The IntClust Package: Vignette

Marijke Van Moerbeke

$\mathrm{May}\ 12,\ 2015$

Contents

1	Intr	roduction	2
2	Dat	ta	2
3	Prin	mary Analysis	2
	3.1	Single Source Clustering	2
	3.2		3
		3.2.1 Aggregated Data Clustering (ADC)	4
		3.2.2 Aggregated Data Ensemble Clustering (ADEC)	4
		3.2.3 Complementary Ensemble Clustering (CEC)	4
		3.2.4 Similarity Network Fusion (SNF)	5
		3.2.5 Weighted Clustering (WeightedClust)	5
		3.2.6 Weighted Similarity Clustering (WeightedSim)	6
		3.2.7 Weighting on Membership Clustering (WonM)	6
	3.3	Comparison of the results	6
	3.4	Selection of a Specific Cluster	12
4	Seco	condary Analysis	14
	4.1	Differential Gene Expression	14
	4.2	Pathway Analysis	15
	4.3	Characteristic Features	16
5	Soft	tware used	18

1 Introduction

Discovering the exact activities of a compound in drug development is important. A single data source only provides insight into the behavior on one layer of the underlying biology. Therefore it is encouraged to rely on multiple data source simultaneously. The R package IntClust is the result of an integrated data analysis project where several multi-source clustering procedures have been investigated. Further, differential gene expression, pathway analysis and visualization tools for the results are available. This vignette is a user guide for the package but does not show all possible functions and tricks. It contains a general overview, several examples and is accompanying the helpfiles of the IntClust package.

2 Data

First, we load the package and the included data set. The data set is the publically available CMAP MCF7 data set containing 56 compounds. Further, information on 250 fingerprints, 477 target predictions and gene expression of 2434 genes is included. The fingerprint and target prediction matrices are the available data sources of the MCF7 data and are both binary. It is noted that the number of data sources is not limited and that any type (binary, continuous, discrete) of data can be used. It is important to check that the compounds are in the rows of the data sources and in the columns of the gene expression matrix.

```
> library(IntClust)
> data(fingerprintMat)
> data(targetMat)
> data(geneMat)
```

3 Primary Analysis

The primary analysis consist of performing several clustering, single and mult-source, techniques. The single clustering results will reveal whether or not there is a high degree of resemblance between the data sources. If multi-source clustering are executed, the results can be compared and the influence of each data source can be seen.

3.1 Single Source Clustering

Clustering can be performed on each data source separately with the *Cluster* function. The available distance measures are euclidean for continuous data and jaccard or tanimoto for binary data. The function can be easily extended to contain more distance metrics. The implemented method for clustering is agglomerative hierarchical clustering (Hastie et al. (2009)) as coded by the *agnes* of the cluster package with the ward link. The option for normalizing the data is readly available. Normalization is performed with the Normalization function and can be called on separately. The normalizing methods are: Quantile-Normalization, Fisher-Yates Normalization, standardizzation and range normalization. See the helpfiles for more information. No normalization is necessary for the example since both data matrices are binary and their distances will all fall between zero and one.

If the number of clusters to work with is unknown, the option gap in the *Cluster* function can be put to TRUE. Relying on the gap statistic (Hastie et al. (2009)), an optimal number of clusters will be determined. Another option is to determine the number of clustes with Selector Clusters. Medoid clustering is performed for a sequence of numbers of clusters for each provided data source. The number correspinding with maximal average silhouette widths over the data sources can be taken as an optimal number of clusters.

```
> List=list(fingerprintMat,targetMat)
```

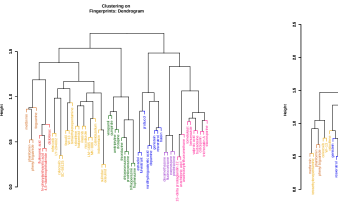
```
silhouette-optimal number of clusters: 20 silhouette-optimal number of clusters: 10 silhouette-optimal number of clusters: 20
```

In the specific case of these data sources, the average silhouette width will only increase as the number of clusters increases. Therefore, we will rely on the gap statistic and conclude on 7 clusters.

Colors are an important aspect of the visualization tools. The function *ColorPalette* provides us with the HEX codes of shades if colors and a desired number of colors are given. It is necessary to provide a color for each cluster and in some cases extra colors might be necessary.

One method to plot clustering results is to plot their dendrograms. A function called *ClusterPlot* is available to plot the results from the *Cluster* and any other clustering function. The option is available to color the leaves according to another clustering results. An example is presented for further clarification.

- > ClusterPlot(Data1=MCF7_F,Data2=MCF7_F,nrcluster=7,cols=Colors,main="Clustering on Fingerprints: Dendrogram",ylim=c(-0.1,1.8))
- > ClusterPlot(Data1=MCF7_T,Data2=MCF7_F,nrcluster=7,cols=Colors,main="Clustering on Targets: Dendrogram",ylim=c(-0.1,2.5))



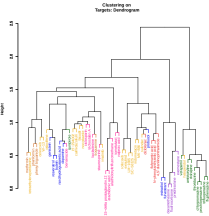


Figure 1: FP - MCF7

Figure 2: TP - MCF7

Figure 10 on the left is the results of the fingerprint clustering of the MCF7 data colored by the result of the fingerprint clustering. This implies that the dendrogram was cut into 7 clusters and each cluster was given a color. On the right in figure 2 the result of the target prediction clustering is depicted. The dendrogram was also cut into 7 clusters but each leaf (each compound) was given the same color it had in the fingerprint clustering to make a comparison possible. We could for example investigate the green cluster determined by figure 10. It is seen that in figure 2 the cluster does not undergo a lot of changes under the influence of the target predicions. The group is reformed but one compound has dissapeared and has been replaced by a compound of the what was the yellow cluster in figure 10.

3.2 Multi-Source Clustering

Several multi-source clustering procedures have been implemented in the IntClust package. A brief overview is provided here.

3.2.1 Aggregated Data Clustering (ADC)

Aggregated data clustering can only be applied if all data sources are of the same type. This is because the first step is to fuse all data matrices into one larger matrix such that in the end only one data matrix remains. Next, clustering is performed on this single matrix.

3.2.2 Aggregated Data Ensemble Clustering (ADEC)

Aggregated data ensemble clustering starts as ADC with the fusion of the data matrices can thus only be performed if all data matrices are of the same type. Then ensemble clustering is performed on the fused data matrix. This comes down to repeatedly applying hierarchical clustering. The original version of the method (Fodeh et al. (2013)) is implemented as version ADECa and two adaptions were made as ADECb and ADECc. A more detailed explanation follows.

Call m the number of features of the fused data matrix A.

- 1. In every iteration, a random sample of features r of A is taken and form matrix A. The number of features is randomly set between $\frac{m}{2}$ and m-1 each time.
- 2. Hierarchical clustering is performed on A'. The dendrogram is cut into a specific number of clusters k.
- 3. The incidence matrix C is computed. This binary matrix has as rows and columns the objects of the data set. A value of 0 indicates that these objects belong to the same cluster while 1 implies that they do not. This seems counterintuitive but was done to ensure distances rather than similarities.
- 4. The co-association matrix S is iteratively computed as

$$S^{(t+1)} = S^{(t)} + C$$

and indicates the number of times a pair of objects belong to the same cluster.

- 5. The steps in 1-4 are repeated t times.
- 6. Finally, hierarchical clustering is performed on the resulting co-association matrix S which yields the result of the ensemble clustering.

This is the description of ADECa. In version b a random sample of features will not be taken but instead all features will be used in every iteration. Variation is ensured by cutting the resulting dendrogram multiple times for a range of values for k. Version c is a combination of version a and b. Every iteration a random sample of features will be taken and each resulting dendrogram will be cut into cluster several times for a range of values for k.

```
> L=list(fingerprintMat,targetMat)
```

```
> MCF7_ADECa=ADECa(List=L,distmeasure="tanimoto",normalize=FALSE,method=NULL,
t=20,r=NULL,nrclusters=7,clust="agnes",linkage="ward")
```

- > MCF7_ADECb=ADECb(List=L,distmeasure="tanimoto",normalize=FALSE,method=NULL, nrclusters=seq(5,25,1),clust="agnes",linkage="ward")
- > MCF7_ADECc=ADECc(List=L,distmeasure="tanimoto",normalize=FALSE,method=NULL, t=20,r=NULL, nrclusters=seq(5,25),clust="agnes",linkage="ward")

3.2.3 Complementary Ensemble Clustering (CEC)

Complementary ensemble clustering (Fodeh et al. (2013)) shows similarities with ADEC. Instead of merging the data matrices, ensemble clustering is performed on each data matrix separately as described for ADEC. After the iterative procedure, the co-association matrices are combined via a weigted linear combination into a final weigted co-association matrix. On this final matrix clustering is performed once more which yields the final result. The version CECa, CECb and CECc differ respectively from each other as where ADECa, ADECb and ADECc differ.

3.2.4 Similarity Network Fusion (SNF)

Similarity network fusion (Wang et al. (2014)) consists of two stages and results in a sharing of information over all available data sources.

Stage I - The initial step A similarity network is set up for each data matrix. First, the distances between the objects are calculated. These are then rescaled to similarities relying on the kernel of normal distribution. This results in a matrix with a similarity value for each pair of objects. The similar objects were found, the higher their similarity value. The connection to a network is made when the matrix is visualized as weighted graph with the objects as nodes and the pairwise similarities as weights on the edges.

Stage II - The network fusion step To perform a fusion of the networks, two more matrices are necessary. The first is a normalized version of the similarity matrix and is referred to as a global matrix. The second is also a normalized version of the similarity matrix but only considers the points in the vicinity of objects. This implies that the normalization is performed for a specific number of neighbours and all other values in the row are set to 0. This matrix is called the local matrix. The assumption is that objects in the neighbourhood bring more reliable information forward than objects further away. Each global matrix is now updated via a matrix multiplication of the corresponding local matrices and the average of all other global matrices. After a number of iterations, the average of the updated global matrices is taken as the final fused network. The similarities are converted to distances and a final clustering is performed.

The R package SNFtool accompagnies the paper by Wang et al. (2014). Its functions are combined into SNFa. However, it was noticed that the functions do not coincide with the method described in the paper. Therefore SNFb was written which performs the steps as these were outlined. Versions SNFc shows a high degree of similarity with version b but instead of performing the normalization of the local matrix separately, the local matrix is taken as a submatrix of the global matrix.

3.2.5 Weighted Clustering (WeightedClust)

Weighted clustering is a straigtforward method. From each data matrix, a distance matrix ic calculated. The distance matrices are than combined via a weighted linear combination into one weighted distance matrix on which clustering is performed.

3.2.6 Weighted Similarity Clustering (WeightedSim)

Weighted similarity clustering (Perualila-Tan et al. (in press)) is practically the same as weighted clustering but an optimal weight is determined first. Each single source clustering is compared to every weighted clustering and the jaccard index is calculated as a representative of their similarity. For every weight combination, each pair of jaccard indices is compared to each other by takin their ratios. The weight combinations for which the ratios is closests to 1 is taken as the optimal combination.

An optimal weight can alternatively be determined by the function DetermineWeight_SilClust. For each provdided weight, the distance matrices are combined in a weighted combination and medoid clustering with the pam function is performed. Based on the silhouette widths and the cluster membership an optimal weight is determined. See the helpfiles for more details.

3.2.7 Weighting on Membership Clustering (WonM)

Weighting on membership performs hierarchical clustering on each data source separately. With k the number of clusters, the resulting dendrograms are cut into clusters for a range of values for k. Each time, an incidence matrix is set up. This is a matrix with as rows and columns the objects of the data set. Its values are 0 and 1 with 0 indicating that a pair of object resides in the same cluster to ensure distances. The incidence matrices are first summed over all values of k per data source and then these of the different data sources are added as well. On the resulting consensus matrix, hierarchical clustering is performed once again to obtain the final clustering taking into account information of all data sources.

3.3 Comparison of the results

The clusters of the multi-source methods consist of objects that are similar on multiple layers of the underlying biology represented by the multiple data sources. The interest lies in those clusters that remain stable over the applied methods. If a cluster does not undergo too many changes and is found multiple times, confidence can be put into those objects belonging together. Further, it can be hypothesized that the used data sources are related for those clusters. This can provide more insight into the mechanism of action of the compounds.

A comparison of the results is not done on sight. Different methods cluster the compounds in

a different order and this results in non-corresponding cluster numbers. Therefore, it was decided to take one method as reference and rearrange the cluster numbers of the other results to this reference. The re-appointing of the cluster numbers is based on finding the cluster that relatively has the most in common with one of the reference clusters and taking over this number. The function MatrixFunction was written for this purpose and its algorithm is partly based on the Gale-Shapley algorithm (?).. It creates a matrix of which the columns are the compounds in the order of clustering by reference method. The rows are the different methods and the values of the cells are the rearranged cluster numbers. If each value is associated with a colour, a visualization of the matrix can be made. A similarity measure is handy for the comparison of multiple methods. Given a method to be used as reference, it is observed which compounds are residing in the same cluster for a second method after applying MatrixFunction. The number of compounds that belong to the same cluster is summed and divided by the total number of objects in the data set. This is computed by the SimilarityMeasure function.

The MatrixFunction and SimilarityMeasure functions are combined in the ComparePlot function.

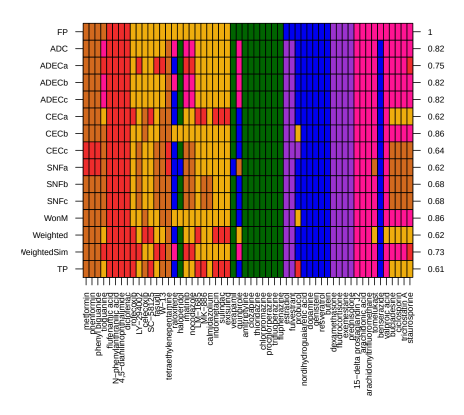


Figure 3: Comparison of the results for the CMAP MCF7 data

Figure 3 shows a comparison of all executed clustering procuderes with the fingerprint clustering as a reference. All other methods were reordered accordingly. We notice that the green cluster remains stable over all other methods since all but one are grouped together every time. Under the influence of the target predictions one compound dissapears and is replaced by another. For the weighted procedures, the result for a weight of 0.5 is shown. This figure could also have been made over the weights in the weighted clusterings procedures as presented below.

```
> L=list(MCF7_F,MCF7_Weighted,MCF7_T)
> names=c("FP",seq(1,0,-0.1),"TP")
>
```

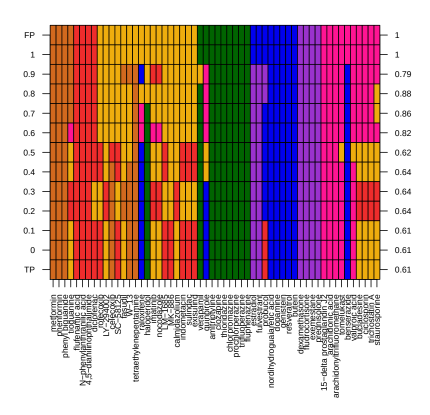


Figure 4: Comparison of the weights in the weighted clustering for the CMAP MCF7 data

It is seen that again, as soon as target prediction information is involved, one compound dissapears from the cluster while all others remain. For higher weights, another compound joins the cluster.

Further, it is possible to cluster the single source clusterings and multi-source clustering next to each other such that a visual comparison is possible. This is mostly interesting of more than 2 data sources are available. The function will here be demonstrated for 2 nonetheless. It is made sure that for both comparison plots the same reference has been used, namely the first element of ListM.

location=NULL)

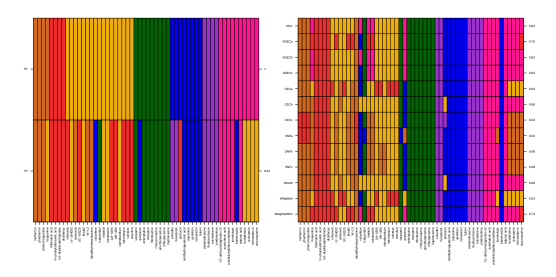
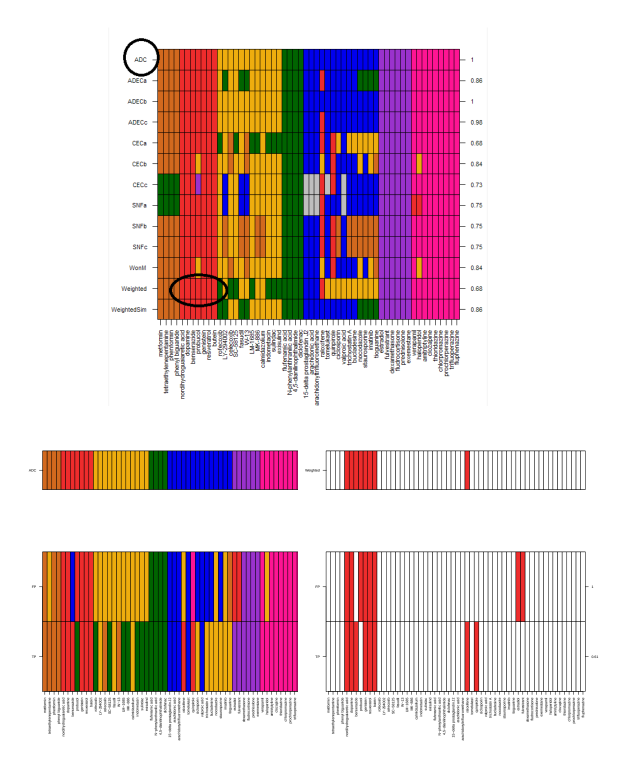


Figure 5: Comparison of single source and multi-source clustering

Further, an interactive function exists as well. In this function a comparison plot will be made with the *ComparePlot* function of the results provided in ListM. Then, there exists the opportunity to click on this figure to either indicate one specific cluster or an entire method. To indicate a cluster, we click on the color of that cluster for the method of choice. To indicate an entire method, we click on the name of the method on the left of the figure. The indicated cluster/method will then be compared to the elements provided in ListS in a new plot. It is made sure that for both comparison plots the same reference has been used, namely the first element of ListM.

=c(8,4.5,2,2.5), Interactive=TRUE, N=2)



 $\label{eq:Figure 6: Interactive Plot - Example} Figure \ 6: \ Interactive \ Plot \ - \ Example$

Two clustering results X and Y can be compared with the help of a test statistic calculated by the CompareSilCluster function. Single data sources can be provided in their data format. A distance matrix is necessary for all multiple source clusterings. A medoid clustering is performed of which the silhouette widths and cluster memberships are retrieved. The silhouette widths of data X are regressed against the corresponding memberships but also against the memberships determined by data Y. The R^2 values are obtained and the statistic is calculated as: A p-value is obtained by bootstrapping the distance matrices.

3.4 Selection of a Specific Cluster

As mentioned above, interest lies in stable clusters. We can now select one specific cluster and perform further analysis. We investigate of which compounds the cluster exists and how it was formed over the different results and/or weights.

To find the compounds of a cluster, we can use the *FindCluster* function. Based on the matrix produced by *MatrixFunction*, we can provide the row and the cluster number (counting from 1 to 7 from left to right with every color being a new cluster in the first row) and the compounds of that cluster will be returned. We will focus on the green cluster of the target predictions.

Further, we can track how this cluster was formed with the help of the *TrackCluster* function. We will indicate to follow the cluster specifically (followClust=TRUE) and not the largest group of compounds of the originally indicated cluster that remain together (followMaxComps=FALSE). This might involve a change of cluster.

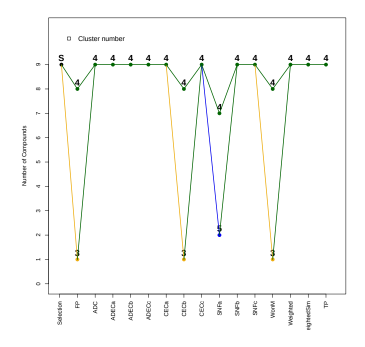


Figure 7: Tracking of the green cluster over the methods

Next, we perform secondary analysis on the cluster of interest.

4 Secondary Analysis

4.1 Differential Gene Expression

Interesting genes are those that behave differently in the selected compounds. The gene may have a higher variation or can be up-regulated or down-regulated. Several methods exist to find such differentially expressed genes such as a two sample t-test or a permutation test. It was opted to work with a method called Linear Models for Microarrays (limma) (Smyth (2004)). The resulting p-values were adapted for FDR and the significance level was chosen to be 0.05. Limma is in fact a regular two sample t-test with an adjusted denominator. This adjustment is made to avoid that genes that have a small fold change and a small variance will be considered significant by the procedure. The denominator is estimated with an empirical bayesian approach.

We could test all clusters with the *DiffGenes* functions but will foucus only on our compounds of interest. These will be tested against all other compounds combined and we ask to collect the top 10 genes. Since we focus on our selection of compounds, there is no need to specify clustering results as the result will be independent from the grouping of the compounds.

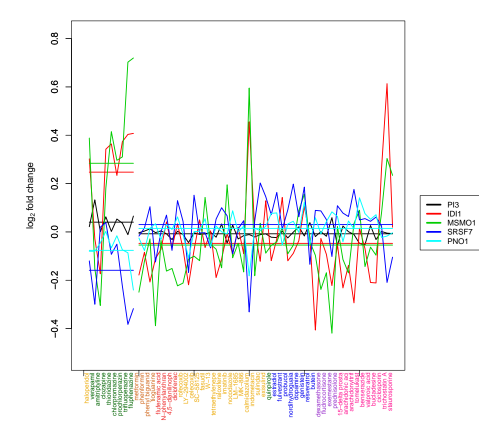


Figure 8: The Top 5 Genes

4.2 Pathway Analysis

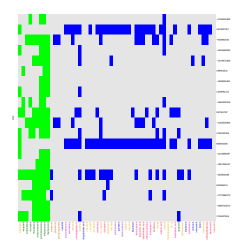
The final step in the analysis is to allocate the genes to a gene set or pathway. If a gene set is enriched, i.e. the probability to see this many significant genes of this gene set by chance is low in the selected compounds, one may be fairly sure that the compounds share an activity on this pathway. The selected database for pathway analysis is the Gene Ontology (GO) database. This is an hierarchical database which starts from more general terms (annotations) to very specific ones. The pathway analysis performed was functional class scoring, also known as MLP. The search for enriched gene sets starts from the p-values of all genes. For a predefined gene set, it calculates a score that summarizes the significance of all the genes included in that specific set. This score is the mean of the negative logarithm of the p-values (MLP).

It is then determined how likely it is to see the MLP value by chance. This is done by a comparison with the empirical null distribution. To attain this distribution, the labels are permuted across the samples and the MLP is recalculated. This is repeated a few times for all gene sets. The null distribution of MLP and the observed value of MLP are compared and if there is a small probability to find the observed MLP (a small p-value for this distribution), the score is deemed significant. The MLP method to perform pathway analysis is based on resampling of the data. Therefore it is recommended to perform the pathway analysis multiple times to observe how much the results are influenced by a different sample. We will perform iterative pathway analysis for our selection of compouds with the *PathwaySelectionIter* function. Finally, we will found out how many of these were shared over the different iterations with the *Genseset.intersectSelection* function.

4.3 Characteristic Features

It is seen that genes were found to be differential expressed for the selected compouds which form the green cluster based on target predictions. We could now investigate whether there are fingerprints or target predictions that define this cluster. This can be done with the *ChooseFeatures* function. The function has the option to provide the input interactively. A dendrogram of a provided clustering result will be produces and the user can identify the cluster of interest by clicking. However, the input can also be provided manually as we will do now.

We plot the top identified features with the FeaturesPlot.



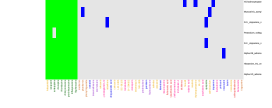


Figure 9: Top 20 Fingerprints - MCF7

Figure 10: Top 20 target predictions - MCF7

References

- Fodeh, J., Brandt, C., Luong, B. T., Haddad, A., Schultz, M., Murphy, T., and Krauthammer, M. (2013), "Complementary Ensemble Clustering of Biomedical Data," *Journal of Biomedical Informatics*, 46, 436–443.
- Hastie, T., Tibshirani, R., and Friedman, J. (2009), The Elements of Statistical Learning: Data Mining, Inference, and Prediction, Springer, chap. Unsupervised Learning, pp. 501–527.
- Perualila-Tan, N., Shkedy, Z., Talloen, W., Goehlmann, H. W. H., Consortium, Q., Van Moerbeke, M., and Kasim, A. (in press), "Clustering of Chemical Structure and Bioactivity Data in Early Drug Discovery." *Journal of Bioinformatics and Computational Biology*.
- Smyth, G. K. (2004), "Linear Models and Empirical Bayes Methods for Assessing Differential Expression in Microarray Experiments," *Statistical Applications in Genetics and Molecular Biology*, 3.
- Wang, B., Mezlini, M. A., Demir, F., Fiume, M., Tu, Z., Brudno, M., Haibe-Kains, B., and Goldenberg, A. (2014), "Similarity Network Fusion for Aggregating Data Types on a Genomic Scale," *Nature*, 11, 333–337.

5 Software used

- R version 3.1.3 (2015-03-09), x86_64-w64-mingw32
- Locale: LC_COLLATE=Dutch_Belgium.1252, LC_CTYPE=Dutch_Belgium.1252, LC_MONETARY=Dutch_Belgium.1252, LC_NUMERIC=C, LC_TIME=Dutch_Belgium.1252
- Base packages: base, datasets, graphics, gr
Devices, methods, parallel, stats, stats4, tools, utils
- Other packages: AnnotationDbi 1.28.2, Biobase 2.26.0, BiocGenerics 0.12.1, biomaRt 2.22.0, cluster 2.0.1, DBI 0.3.1, GenomeInfoDb 1.2.4, ggplot2 1.0.1, gtools 3.4.1, IntClust 0.0.1, IRanges 2.0.1, org.Hs.eg.db 3.0.0, plotrix 3.5-11, pls 2.4-3, rj 2.0.3-1, RSQLite 1.0.0, S4Vectors 0.4.0, SNFtool 2.2
- Loaded via a namespace (and not attached): a4Base 1.14.0, a4Core 1.14.0, ade4 1.7-0, bitops 1.0-6, caTools 1.17.1, colorspace 1.2-6, digest 0.6.8, gdata 2.13.3, gplots 2.16.0, grid 3.1.3, gridExtra 0.9.1, gtable 0.1.2, heatmap.plus 1.3, KernSmooth 2.23-14, lava 1.4.0, limma 3.22.7, MASS 7.3-40, MLP 1.14.0, munsell 0.4.2, plyr 1.8.1, prodlim 1.5.4, proto 0.3-10, Rcpp 0.11.5, RCurl 1.95-4.5, reshape2 1.4.1, rj.gd 2.0.0-1, scales 0.2.4, splines 3.1.3, stringr 0.6.2, survival 2.38-1, XML 3.98-1.1