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Applications Note

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Flashpca: Fast sparse canonical correlation analysis of genomic data

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Abstract

Summary: Sparse canonical correlation analysis (SCCA) is a useful approach for correlating one set of measurements, such as single nucleotide polymorphisms (SNPs), with another set of measurements, such as gene expression levels. We present a fast implementation of SCCA, enabling rapid analysis of hundreds of thousands of SNPs together with thousands of phenotypes. Our approach is implemented both as an R package flashpcaR and within the standalone commandline tool flashpca.

Availability and implementation: https://github.com/gabraham/flashpca

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 Introduction

Canonical correlation analysis (CCA) is a well-known statistical approach for multivariate analysis of two datasets (Hotelling, 1936). In the context of large-scale genomics and multi-omic data, CCA can prove useful in identifying relationships amongst complex data, for example single nucleotide polymorphisms (SNPs) and gene expression levels. Approaches that consider one SNP at a time together with multiple phenotypes have been shown to increase power to detect QTLs over the simpler but commonly utilised single-SNP/single-phenotype approach (Ferreira et al., 2009; Inouye et al., 2012), particularly when analysing correlated phenotypes that are modulated by the same genetic variants.

Analysis of multiple SNPs simultaneously is an attractive extension of the single-SNP multiple-phenotype approach, however, standard CCA is not well-defined when the number of samples is lower than the number of SNPs or phenotypes ($n < \min\{p,m\}$). One solution is Sparse CCA (SCCA) (Witten et al., 2009a,b; Parkhomenko et al., 2009), an L_1 -penalised variant of CCA which allow for tuning the number of variables that effectively contribute to the canonical correlation, thus making the problem well-defined. Owing to the induced sparsity, SCCA can be useful for identifying a small subset of SNPs and a small subset of the phenotypes exhibiting strong correlations. However, the rapidly increasing size and coverage of genotyping arrays (exacerbated by genotype imputation), together with the availability of large phenotypic datasets (transcriptomic,

metabolomic, and others; e.g., Bartel *et al.* (2015); The GTEx Consortium (2015)), makes it challenging to perform analyses such as SCCA using existing tools.

We have developed an efficient implementation of SCCA that is capable of analysing genome-wide SNP datasets (1 million SNPs or more) together with thousands of phenotypes, as part of the tool flashpca (Abraham *et al.*, 2014). The tool is implemented in C++ using the Eigen 3 numerical library (Guennebaud *et al.*, 2010), as well as an R interface (package flashpcaR) based on ReppEigen (Bates *et al.*, 2013).

Here, we compare the SCCA implementation in flashpcaR and flashpca with a widely-used implementation (PMA, by Witten *et al.* (2013)), and demonstrate the substantial improvements in speed of our tool, allowing for large analyses to be performed rapidly.

2 Methods

In standard CCA, we assume that we have two matrices \mathbf{X} $(n \times p)$ and \mathbf{Y} $(n \times m)$, measured for the same n samples. We further assume that both \mathbf{X} and \mathbf{Y} have been column-wise standardised (zero mean, unit variance). For a single pair of canonical variables a and b, CCA involves solving the problem

$$\underset{a,b}{\arg\max} \frac{a^T \Sigma_{XY} b}{\sqrt{a^T \Sigma_{XX} a \ b^T \Sigma_{YY} b}},\tag{1}$$

where Σ_{XX} and Σ_{YY} are the covariance matrices of X and Y, respectively, and Σ_{XY} is the covariance matrix of X and Y. The

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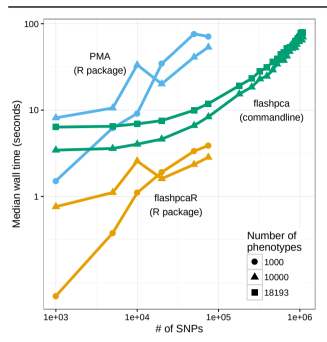


Fig. 1. Timing (median of 30 runs) of SCCA implemented in (i) the flashpcaR (R package) and (ii) flashpca (stand-alone commandline tool), compared with SCCA from PMA, using subsets of the HapMap3 dataset with gene expression levels as phenotypes.

solution is given by the singular value decomposition (SVD) of $\Sigma_{XX}^{-1/2}\Sigma_{XX}\Sigma_{XY}^{-1/2}$, with $a=\Sigma_{XX}^{-1/2}u_1$ and $b=\Sigma_{YY}^{-1/2}v_1$, where u_1 and v_1 are the first left and right singular vectors from the SVD.

SCCA is typically run on high-dimensional data, where a useful assumption is that the variables within ${\bf X}$ and ${\bf Y}$ are uncorrelated, i.e., ${\bf \Sigma}_{XX}={\bf \Sigma}_{YY}={\bf I}$ (Parkhomenko *et al.*, 2009), hence, a=u and b=v. Thus, SCCA involves solving a simplified form of CCA,

$$\underset{u,v}{\operatorname{arg\,max}} \quad u^T \mathbf{\Sigma}_{XY} v$$

s.t.
$$||u||_2^2 = 1$$
, $||v||_2^2 = 1$, $||u||_1 \le s_u$, $||v||_1 \le s_v$, (2)

where u and v are the left and right canonical vectors, and s_u and s_v are constraints on the L_1 norms of the canonical vectors.

The problem can be converted into the penalised (Lagrangian) form, making it solvable using iterative soft-thresholding (Parkhomenko et~al., 2009), which we employ here. Unlike standard CCA, SCCA is well-defined even when $n < \min\{p, m\}$, and induces sparse canonical vectors, depending on the choice of L_1 penalties (higher penalties lead to higher sparsity). The optimal set of penalties can be found via cross-validation: the data (both ${\bf X}$ and ${\bf Y}$) are split into training and test sets, SCCA is run on the training set $({\bf X}_{train}, {\bf Y}_{train})$ using a 2D grid of penalties, and the pair of penalties that produce the highest correlations in the test set, ${\rm Cor}({\bf X}_{test}u, {\bf Y}_{test}v)$, are selected. Optionally, a new model may be fitted to the entire data using these penalties.

3 Results

We utilised the HapMap3 phase III genotypes (International HapMap 3 Consortium, 2010), together with gene expression data of 709 individuals (Stranger *et al.*, 2012), with 1.4M SNPs and 47,000 gene expression probes (out of which 21,800 probes were analysed by Stranger *et al.* (2012) and used in our analysis). After quality control (see Supplementary Material) and taking the intersection of post-QC SNPs within each population, the data consisted of 709 individuals, 973,983 SNPs, and 18,379 gene expression probes.

We first confirmed that flashpcaR::scca produced models comparable with PMA::CCA, by comparing the results in crossvalidation on HapMap3 genotypes together with simulated gene expression levels (Supplementary Material). Next, to further assesss the relative speed improvement using real-world data, we used subsets of the HapMap3 genotypes and real gene expression levels (Stranger et al., 2012) and compared the runtime of: (i) PMA::CCA (R package), (ii) flashpcaR::scca (R package), and (iii) flashpca (commandline tool). Whereas both PMA::CCA and flashpcaR::flashpca are bound by the memory limitations of R, the commandline tool flashpca allows much larger analyses; hence we also ran larger analyses of increasing size: chromosomes 1-2, 1-3, ..., 1-22, up to all 973,983 SNPs. Figure 1 shows that flashpcaR::scca was $8-23 \times$ faster than PMA:: CCA, with an analysis of 75,000 SNPs and 10,000 gene expression levels completing in 3s and 54s, respectively. Note that runtime is not monotonic in the number of SNPs/phenotypes, since the number of iterations to convergence is affected by factors including LD structure. The commandline flashpca was faster than PMA::CCA as well, and completed an analysis of 709 individuals, 973,983 SNPs and 18,379 gene expression levels in median wall time of ~ 74 s (including all overheads). using $\sim \! 10 \text{GiB}$ of RAM. Note that cross-validation over a grid of penalties will increase these times considerably, in which case we recommend parallelisation over several cores (Supplementary Material).

4 Conclusion

flashpca provides a fast implementation of sparse canonial correlation analysis, making it possible to rapidly analyse high dimensional datasets. flashpca is available both as an R package, which enables analysis of metabolomic, transcriptomic, or any other quantitative set of measurements. For the case of large SNP datasets too large to fit in R, the commandline tool is available as well, enabling large QTL analyses of >1M SNPs and thousands of phenotypes.

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