RESEARCH ARTICLE

Replication of the principal component analyses of the human genome diversity panel [version 1; referees: 1 approved, 1 approved with reservations]

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Abstract

Background. In 2008, several principal component analyses (PCAs) applied on 660,918 single-nucleotide polymorphisms (SNPs) from 938 individuals from 51 worldwide populations of the Human Genome Diversity Panel were published by Li et al. PCAs were applied on subsets of individuals sharing a common geographic origin and showed that in several geographic regions, genome-wide variations of SNPs grouped individuals by populations in the two first principal components. In this study, we replicated the PCAs applied on two geographic subsets, first on individuals from Europe and second on individuals from the Middle East & North Africa. Methods. Quality control, feature selection, and PCA were applied on each geographic subset. The results were displayed on the two first principal components and compared to the original figures. Results. The replicated figures were found to match closely to the original figures. Conclusions. Therefore, the main results were replicated and can be independently reproduced by using publicly available data, source code, and computing environment.

This article is included in the Preclinical Reproducibility and Robustness gateway.
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Competing interests: Thomas Charlon, Alessandro Di Cara, and Jérôme Wojcik are employees of Quartz Bio S.A., Switzerland. The authors declare no competing interests related to this commercial affiliation. This does not alter the authors’ adherence to F1000Research policies on sharing data and materials.

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Introduction
Quartz Bio and the Stochastic Information Processing group are involved in the PRECISESADS project (http://www.precisesads.eu), which aims at reclassifying Systemic Autoimmune Diseases (SADs), a group of chronic inflammatory conditions characterized by the presence of unspecific autoantibodies in the serum and resulting in serious clinical consequences, based on genetic and molecular biomarkers rather than clinical criteria.

In order to use genetic similarities to deliver personalized treatments to patients affected by SADs as well as other diseases, it is important to first understand the genetic structures in healthy populations.

In 2008, Li et al. showed that although specific world regions have different genetic origins, all revealed population structures in principal component analyses (PCAs). Similar population structures were also observed in studies using other genome-wide variations datasets.

Li et al. applied PCAs on subsets of individuals from two geographic regions, Europe and the Middle East & North Africa, and displayed the results on the two first principal components in their article as Figures 2A and B, respectively, (with the latter labeled only Middle East).

In an attempt to replicate these two figures, we performed quality control, minor allele frequency filtering, tag SNP selection, and PCAs on both regional subsets of the SNP microarray data. The PCAs were then displayed on the first two principal components.

The replicated figures were found to match closely to the original figures, and therefore confirmed a successful replication.

Methods
Genotype data
The dataset consisted of two files: a zip file including the genotype data of 660,918 SNPs from 1,043 individuals with the annotations of the SNPs, and a text file composed of the annotations of 953 individuals (see Data and software availability).

The annotations of individuals were used to create two subsets of the data. The first contained 157 individuals from Europe and the second contained 163 individuals from the Middle East & North Africa.

Analysis sets
For each geographic region subset of the data, we verified that no individuals had missing value rates above 3% and excluded SNPs with missing value rates above 1%. An additive genetic model was then used to encode each A/B SNP (A/A = 0, A/B = 1, B/B = 2), which converts categorical SNP values to numerics by assuming that the effect of the A/B heterozygote and B/B homozygote are proportional to the number of B alleles. SNPs with minor allele frequency below 5% were excluded to remove rare variants, which are more prone to genotyping errors. In addition, in order to decrease the required computation time and memory usage, redundant SNPs were removed by applying TagSNP (r2 > 0.8, window of 500,000 base pairs). The missing values were imputed by random sampling of each SNP. Then each SNP was centered and scaled to unit variance. All steps were performed using the SNPClust R package v1.0.0.

For the Europe subset, a total of 375,164 SNPs from 157 individuals were selected for analysis. This defines our Europe analysis set.

For the Middle East & North Africa subset, a total of 412,979 SNPs from 163 samples were selected for analysis. This defines our Middle East & North Africa analysis set.

For comparison, the supporting online material of Li et al. reported that individuals with missing value rates above 2.5% and SNPs with missing value rates above 5% were excluded. Table S1 of Li et al. reports that 156 individuals from Europe and 160 from the Middle East & North Africa were used and the supporting online material reports that 642,690 SNPs were used.

Principal component analyses
PCAs were applied on the two analysis sets and displayed using the SNPClust R package v1.0.0. Principal component analysis (PCA) is a dimensionality reduction method, which projects SNPs by linear combination to maximize the variance on successive axes, i.e. principal components, while constraining the axes to be orthogonal.

The supporting online material of Li et al. reports that they first computed the Identity-by-State (IBS) matrix among the 938 individuals by using PLINK (version not provided) and then performed PCAs on the IBS matrix for each region separately. In this study, PCAs were applied on the analysis sets and not on IBS matrices.

Results
PCA of the Europe analysis set
The PCA of the Europe analysis set was displayed on the two first principal components (Figure 1). Individuals were grouped by population and the replicated figure matched closely with Li et al.’s Figure 2A.

The explained variance was almost identical, as the replication stated 2.1% in PC1 and 1.6% in PC2, while Li et al.’s Figure 2A stated 2.4% and 1.6%, respectively.

PCA of the Middle East & North Africa analysis set
The PCA of the Middle East & North Africa analysis set was displayed on the two first principal components (Figure 2). Individuals were grouped by populations and the replicated figure matched closely with Li et al.’s Figure 2B.

Two differences from Li et al.’s analysis were noted, first the Bedouin and Druze populations exhibited a larger spread on PC1 in the original figure. Second, one Bedouin individual was located with Mozabite individuals, which did not appear in Li et al.’s Figure 2B.
Figure 1. **Two first principal components of the Europe analysis set.** Visualization of the principal component analysis on 375,164 SNPs from 157 individuals from Europe. Individuals from North and South were differentiated in the first principal component and located in the lower and upper sides, respectively. Individuals from East and West were differentiated in the second and located in the right and left sides, respectively.

Figure 2. **Two first principal components of the Middle East & North Africa analysis set.** Visualization of the principal component analysis on 412,979 SNPs from 163 individuals from the Middle East & North Africa. Individuals from East and West were differentiated in the first principal component and located in the right and left sides, respectively. Individuals from North and South were differentiated in the second and located in the lower and upper sides, respectively.
The explained variance was slightly smaller, as the replication stated 3.1% in PC1 and 2.2% in PC2, while Li et al.’s Figure 2B stated 5.0% and 2.6%, respectively.

Discussion
The replicated figures matched closely to the original figures, although two differences appeared when examining the Middle East & North Africa subset: the smaller spread of two populations and the presence of an outlier.

Therefore, the main results were replicated and can be independently reproduced by using publicly available data, source code, and computing environment.

We successfully confirmed that although the two geographic regions studied had different genetic origins, both exhibited population structures in PCAs.

Understanding the genetic structure of healthy populations will enable us to use genetic similarities to deliver personalized treatments to patients affected by SADs. Using this replication, the PRECISESADS project will be able to compare clusters of patients affected by SADs to clusters of healthy individuals, independently from their ancestry-driven genetic structure.

Data and software availability
As stated in Li et al., the data sets are freely available online. Although the links that were provided are now outdated, the two data files are available from HGDP-CEPH: http://www.hagsc.org/hgdp/files.html (download link: http://www.hagsc.org/hgdp/data/hgdp.zip) and http://www.cephb.fr/en/hgdp_panel.php#serie2; ftp link: ftp://ftp.cephb.fr/hgdp_v3/hgdp-ceph-unrelated.out).

The PCAs were computed and displayed using the previously published R package SNPClust v1.0.0.

Computing environment in a Docker container is available from: https://hub.docker.com/r/thomaschln/reproducible-hgd

Source code required to generate this article and the definition of the corresponding computing environment, in which all required software are installed: https://github.com/ThomasChln/reproducible-hgd

Archived source code as at time of publication: doi, 10.5281/zenodo.345137

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Ethical statement
The data were previously published and approved by ethics committees. No samples were used and records were de-identified.

Author contributions
Conceptualization: JW SV; Formal analysis: TC; Funding acquisition: JW; Investigation: JW ADC; Methodology: TC JW; Project administration: JW; Software: TC; Supervision: JW SV; Validation: TC JW ADC; Visualization: TC; Writing - original draft: TC; Writing - review & editing: JW ADC SV.

Competing interests
Thomas Charlon, Alessandro Di Cara, and Jérôme Wojcik are employees of Quartz Bio S.A., Switzerland. The authors declare no competing interests related to this commercial affiliation. This does not alter the authors' adherence to F1000Research policies on sharing data and materials.

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References

Open Peer Review

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Version 1

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Michael G. B. Blum
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The authors replicate the ascertainment of worldwide population structure obtained by Li et al. (2008). They perform PCA to capture population structure. The PC axes closely match the ones obtained by Li et al.

However, the authors found that some Bedouin individuals don't belong to the population they should belong to. The authors should read and cite the 2 following papers that found related results


Additionally, I run the provided docker command (docker pull thomaschln/reproducible-hgdp) to reproduce the analysis but I don't find the generated results. The webpage (https://github.com/ThomasChln/reproducible-hgdp) should be improved and should include a more detailed tutorial.

References

Is the work clearly and accurately presented and does it cite the current literature?
Partly
Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Referee Expertise:** Population genetics, biostatistics, bioinformatics

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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**Referee Report 28 March 2017**

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This manuscript reports on the re-running of two PCA analyses presented in an earlier publication Li *et al.* 2008). The authors confirm the PCA results presented in the original paper and point out two minor differences.

The analysis looks solid and carefully executed. There are a few aspects that could be improved:

1. What I missed a bit was the justification why only the middle Eastern and European subsets were reanalysed. Also, the authors motivate their reanalysis so that they can use these individuals as controls for their PRECISESADS study. I was expecting the authors to go slightly further: do they have control samples? Where do they map on these PCA plots? If they match the location of those from the HGDP, I agree that it is an excellent indication to go further with their study cases. I think these points would further our understanding and go beyond the partial re-analysis of a published data and reporting identical findings.

2. Would be very helpful for the readers to see for every analysis step where did the authors use exactly the same tool as Li *et al.* and where do they differ? If at some point different tools were used, were the parameters set to be identical? How close was the pruned subset of SNPs when analysed by them and by Li *et al.*?
The title and abstract reflect well the study content. The methods and results are clearly explained, the data are available and the analysis is provided in full details in a Docker container. Study motivation could be better explained and the conclusions in terms of consequences for their future study could be more detailed.

**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.