how one can make use of a constrained ordination $_{152}$ method namely Redundancy Analysis (RDA) as 153 an efficient and robust genome scan method. $_{\scriptscriptstyle 154}$ We discarded the other constraint ordination 155 methods such as CCA since they are very similar 156 in their principles. RDA has already been used $_{\scriptscriptstyle 157}$ for instance by ? to perform genome scan in $_{158}$ order to detect loci involved in the adaptation 159 to climate in Arabidopsis thaliana. Outliers were 160 identified as SNPs with the greatest squared scores $_{161}$ along the first RDA axis (i.e. those in the 0.5_{162} % tail). We build on this idea to develop a $_{\scriptscriptstyle 163}$ comprehensive and robust statistical test that $_{164}$ allows to search for outliers on an arbitrary 165 number of RDA axis simultaneously and allows 166 to control precisely for the false discovery rate. 167 Using simulations, we show that it has better 168

gradient that better explain the adaptive variation in the data. It is therefore a proof of concept of the idea of using constrained ordination method as an environmental genomic tool to identify relevent selective gradient in the environmental data. Finally, to give a concrete illustration of RDA approach in population genomics, we apply this method to the detection of outliers on a real data set.

Material and method

Genome scan

Redundancy analysis (RDA) was first introduced by (Rao, 1964) and is clearly described in (Legendre Pierre & Legendre, 2013) section 11.1. It is the direct extension of multiple regression to the modeling of multivariate response data. Typically the data to be analysed are separated in two sets, a response matrix Y of variable to be explained (e.g. species abundance in a set of sites; m sites and n species) and an explanatory matrix X (e.g. a set of environmental variable within each site; m sites and p environment). In the following analysis, species are replaced by loci and sites by individuals. In other word, we wish to project on a reduced space the proportion of variance in genetic difference between individuals which is better explained by environmental data. After this ordination, we follow the Luu et al. (2016) methodology to compute pvalues. First we





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compute the test statistic by regressing each of the 200 in the differentiation pattern (Frichot et al., p SNPs by the K ordination axis $X_1,...,X_K$. $G_j = \sum_{k=1}^K \beta_{jk} X_k + \epsilon_j, j = 1, ..., p$ 171 where β_{jk} is the regression coefficient 203 corresponding to the j-th SNP regressed by 204 multivariate method for genome scan is based. 173 the k-th ordination axis, and ϵ_i is the residuals 205 vector. To summarize the result of the regression 206 analysis for the j-th SNP, we return a vector of 207 z-scores $z_j = (z_{j1},...,z_{jK})$ where z_{jk} corresponds 208 using scatter plots. To control for false positive, to the z-score obtained when regressing the 209 j-th SNP by the k-th ordination axis. The test 210 statistic is a robust Mahalanobis distance D $_{211}$ Once outliers have been identified, we isolate them computed using covRob function of the robust R $_{\tiny 212}$ package. D should be Khi2 distributed after $_{213}$ a correction with inflation factor (Luu et al., 214 2016). Pvalues are computed using K degree of $_{215}$ freedom. We use the FDR approach to control 216 for false positives. Qvalue are computed with $_{217}$ qvalue R package and a loci is considered as an $_{\tiny 218}$ outlier if its qualue is less than 10%. For the 219 analysis of simulated dataset (see below), we $_{220}$ retain the first four ordination axis to compute 221 Mahalanobis distances as they seem to explain 222 most of the variance in the data. To peform 223 the ordination, we use the 10th environmental $_{224}$ variables as input in the explanatory matrix. In 225 the following example, we don't use phenotypic 226 informations since these informations are often 227 laking in environmental genomics. Neither we use geographical coordinates (i,j) which is sometimes $_{229}$ added to control for the geographical covariation $_{\scriptscriptstyle{230}}$

To emphasize the utility of RDA, we compared to pcadapt from which the idea of using On the simulated dataset, we retain K=3 axis to compute Mahalanobis distances as it seems to explain the main amount of variance in the data we used the same qualue threshold (i.e. j=10%). Environmental genomic

in a separate matrix A defining an "adaptively enriched genetic space" as coined by Steane et al. (2014). Following their methodology, we perform a second constrained ordination (RDA) on matrix A against environmental data. The rational of this analysis is to remove neutral variation before performing ordination in order to have a better picture of which environmental gradients have the strongest association with the adaptive genetic space. On the simulated dataset, we report the R^2 statistics between env1, env2 and env3 and the first three ordination axis to have an idea of which they are better associated with and if the ordination space succeed in seperating the environmental effect on different axis.

Simulations

To test for the efficiency of RDA in population genomic, we performed simulations using simuPop python library (?). We compared our approach to PCAdapt method to perform genome scans. Both









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approach are equivalent except their ordination 264 10 different environmental variables. The first one method. Finally we use these simulations to 265 determines the selective pressure on trait 1, the evaluate RDA approach as a mean to detect 266 second one on trait 2 and the third one on trait selective environmental gradient. A lattice of 267 3. The first environment variable is a quadratic 8x8 populations is simulated (i.e. 64 populations 268 gradient coded by function $env1 = -(\cos(\theta))$ * in total). Each population is initialized with $_{269}$ $(i-3.5))^2-(sin(\theta)*(j-3.5))^2+18,\theta=\pi/2$, i and 200 diploid individual with random genotypes. 270 j being the population indicator on the 8x8 Migration is set to 0.5 so that population structure 271 lattice. The second one is a linear plan gradient must be very smooth and genetic differentiation 272 coded by function $env2 = h*\cos(\theta)*(i-1) + h*$ must show an isolation by distance pattern over $\sin(\theta) * (j-1) + k$ with h=2, $\theta = \pi/4$ and k=1the 64 populations. This is where pcadapt is 274 3. The third environment variable simulates best designed for. Loci are biallelic (0 or 1) like $_{275}$ a coarse environment with value env3=2 for SNPs. Allele frequency of the whole population is $_{276}$ all populations except population (i,j) = (2,2), initialized at 0.5. 1000 loci are defined. They are 277 (2,3), (3,2), (3,3), (6,2), (6,3), (7,2), (7,3), (2,6), separated in 200 chuncks of 5 SNPs in physical 278 (2,7), (3,6), (3,7), (6,6), (6,7), (7,6), (7,7) for linkage with recombination rate between adjacent 279 which env3=18. Env4, env5 and env6 have loci fixed at 0.1. 3 different Traits are coded by 280 exactly the same equation than env1, env2 and a group of 10 different loci. The first trait is 281 env3 respectively. The remaining 4 environment coded by loci 1, 11, 21, ..., 91. The trait value is 282 variable are similar to env2 but with different simply the sum of genotype value and therefore 283 value of h and θ . Env7 has h=2, $\theta=0$ and k=1can take value between 0 an 20. For the sake 284 3. Env8 has h=2, $\theta=\pi/4$ and k=0. Env9 has of realism, we add to each trait a random noise 285 $h=1, \theta=\pi/4$ and k=4. Env10 has $h=0.5, \theta=$ (non heritable variation) drawn from a normal 286 $\pi/4$ and k=8. Graphical representation of mean distribution N(0,2). The second trait is coded by 287 environmental value for environment 1, 2 and loci 101, 111, ..., 191 and the third is coded by loci 288 3 is given in Fig. ??. Environment 4, 5 and 6 201, 211, ..., 291. Each trait is therefore coded by 289 have respectively the same mean value spatial free recombining SNP loci. In other words, there 290 distribution. For a graphical representation of are 30 coding SNPs among 1000. Selection can 291 environment 7 to 10, see supplementary material. have an effect on linked loci, for instance, loci 2, 292 3, 4 and 5 can be impacted by selection on locus 293 the environment variable. To avoid colinearity 1. However, recombination is high enough (0.1) to 294 between environments variable, we added noise by

Environmental equation gives a mean value of expect a limited linkage effect. We have defined 295 drawing an environment value within a normal









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distribution $N(\mu = env, \sigma = 1)$. Fitness for each 328 trait is set to be $-e^{((x-env)^2/(2*\omega^2))}$, x being the 329 quantitative trait value, env the environmental 330 value and ω is defining selection strength and 331 has been set to 10 which in our experience seems 332 sufficient for loci to be often detected. To get 333 the overall fitness for a given individual, fitness 334 associated to each trait are multiplied. Fitness 335 are relative and selection arises on parents and 336 determine their number of offsprings. Simulations 337 are made across 500 generations. At the end 338 of simulation, we sample 10 individuals per 339 population. Therefore, we have a sample of 640 340 individuals with 1000 SNP-like loci. Real dataset 310 The Loblolly pine dataset is a sample of 682 343 individuals genotyped on 1,730 SNPs selected in 344 ESTs (Eckert et al., 2010). 60 climatic variables 345 were available and summarized by the authors 346 in the five first axis of a PCA. The first axis, 347 PC1 is mainly linked to latitude, longitude, 348 316 temperature, and winter aridity. PC2 is linked 349 to longitude, spring-fall aridity, and precipitation. 350 We inputed the missing data using a very simple 351 algorithm implement in function sing.im of the R $_{352}$ package linkim (Lachenbruch, 2011). It imputes 353 the missing value based on the observed data $_{354}$ proportions. We used K=4 axis to compute 355 Malahanobis distances. The Chinook salmon consists of 19 703 SNP 357 loci genotyped on 1956 total individuals pooled 358

StreamOrder, bio03, bio17 and bio18) been used among 24 different climate and environmental variables because they have been tested as significantly associated with the SNP variation rangewide citepHecht2015. MigDistKM stands for Migration distance from collection site to ocean (km), StreamOrder for Stream Order of collection site using Strahler method, bio03 for Isothermality, bio17 for Precipitation of Driest Quarter (mm) and bio18 for Precipitation of Warmest Quarter (mm). We could have tested more variable but this is just an illustration and is by no mean an extensive study of this species. Since data are pooled, we have randomly created a sample of 100 individuals for each collection based on the allele frequencies to be able to analyze the data following our individual based pipeline. We used K=4 axis to compute Malahanobis distances.

Results

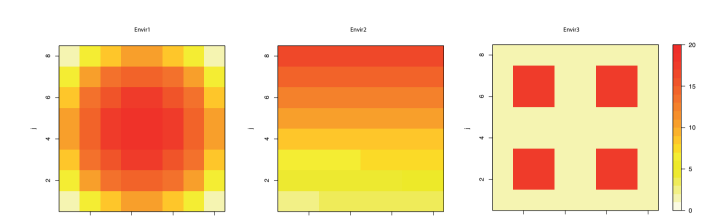
Genome scan

temperature, and winter aridity. PC2 is linked 349 When looking at the analysis on one simulation, to longitude, spring-fall aridity, and precipitation. 350 the pcadapt method seems successful at detecting We inputed the missing data using a very simple 351 QTL2 SNPs (Fig. 2) but fails at detecting algorithm implement in function sing.im of the R 352 QTL1 and QTL3 SNPs. On the other hand, package linkim (Lachenbruch, 2011). It imputes 353 RDA succeeds at detecting QTL2 SNPs and the missing value based on the observed data 354 also some of the QTL1 and QTL2 SNPs (Fig. proportions. We used K=4 axis to compute 355 3). The ordination seems to correctly detect Malahanobis distances. 356 environmental variable 1 and 3 as drivers of genetical variance in the data. Over the 100 loci genotyped on 1956 total individuals pooled 358 simulations, we have measured the average FDR in 46 collections. Five variables (MigDistKM, 359 and power for both pcadapt and RDA (Fig 4).









 ${\bf FIG.~1.}$ Graphical representation of mean environmental value for environment 1, 2 and 3

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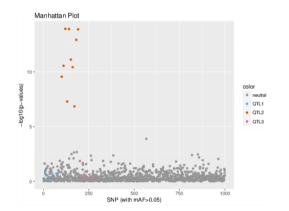


FIG. 2. Manhattan plot of the result of pcadapt on a simulated data set.

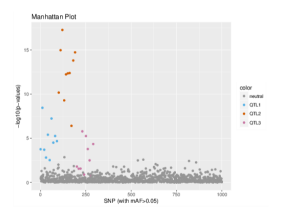


FIG. 3. Manhattan plot of the result of genome scan using RDA on a simulated data set.

Environmental genomics

We then performed a second RDA on the $_{382}$

dataset as in Fig. 2 and 3 and display its results on Fig. 5. We did the same analyis and measured the mean R^2 between env1, env2 and env3 and each of the first three ordination axis. This is summarized

Loblolly Pine

in Fig. 6.

Discussion

Fig 2 shows that pcadapt approach works well when the environmental gradient and the selective pressures are acting in the same direction than the geographical pattern of isolation by distance. Whereas when the environmental gradient is quadratic on the geographical range (QTL1) or when it is a coarse environment (QTL3). Indeed, we can hypothesize that the PCA ordination fails at orienting the genetic space differentiation into the direction of environment 1 and environment 3 therefore leaving no chance to detect any outliers on the QTL influenced by these environmental "adaptively enriched genetic space" as performed $_{383}$ variables. Fig 3 shows that RDA has a much by Steane et al. (2014) on the same simulated 384 better behavior than pcadapt by taking advantage









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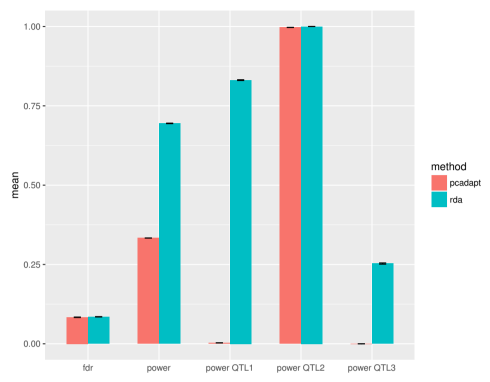


FIG. 4. Performance results of rda and pcadapt methods. Each performance value is averaged over 100 simulated dataset (error bars are displayed but hardly visible since they are very scarce). fdr, is the false discovery rate. power is given for the whole selected loci and seperately for loci coding for quantitative trait 1, 2 and 3.

of using informations of environmental local 402 conditions.

Results summarized on Fig 4 is confirming 404
that both methods have a good control of false 405
discovery rate (≈0.1) and that overall RDA shows 406
better performance at detecting true outliers since 407
it succeeds to detect quite often QTL1 and 408
QTL3 SNPs. It seems however less efficient at 409
detecting QTL3 outliers but this might be due 410
to the fact that local adaptation on a coarse 411
environment is more difficult that adaptation on 412
a smooth environmental gradient as environment 413
1 and 2. These simulations plead in favor of using 414
constrained ordination method instead of PCA 415
when non genetic data such as environmental 416
variable are available in order to orientate the axis 417
in the direction of informative gradients. 418

When performing an RDA on the "adaptively enriched genetic space", Fig. ?? and ?? show that the method succeed at detecting the relevant selective gradient and separating them on different axis at least on our simulations. This therefore serves as a proof of concept of Steane et al. (2014)'s approach to represent multilocus selective gradient and the possibility to use the ordination axis it to devise a metric that provides a holistic measure of genomic adaptation. Indeed, in RDA1 is strongly associated with envir2, RDA2 with envir1 and RDA3 with envir3 whereas poorly associated with the other axis. As expected, the correlated environment are also strongly associated with this respective axis. This is reflecting the fact that in reality it is difficult on an environmental gradient to distinguish









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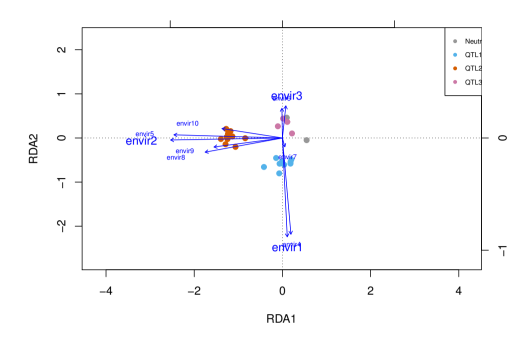


FIG. 5. RDA on the adaptively enriched genetic space. We discarded the individual points for readability. Dots represents outliers SNPs. R^2 of envir1 with the first, second and third axis is (0.02%, 77.5%, 14.5%), envir2 is (99.3%, 0.003%, 0001%) and envir3 is (0.009%, 0.82%, 64.7%)

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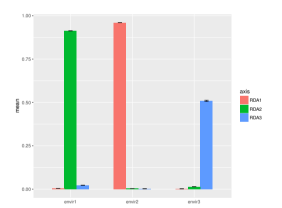


FIG. 6. \mathbb{R}^2 between envir1, envir2 and envir3 and each of the first three ordination axis. Values are averaged across the 100 simulated datasets.

among the covariable which one has a causal and effect on the individual fitness. However, it is often sufficient for biologists when performing an exploratory analysis to identify combination of environment variable having a strong association with adaptive variation without knowing precisely the underlying mechanical process.

Supplementary Material

Acknowledgments

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