Master of Science in Omics Data Analysis

Master Thesis

Bioinformatic tools for Big Data in Omic studies with application to genomic inversion calling and multi-omic data integration

by

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Application Note

Bioinformatic tools for Big Data in Omic studies with application to genomic inversion calling and multi-omic data integration

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Abstract

Motivation: The diversity and huge omics data take biology and biomedicine research and application into a big data era. Most of the current statistical analyses required to analyze omic data are not designed to deal with big data. Principal component analyses and multivariate methods to integrate multi-omic data are one of those examples. Therefore, having efficient and scalable functions are required to exploit the large amount of omic data which is currently available.

Results: We developed a library called BigDataStatMeth which includes functions to perform basic matrix operations and linear algebra for big matrices using HDF5 and DelayedArray Bioconductor’s infrastructure. We tested its performance by comparing the computational time with the one obtained with R base functions. Our results showed that our implementation outperforms existing functions and that the improvement increases when sample size is also increasing. This package can be the basis for implementing statistical methods required in omic data with large number of samples or features. As a proof-of-concept, we implemented PCA and Lasso regression within the same package and we also created another Bioconductor package, mgcca, which implements Generalized Canonical Correlation Analysis (GCCA) that is used in multi-omic data integration. We implemented an algorithm that allows the possibility of having missing individuals in one or more tables. The implemented methods have been used to analyze real omic data. We first used PCA to call genotype inversions of more than 400K individuals from UKBiobank. Then, data from TCGA was used to integrate multiple omic layers using GCCA.

Availability: Both packages are available at BRGE’s GitHub repository: https://github.com/isglobal-brge
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Supplementary information: We have four supplementary material files. One of them (Supp_Mat.pdf) includes supplementary information about the methods used in this work as well as suplementary tables and figures corresponding to the benchmarking. The other three files correspond to one vignete describing BigDataStatMeth package another for mgcca and a last one having a real data example to integrate multi-omic data using GCCA.
1 Introduction

The diversity and huge omics data take biology and biomedicine research and application into a big data era. Encouraged by constant cost reduction, data-sharing initiatives, and availability of public data large amounts of genomic, transcriptomic, and other omics data have become available ready to be analyzed. Therefore, omics data analyses require scalable and computationally efficient algorithms. These include methods to analyze a single omic dataset as well as methods to get an integrated view from the different samples and omic tables with the aim of getting an accurate and comprehensive view about the diseases and different biological processes (Subramanian et al., 2020).

Multivariate methods designed to analyze one or several high-dimensional datasets are widely used in omics data analyses. For instance, Principal Component Analysis (PCA) through Single Value Decomposition (SVD) has been used in genomics to address population stratification (Price et al., 2006)(Price et al., 2010a) or to genotype polymorphic inversions (Cáceres and González, 2015a). In transcriptomics and epigenomics SVD is used to estimate surrogate variables that is required to correct for batch effect (Leek et al., 2012) or to estimate cell proportions (Houseman et al., 2014)(Alquicira-Hernández et al., 2018).

On the other hand, Canonical Correlation Analysis (CCA) (Hotelling, 1936), Generalized Canonical Correlation (GCCA) (Kettenring, 1971), multi factorial analysis (MFA) (Abdi et al., 2013), co-inertia analysis (CIA) (Culhane et al., 2003) and Multi-Omics Factor Analysis (MOFA) (Argelaguet et al., 2018) have been used to perform the multivariate analysis of multiple omic tables (see Subramanian et al., 2020) and (Csala A, 2019) for a review of these methods applied to multi-omic data integration.

Bioconductor includes several packages that are designed to performed most of these multivariate methods such as BioSimular (Lun, 2020), PCAtools (Blighe and Aaron, 2020), SVA (Leek et al., 2019) for surrogate variables and omicade4 (Meng et al., 2014) which implements multiple co-inertia analysis. Most of these implementations, mainly those created to integrate multiple tables, were not designed to deal with big data sets, and hence, they are not computationally efficient. Another limitation that current multivariate methods have to integrate multi-omic data is how to handle missing data individuals. The presence of missing values in multi-omics data is inevitable since, in most cases, omic data are obtained in different time points and quality control can remove individuals form a single omic dataset. Most of the currently implemented methods works only with complete cases which is an underpowered approach (van de Velden and Takane, 2012). missRows is a Bioconductor library based on multi factorial analysis that addressed this issue (I and V, 2020). However, the analysis of large datasets cannot be performed using this method. Actually, the authors recommend to filter out those features with less variability to reduce the dimensionality (Voillet et al., 2016).

Most of the existing inefficient implementations are due to the fact that developers use base R functions to perform basic matrix operations or algebra. In order to overcome these difficulties and provide the user with efficient and scalable functions to implement any statistical method required to analyzed large omic dataset, we have developed BigDataStatMeth Bioconductor library that uses C++ language with Rcpp (Eddelbuettel and François, 2011)(Eddelbuettel and Balamuta, 2017) and RcppEigen (Bates and Eddelbuettel, 2013) from R-CRAN that provides an efficient tool for process and analyze omics data. BigDataStatMeth also works with HDF5 file format (Koranne, 2011) (Fischer et al., 2019) and Delayed Arrays (Pagis et al., 2020) directly from C++ using APIs developed from Bioconductor and other specific to C++.

The implemented algorithms also use parallel methods that will make our functions scalable using OpenMP - OMP (Dagum and Menon, 1998). This library will allow us to implement a new package called mgcca that will allow us to analyze multi-omic data using GCCA including a method that is designed to analyze data with missing individuals (Velden and Bijmolt, 2006). (van de Velden and Takane, 2012).

In order to demonstrate the usability and the good performance of our proposed method, a benchmark analysis is performed to compare the behavior of our functions with those implemented in base R. Our libraries also include a vignette where the use of the functions is shown in a practical way as well as a brief theoretical explanation. The practical use is illustrated using data from two public databases: the UK Biobank (UKB) (Sadlow et al., 2015) and The Cancer Genome Atlas (TCGA) (Tomczak et al., 2015). The UKB data is used to perform inversion calling in about 500K samples by using PCA. The TCGA dataset is used to illustrate how to integrate different omic data when each table have information on different individuals.

2 Methods

2.1 Databases

The Cancer Genome Atlas

The Cancer Genome Atlas (TCGA) (https://portal.gdc.cancer.gov), is a project, to catalogue genetic mutations responsible for cancer, using genome sequencing and bioinformatics. TCGA began as a three-year pilot in 2006 with an investment from the National Cancer Institute (NCI) and National Human Genome Research Institute (NHGRI). The TCGA pilot project proved that making the data freely available would enable researchers anywhere around the world to make and validate important discoveries. In our case, the TCGA data serves as an illustrative example of how to analyze big genomic datasets using our scalable algorithms. TCGA has one of the largest collections of multi-omics and clinical data sets for more than 33 different tumor types chosen because of their poor prognosis and availability of samples. The project contains molecular data from multiple types of assays including DNA and RNA sequencing, array-based expression and DNA methylation among others. Several pre-processed omic data tables are available for each tumour. Clinical data along with
molecular tables can be used to decipher the role of different omics in cancer survival, prognosis; to find biomarkers of treatment response; or to determine individuals with different multi-omics profiles that can be used in personalized medicine.

TCGA data was downloaded using TCGAutils package (Ramos et al., 2020). Gene annotations were made using the Bioconductor package biomaRt (Durinck et al., 2005) using the ensemble database (Yates et al., 2020) and the hsapiens_gene_ensembl dataset. CpGs annotations were made using 450K Human Illumina methylation dataset.

**UK Biobank**

UK Biobank (https://www.ukbiobank.ac.uk) is an international health resource supported by the UK National Health Service (NHS). UK Biobank aims to improve the prevention, diagnosis and treatment of a wide range of serious and life-threatening illnesses – including cancer, heart diseases, stroke, diabetes, arthritis, osteoporosis, eye disorders, depression and forms of dementia.

The UK Biobank is a prospective cohort of 502,536 adults aged between 40 and 69 years (229,182 men and 273,474 women). At recruitment, participants provided electronic signed consent, answered questions on socio-demographic, lifestyle and health-related factors, and completed a range of physical measures. They also provided blood, urine and saliva samples, which were stored in such a way as to allow many different types of assay to be performed (for example, genetic, proteomic and metabolomic analyses). Once recruitment was fully underway, further enhancements were introduced to the assessment visit, including a range of eye measures, an electrocardiograph test, arterial stiffness and a hearing test. The baseline information has been, and will continue to be, extended in several ways, for example, repeat assessments are planned to be conducted in subsets of the cohort every few years. Here we are focusing on a total of 488,377 individuals with European ancestry to whom invocation calling will be performed using PCA (Cáceres and González, 2015a).

### 2.2 Library Implementation

In omics data, we used to deal with large datasets with thousands of variables and a small/moderate number of samples. Currently, this paradigm has become even more challenging since we also have information for thousands of individuals. The analysis of this data requires a great amount of computational resources and optimized algorithms. Bioconductor is software project for the analysis and comprehension of genomic data generated by wet lab experiments in molecular biology. It is based primarily on the statistical R programming language, but does contain contributions in other programming languages.

The challenges of dealing with big data sets in Bioconductor are those found in R. By default, R runs only on data that can fit into your computer’s memory that is the biggest issue that researchers face when trying to use Big Data in R. Another big issue for doing Big Data work in R is that data transfer speeds are extremely slow relative to the time it takes to actually do data processing once the data has transferred. Finally, R is an interpreted programming language which means that it is not translated into machine language in a process prior to execution. R has a process that interprets the code in real time, this affects the efficiency at execution time and sometimes R code is not as fast as you would expect. Nevertheless, there are effective methods for working with big data in R that will allow the efficient and scalable implementation of omic data analyses. These include methods to: 1) program in low level language (C or C++), 2) work directly on disk and load in memory only the required data to be analyzed, and 3) implement parallel algorithms.

We have developed a Bioconductor package, BigDataStatMeth, based on previous methodologies that allows efficient and scalable computation of matrix operations and basic algebra required to implement statistical method in omic data analyses. First, in order to optimize the use of available resources for data processing, we have use Eigen (Guennebaud et al., 2010) which is a C++ template library for linear algebra, matrices, vectors, numerical solvers, and related algorithms using ReppEi gene package (Bates and Eddelbuettel, 2013). In conjunction with eigen we used LAPACK (Linear Algebra Package) which is a library for numerical linear algebra with low-level functions (Anderson et al., 1999). Second, in order to optimize memory usage, Hierarchical Data Format (HDF) (Forte, 1998) in its version 5 (HDF5) (Koranne, 2011) and Delayed Array data objects (Pagès et al., 2020) were used. HDF5 and Delayed arrays allow the effective management of extremely large and complex omics data collections including genomic, RNA-seq, methylation, copy number, mutations or microRNA among others. It also allows to deal with other type of data and metadata associated with an assay like clinical or pathological data. The link between Delayed Arrays and C++ functions have been performed using beachmat library. Finally, in order to speed up computation, OpenMP (Dagum and Menon, 1998) have been used to implement parallel algorithms. Section 1 in Supplementary Material provides a general overview of these methodologies including some figures describing two methods to parallelize matrix multiplication and single value decomposition (SVD) (Figures S1 and S2).

### 2.3 Omics data Analysis - Statistical methods

Methods implemented in BigDataStatMeth can be used to program efficient and scalable statistical methods required in omics data analysis. Principal Component Analysis (PCA) is one of the widely used methods in several omics data analyses. PCA can serve, not only as a dimensional reduction technique, but also to visualize cluster of individuals that are created given a hidden structure. For instance, PCA has been used for a long time in population genetics studies to produce maps summarizing human genetic variation across geographic regions (Menozzi et al., 1978). recently it can be used to explore the potential of disease identification in high dimensional blood microRNA data (SL et al., 2020) or to cluster subjects depending on their genomic inversion status (Cáceres and González, 2015b). In BigDataStatMeth we have implemented an scalable and efficient function to perform PCA using our parallel implementation of SVD that allows, among other, to visualize cluster of individuals given their
We developed a library called BigDataStatMeth which integrates functions to perform basic matrix operations and linear algebra for big matrices using HDF5 and DelayedArray Bioconductor’s infrastructure. This package can be the basis for implementing statistical methods required in omic data with large number of samples or features. As a proof-of-concept, we implemented PCA and Lasso regression within the same package. We also created another Bioconductor package, mgcca which implements GCCA to be used in multi-omic data integration allowing the possibility of having missing individuals in one or more tables.

**BigDataStatMeth library**

The methods described in previous section were used to implement BigDataStatMeth library that allow us to work in a efficient manner with omic data. BigDataStatMeth aims to be a practical, versatile and easy-to-use tool for researchers and omic data analysts. In omics data we can have different assays for the same set of samples with different omics data. For that reason, BigDataStatMeth allows the user to store different omic data in the same file. Additionally, BigDataStatMeth internally store all the results obtained with omics data in the same file as original data in an organized way. Figure 1 shows an example of how data is organized within BigDataStatMeth using a hdf5 file.

As stated in methods section, due to large amount of information associated with omic data, working with data blocks is a key issue. One of the biggest challenges in implementing BigDataStatMeth has been working with small blocks of data because: a) we have to take in to account at each moment the precise coordinates where we have to start to read and the exact block size to read; and b) not all operations can be performed in, since existing algorithms are highly complex. To the challenge of working in blocks we must add the complexity of working directly on files. It is possible when working with data in memory compute calculations directly with the complete dataset, but working with blocks of data from files only allows to have in memory the last read block. This requires to know the data we have on memory and which data we need to load or unload.
Fig. 2. Summary table with the functions implemented in BigDataStatMeth and the methods and types of resources used to be implemented.

compute calculations. This is needed to avoid overloading memory and worsening the overall performance of the function.

In BigDataStatMeth we have developed different functions to work with omic data. The implemented functions can be classified into five groups 1) basic functions with vectors and matrices; 2) linear algebra functions 3) methods for omics data analyses; 4) pre-processing data analyses; and 5) utilities to work with HDF5 data files that allows basic omics data organization. Figure 2 provides a detailed summary with all the implemented functions and methodologies for each group.

We have creates a reproducible vignette illustrating how to use BigDataStatMeth that can be downloaded from (https://github.com/isglobal-brge/BigDataStatMeth/blob/master/vignettes/). The vignette document explains in detail how BigDataStatMeth operates. The vignette also contains detailed examples using real datasets as well as some benchmarking to compare its performance with other existing approaches.

mgcca library

BigDataStatMeth can be easily used to implement any statistical method that requires any of the basic functions described in Figure 2 when analyzing omic data. As a proof-of-concept, we have implemented functions to integrate multi-omic data using GCCA as well as an algorithm that allow integrating multiple omic tables having missing individuals (van de Velden and Takane, 2012). The library can be found in (https://github.com/isglobal-brge/mgcca). The package also includes a vignette having a complete pipeline to integrate transcriptomic and epigenomic data from TCGA using MultiAssayExperiment object which is the default method to handle with multi-omic data in Bioconductor.

3.2 Real data analyses

We have applied the implemented methods in BigDataStatMeth and mgcca to analyze real omics data. We first used PCA to call genotype inversions of more than 400K individuals from UKBiobank. Then, data from TCGA was used to integrate multiple omic layers using GCCA.

PCA with UKBiobank omics dataset

The PCA implementation in BigDataStatMeth was used to analyze genome-wide SNP data. We used PCA to call polymorphic inversions in to well known inversions located at 8p23.1 and 17q21.31. We recall that a genomic inversion is a specific DNA interval that runs backward with respect to a reference genome and that it can be genotype from SNP data using bioinformatic tools based on PCA methodologies (Cáceres and González, 2015a)

Genomic data between coordinates 8,055,789 to 11,980,649 in chromosome 8 was obtained from UKB data. This data was downloaded in Genomic Data Structure format (GDS), and treated with gdsfmt library (Zheng et al., 2012)(Zheng et al., 2017).

BigDataStatMeth has functions to transform GDS data to hdf5 datasets. Initially, data for 1,591 SNPs and 488,377 samples were available. We perform a quality control step by removing those individuals having more than 10% of missing data (n=1411) letting the total samples in 486966. The remaining missing data were imputed using the observed allele frequency at each SNP. After
removing individuals with high percentage of missing values, we impute the rest of the missing values. Figure 3 shows the results of the calling procedure where a perfect clustering is observed.

The same procedure was applied for inversion at 17q21.31. In that case, we selected coordinates 43,661,775 to 44,372,665 from chromosome 17 that includes 462 SNPs. In this case, 15,147 samples were removed due to the large number of missing information letting the total samples in 473230. Figure 4 depicts the results for this inversion.

GCCA with TCGA data

GCCA method with missing individuals was used to analyzed data from TCGA. Our aim was to illustrate how to perform multi-omic data integration with GCCA. Bioconductor library curateCTCGADat (Ramos et al., 2018) was used to download the data, this library provides available data from TCGA as a MultiAssayExperiment object. We illustrate how to perform multi-omic data integration at a whole genome level and then, in a region of interest given by a specific genomic coordinate range that the researcher may be interested in.

We downloaded data from Adrenocortical carcinoma (ACC) which is a rare endocrine malignancy. In particular, we analyzed RNA-seq (Normalized) and Methylation data. One of our supplementary materials also available at ([https://github.com/isglobal-brge/mgcca]) contains a complete description of how to get this data. The downloaded MultiAssayExperiment encapsulates gene expression (RNA-seq) data for 79 samples and 20,501 genes, while methylation data has data from 80 samples and 485,577 CpGs. We first imputed missing values from methylation data (gene expression did not have any). To this end, we use impute function implemented in mgcca package that uses a knn algorithm with 10 neighbours. This function is a wrapper of impute package form Bioconductor adapted to MultiAssayExperiment objects. GCCA analyses revealed a total of 3,906 genes and 6,335 CpGs associated with either first or second global axis with a false discovery rate (FDR) lower than 1% (Figure 5 A and B). In order to interpret the axes we can project the scores of the individuals and color them using any illustrate variable. Figure 5 C depicts the individuals given their vital status. We are aware that this analysis would deserve another type of method (i.e. survival analyses) but we are using it as an illustrative example. We can observe as individuals who died are located in the top-right part of the figure. Therefore, features associated with those axes will be important for survival status. The top five genes related to survival are Protein Coding genes. Main genes are shown in the top right part of the Figure 5 A. These protein coding genes are the Lysine Demethylase 4B (KDM4B), the Poly (ADP-Ribose) Polymerase 2 (PARP2), the Nicalin (NCLN), the Autophagy Related 4D Cysteine Peptidase (ATG4D) and the RNA Exonuclease 1 Homolog (REXO1). Three of the five most statistically significant CpGs are located near a Protein Coding gene. These CpGs are cg00161225, cg00330929 and cg00256231 that are near Purinergic Receptor P2X 1 (P2RX1), Complement C1q Like 1 (C1QL1) and TBC1 Domain Family Member 16 (TBC1D16) respectively. The fourth CpG is cg00362657 located near pseudokinase PEAK3/C19orf35 and the last is cg00164949 with an unknown near gene. Section 3 (tables S1 and S2) in our Supplementary Material provides an extended annotated list about top significant genes and CpGs obtained with mgcca associated with either first or second global axis. Figure 5 shows the main results of this analysis.

Adrenocortical tumors occur as sporadic tumors, as part of the multiple endocrine neoplasia type 1 (MEN1), syndrome or as part of other hereditary disorders (Heppner et al., 1999), (Griniatsos et al., 2011), (Wang...
et al., 2019), Menin 1 gene (MEN1), is a tumor-suppressor gene located on chromosome 11q13 with genomic coordinates (11:64570986-64578766).

Therefore, researchers may be interested in performing analyses in that region to find new biomarkers of ACC. We use the MEN1 gene coordinates with 2kb upstream and downstream for subsetting features in both RNA-seq and methylation. We have data for MEN1 and mitogen-activated protein kinase kinase kinase kinase 2 (MAP4K2), and 104 CpGs. The GCCA analysis in that region ended up with 19 statistically significant CpGs close to MEN1 genomic coordinates that are associated with the two first global components (Figure 5 D and E). Figure 5 D depicts the CpG methylation specific immunoprecipitation in each individual. We can observe as individuals with CpG island methylator phenotype-low (CIMP-low) are located in the top right part and slightly extended to the left in figure. Individuals with CpG island methylator phenotype-intermediate (CIMP-intermediate) are located in the top right part and slightly extended to the bottom right. Finally, individuals with CpG island methylator phenotype-high (CIMP-high) are located in the left part. The significant CpGs related to CIMP-low and CIMP-intermediate are those CpGs depict mainly in right part of Figure 5 C and those significant CpGs related to CMP-high are those CpGs depict on left part in the figure. Annotated list of genes and CpGs can be found in section 3 (tables S3 and S4) in our Supplementary Material.

3.3 Benchmarking

We tested the performance of some of the functions implemented in BigDataStatMeth with respect to those implemented using the basic functions or even more advanced in R. To perform the benchmark, we use the microbenchmark function (Mersmann, 2019), a program or routine to measure and test the performance of a single component or task, this function is implemented in microbenchmark package available in CRAN. The device used for the benchmark was an iMac with a quad-core i5 processor (I5-6500) at 3.2GHz, 24Gb 1867 MHz DDR3 of RAM and a fusion disk drive (hybrid drive that combines a hard disk drive with a NAND flash storage).
FirstAuthorLastName et al.

Fig. 6. Graphic performance for different functions implemented in BigDataStatMeth and R. Panel A compares matrix multiplication times using BigDataStatMeth’s functions that uses C++ with basic R multiplication function and crossproduct function implemented in R. Panel C compares the computing time using mgcca function implemented in R and in BigDataStatMeth. The multiplications with R were carried out until reaching the data size of 4500x4500, the operations implemented in BigDataStatMeth continued to be applied until reaching a data size of 7500x7500. It was observed that the execution time with BigDataStatMeth was still much lower than that obtained with the R functions.

Figure 6 compares the performance for some of the implemented method. More results can be found in Benchmarking in Section 4 in our Supplementary Material. In general, our implementations outperformed others available in R. We can also observe that the improvement increases when data dimension increases.

4 Conclusion

Omics technologies are bringing a revolution in transforming the medicine and the health care sector, especially with regard to personalized medicine. This work is only a very basic approximation of the tool that can be developed to support omics data analysis, and to advance in the field of personalized medicine where are needed tools capable of analyzing big data efficiently and accurately in a few seconds. With methods and technologies applied in BigDataStatMeth it has been seen that there are important improvements in terms of performance and system resource management that can help in personalized medicine to obtaining the results derived from omics data analysis with effectively and accurately results.

Considering that BigDataStatMeth is a scalable library, future work would go through creating more functionalities adapted to new requirements and methods in omics data analysis field that allow progress in biomedicine and the personalized medicine. Some of these features would be a) managing the missing data in different ways in order to prevent distorted results in omics analysis, b) creating more statistical methods able to analyze the distinct omics and multi-omics datasets from distinct perspectives and c) generating complex plots from omics analysis to help the scientific and medical community to understand the complex underlying biology.

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