Two criterions to benchmark batch effect correction algorithms with an application to the integration of Baseline RNASeq datasets from Expression Atlas

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11/06/2019

I present here two different methods to be used in the context of the integration of datasets from different sources, subject to batch effect. I extracted one of them from an existing paper and I introduce a new one in this paper. Although both of them are based on Principal Components Analysis, the first one has what I would call a "sample-based" approach while the second one focuses more on the information given about genes.

The final aim of this work is the creation of several Expression Atlases by merging the RNASeq datasets available on the existing Expression Atlas. The building of such atlases shall be done using heterogenous datasets which can definitely not be supposed to follow the same distribution. Therefore many batch effect correction algorithms can already be considered as irrelevant for this problem (batch-mean centering to name but one, as well as every method which doesn't take biological factors into account).

The data: Mouse Expression Atlas

In this section, I show a typical workflow of data preparation before integration.

```
#relative path to the data of all the experiments
batch data<- "Mouse Expression Atlas/batch data/"
#list of experiments
experiments<-list()
for(filename in dir(batch_data)){
  experiments[[filename %>% str_split("-") %>% unlist %>% extract(2:3) %>% paste(collapse="")]] <- get(
}
#tissues investigated in each experiment
tissues<-experiments %>% map(~.$organism_part)
#matrix of intersections between batches
intersections<-NULL
for(i in tissues %>% seq_along){
  for(j in tissues %>% seq_along){
    intersections%<>%c(length(intersect(tissues[[i]],tissues[[j]])))
  }
intersections %<>% matrix(length(tissues)) %<>% set_colnames(names(tissues)) %<>% set_rownames(names(ti
library(igraph)
intersections %>% graph_from_adjacency_matrix %>% plot
```

As three experiments are isolated, one shall not try to integrate them, as it is not possible to correct batch effect with these experiments (since it is not possible then to dissociate batch effect from biological variations).





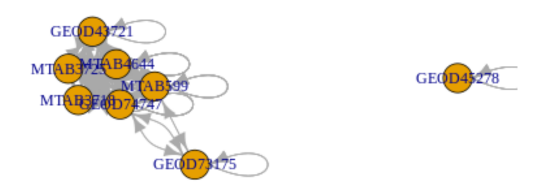


Figure 1: "Graph of intersections between the experiments"

```
experiments[c('GEOD45278','GEOD44366','ERAD169')]<-NULL
```

After these steps, the package eigenangles can be used to integrate the datasets. The datasets are merged within a unique SummarizedExperiment object. If one sets method="none", no batch effect correction is applied. However the count matrix is log-transformed by default (it is possible not to do this transformation by setting log=FALSE within the call to integrate.experiments).

```
library(eigenangles)
integrate.experiments(list=experiments, method="none") -> all
all
```

```
## class: SummarizedExperiment
## dim: 1645 119
## metadata(1): batch
## assays(1): corrected_log_counts
## rownames(1645): ENSMUSG00000000056 ENSMUSG00000000131 ...
## ENSMUSG00000106536 ENSMUSG00000106617
## rowData names(0):
## colnames(119): SRR652245 SRR652246 ... ERR032237 ERR032238
## colData names(16): AtlasAssayGroup sex ... strain_or_line batch
```

One can apply the available batch effect correction methods to integrate these experiments, such as:

• ComBat:

```
integrate.experiments(list=experiments, method="combat", model = ~organism_part) -> combat
```

Standardizing Data across genes

• RUVs:

```
integrate.experiments(list=experiments, method="ruv", model = ~organism_part) -> ruv
```

• MNN:

```
integrate.experiments(list=experiments, method="mnn", model = ~organism_part) -> mnn
```

Please note the presence of a model argument, that gives the biological differences between the samples, to be taken into account during batch effect correction. Here it is a formula ~organism_part referring to the so called column in the colData() fields of all the experiments.

A sample-based criterion

I introduce here the guided PCA which comes originally from an article by Sarah Reese [A new statistic for identifying batch effects in high-throughput genomic data that uses guided principal component analysis]. The eigenangles package has an implementation of this concept in order to compare the different batch effect correction algorithms.

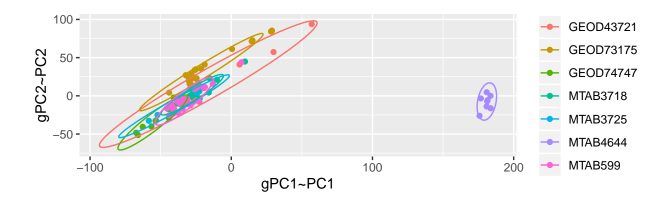


Figure 2: First plan of variance of the guided PCA of the uncorrected dataset

The idea of guided PCA is to perform PCA on a batch-aggregated dataset. All the samples from the same batch are averaged or summed to form one sample and PCA is performed on these new samples (one sample per batch). This PCA yields to some geometrical axes, that we will called guided principal components, different from the ones that standard PCA would give. The original samples (not aggregated) are then projected on these new directions so that the parts of variance of these axes can be estimated.

The guided principal components are found in a way that they represent the directions of batch effect. If these directions are important compared to standard principal components, i.e. if their variance is comparable to the ones of low rank principal components, it would mean that batch effect is important.

To do this analysis, the function gPCA from package eigenangles gives two quantities: - the delta statistics, which are the ratio of variance of the guided principal subspaces and the standard principal subspaces. This quantity is necessarily less than 1, as the standard principal subspaces maximises the variance of the dataset when projected on a subspace of the same dimension. The definition of the delta statistics is denoted this way:

$$\delta_k = \frac{\mathbb{V}gPC_1 + \ldots + \mathbb{V}gPC_k}{\mathbb{V}PC_1 + \ldots + \mathbb{V}PC_k}$$

• the ranks of variance of the guided principal components, defined for a guided principal component as the greatest rank such that the standard principal component of this rank has greater variance than the considered guided principal component.

The eigenangles package provides methods to visualise these quantities after running gPCA function on the dataset.

Let us see the gPCA of the not corrected dataset, to evaluate the importance of batch effect within it:

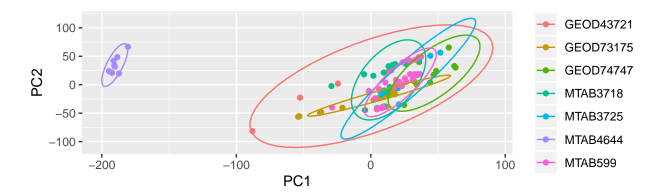


Figure 3: First plan of variance of the standard PCA of the uncorrected dataset

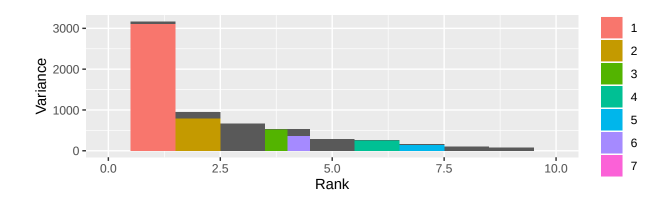


Figure 4: Ranks of variance of the guided principal components of the uncorrected dataset

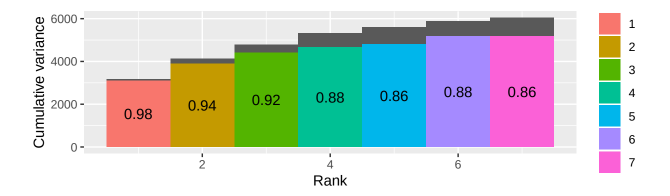


Figure 5: Delta statistics shown with the cumulative parts of variance of both guided and unguided principal components of the uncorrected dataset

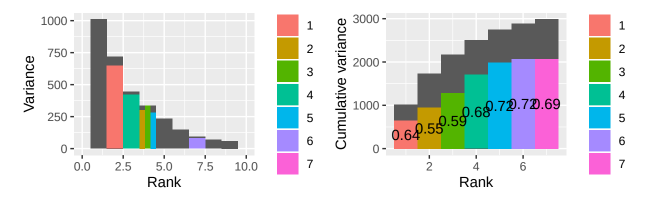


Figure 6: gPCA statistics of the dataset after ComBat correction

```
all.gpca %>% plot("delta")
```

We will now apply this tool to the different corrected datasets, in order to benchmark the algorithms:

A new gene-based criterion

I introduce here a new method to evaluate the importance of batch effect within an integrated dataset. This method is based on some geometric consideration on the principal components of the single datasets compared to those of their merger (with or without correction). As principal components represents geometrical directions, a way to compare them is to estimate the angle between them.

In the simple case where datasets are supposed to have the same distribution, we expect actually the single datasets to have their principal components similar between them and to those of the merged dataset.

As the coefficients of principal components are akin to weights on the genes and represent somehow their involvement in the variance of a dataset, they are a good summary of the information provided by a dataset on the genes. Thus low angles between the respective principal components of each dataset and the ones of their merger means somehow that the information provided by the single datasets has been conserved through their integration.

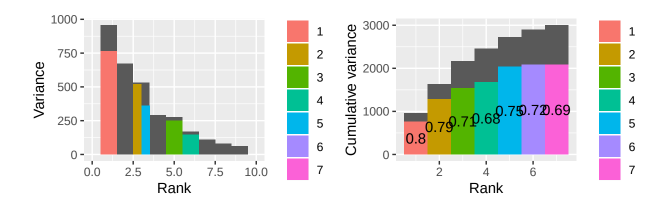


Figure 7: gPCA statistics of the dataset after RUV correction

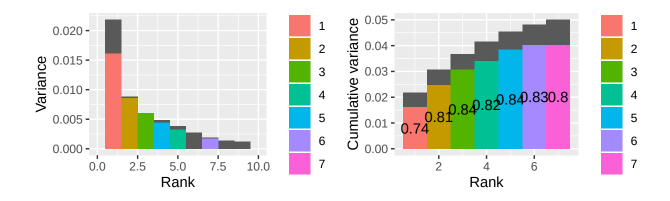


Figure 8: gPCA statistics of the dataset after MNN correction

Mathematical consideration for the calculation of angles

To calculate an angle between two vectors \vec{u} and \vec{v} in a space of any dimension, the most commonly used definition is :

$$\widehat{(\vec{u}, \vec{v})} = \arccos \frac{\langle \vec{u}, \vec{v} \rangle}{\|\vec{u}\| \|\vec{v}\|}$$

where $\langle \vec{u}, \vec{v} \rangle$ denotes the euclidian inner product between vectors \vec{u} and \vec{v} , and $\|.\|$ denotes the euclidian norm

To calculate angles between the first principal component of each dataset and the integrated one, this definition can be used easily as PC1 are 1-dimensional direction. \vec{u} shall be chosen as an orientation vector of PC1 of the considered individual dataset and \vec{v} as an orientation vector of PC1 of the integrated dataset. In R, orientation vectors of principal components are given by the columns of the \$rotation element in the output of a call to prcomp function.

Although this definition doesn't allow to extrapolate this idea to principal components with higher ranks. Indeed, estimating angles between principal components of higher rank doesn't make sense contrarily to angles between PC1, as only PC1 maximises the variance of the dataset projected on a 1-dimensional axis. Principal components of higher rank don't have such properties by themselves. However the plane generated by PC1 and PC2 maximises the variance of the dataset projected on a 2-dimensional subspace, in just the same way as the n-dimensional subspace generated by PC1, ..., PCn is such that the variance of the dataset projected on such a subspace is maximised.

Therefore the good generalisation of this idea is to compute the angle between the n-dimensional subspaces $\operatorname{span}(PC_1^i,...,PC_n^i)$ and $\operatorname{span}(\dot{PC}_1,...,\dot{PC}_n)$. This requires to be able to calculate angles between subspaces, whereas the previous definition only gives a way to calculate angles between vectors and therefore only between 1-dimensional subspaces.

Thus we give the following definition for the angle between two subspaces $U = \text{span}(\vec{u}_1, ..., \vec{u}_n)$ and $V = \text{span}(\vec{v}_1, ..., \vec{v}_m)$, parts of a space of dimension p = m + n and where $(\vec{u}_1, ..., \vec{u}_n)$ and $(\vec{v}_1, ..., \vec{v}_m)$ are orthonormal bases of those subspaces respectively:

where the coordinates of the vectors are given in an orthonormal basis of the space. If one disposes of non-orthonormal bases for U and V, one can use any orthogonalisation process, such as any QR-factorisation method, in order to apply the previous formula legitimately.

Here there is a constraint on the dimension of data, given above by p = m + n, due to application of determinant operator, only defined for a square matrix. In our problem, the dimension p of data is the number of genes considered. Although, still in our problem, one wants to calculate angles between two n-dimensional subspaces for any value of n, so that the condition p = m + n = 2n shall not be satisfied in general.

However, this is not a real issue as the 2n base vectors $\vec{u}_1,...,\vec{u}_n,\vec{v}_1,...,\vec{v}_n$ are themselves situated in a 2n-dimensional subspace, where determinant can be applied as well as in the original p-dimensional space. The issue is then to rewrite the problem in this particular subspace i.e. $U+V=\mathrm{span}(U\cup V)$, using an orthonormal basis of this subspace. One can easily find such a basis by performing a QR-factorisation to the family of vectors $(\vec{u}_1,...,\vec{u}_n,\vec{v}_1,...,\vec{v}_n)$ (ordered as columns in a matrix) where the R matrix contains the coordinates of the original vectors in this new basis.

Thus we adopt the following framework to calculate angles between the subspaces span $(PC_1,...,PC_n^i)$ and span $(\dot{P}C_1,...,\dot{P}C_n)$ for any rank n and any batch i:

- Apply QR-factorisation to the family of p-dimensional vectors $(PC_1^i,...,PC_n^i,\dot{PC}_1,...,\dot{PC}_n)$ to find their coordinates (given in the R matrix) in an orthonormal basis of their 2n-dimensional subspace. Thus we get 2n new vectors of coordinates (although they represent geometrically the same vectors) whose dimension is also 2n, that is to say a square matrix to which determinant is applicable. Let's denote $(\widetilde{PC}_1^i,...,\widetilde{PC}_n^i,\widetilde{PC}_1,...,\widetilde{PC}_n)$ these new vectors of coordinates.
- Apply QR-factorisation to the family $(\widetilde{PC}_1^i,...,\widetilde{PC}_n^i)$ to get an orthonormal basis of its span (given in the Q matrix) : $(\widetilde{PC}_1^{i\perp},...,\widetilde{PC}_n^{i\perp})$. Do the same with the family $(\dot{PC}_1,...,\dot{PC}_n)$ to obtain an orthonormal basis of its span : $(\widetilde{PC}_1^i,...,\widetilde{PC}_n^i)$
- Hence the angle between the subspaces $\operatorname{span}(PC_1^i,...,PC_n^i)$ and $\operatorname{span}(\dot{PC}_1,...,\dot{PC}_n)$ is given by :

$$\alpha_n^i = \arcsin \det(\widetilde{PC}_1^{i\perp},...,\widetilde{PC}_n^{i\perp},\widetilde{PC}_1^{i\perp},...,\widetilde{PC}_n^{i})$$

Discussion on the scaling step before PCA

 \dots to be completed

Benchmark of the correction algorithms using eigengenes angles

```
par(mfrow=c(1,2))
plot(
   uncorrected=all.angles,
   ComBat=combat.angles,
   RUV=ruv.angles,
   MNN=mnn.angles
)
```

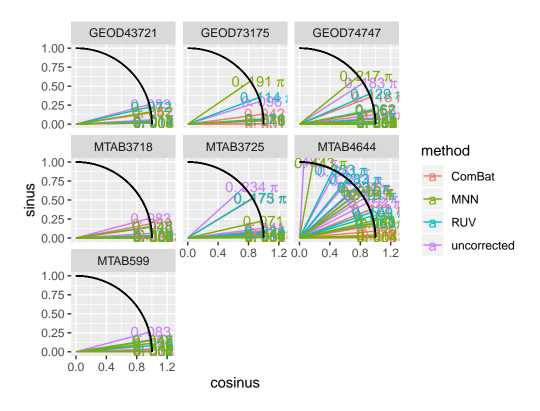


Figure 9: Eigengenes Angles for all corrections

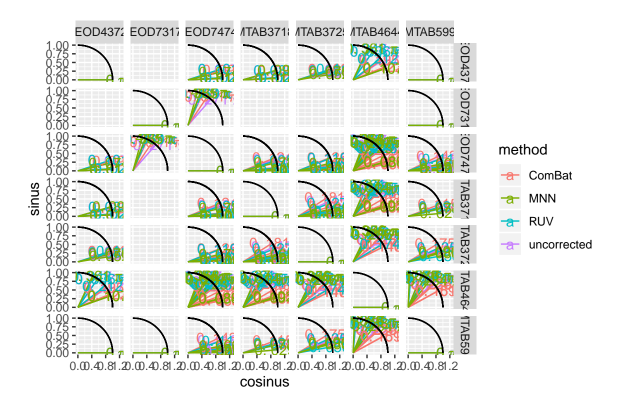


Figure 10: Eigengenes Angles for all corrections