

canceR: A Graphical User Interface for accessing and modeling the Cancer Genomics Data of MSKCC

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1 Introduction

canceR is a graphical user friendly interface to explore, compare, and analyse all available Cancer Data (Clinical data, Gene Mutation, Gene Methylation, Gene Expression, Protein Phosphorylation, Copy Number Alteration) hosted by the Computational Biology Center (cBio) at Memorial-Sloan-Kettering Cancer Center (MSKCC). **canceR** implements functions from various packages:

1. to access, explore and extract Genomics Cancers Data Base of MSKCC (**cgdsr**, [2, 4]),
2. to associate phenotypes with gene expression (**phenoTest**, [8]),
3. to predict which biological process or pathway or immune system are significantly different under the phenotypes and which genes are associated ((GSEA-R) [10, 9]),
4. to predict the most up/down regulated gene sets belonging to one of MSigDB collections [10] (**GSEAlm**, [7]),
5. to classify genes by diseases (**geNetClassifier**, [1]), or
6. to classify genes by variable or phenotype (**rpart**, [11]),
7. to plot genes correlations.

2 Installation

No R packages:

- tktable <http://tktable.sourceforge.net/>
- BWidget <http://sourceforge.net/projects/tcllib/files/BWidget/>
- GnuWin32 (for Windows) http://sourceforge.net/projects/getgnuwin32/?source=typ_redirect

For Debian distribution (GNU/Linux)

```
sudo apt-get install ("r-cran-tcltk2", "r-cran-tkrplot")
sudo apt-get install('Tk-table', 'BWidget')
sudo apt-get install libcurl4-openssl-dev
sudo apt-get install r-cran-xml
```

For Windows distribution

LibXml2: parser for XML <http://sourceforge.net/projects/gnuwin32/files/>

For OS X distribution X11: graphics device

run R and write these lines in console to install dependencies.

```
## dependencies from CRAN
install.packages("RCurl", "XML")
install.packages(c("cgdsr", "tkrplot", "Formula", "RSvgDevice", "RCurl" ))
## dependencies from Bioconductor
source("http://bioconductor.org/biocLite.R")
biocLite("GSEABase", "GSEAlm", "geNetClassifier", "Biobase", "phenoTest")
biocLite("canceR")
library(canceR)
canceR()
```

Get the development version from github

```
devtools::install_git("kmezoud/canceR")
```

3 Starting Window

run R and write theses lines in console to run **cancerR** package.

```
library(cancerR)
cancerR()
```

The starting window (Figure 1) loads all available Cancer Studies (Figure 1 3) or search some ones by key word (Figure 1 4). Before to get Cancers Data (Figure 1 7), it is important to set workspace for output files (Figure 1 1). The starting window displays Help menu where user can get this vignette (Figure 1 2).

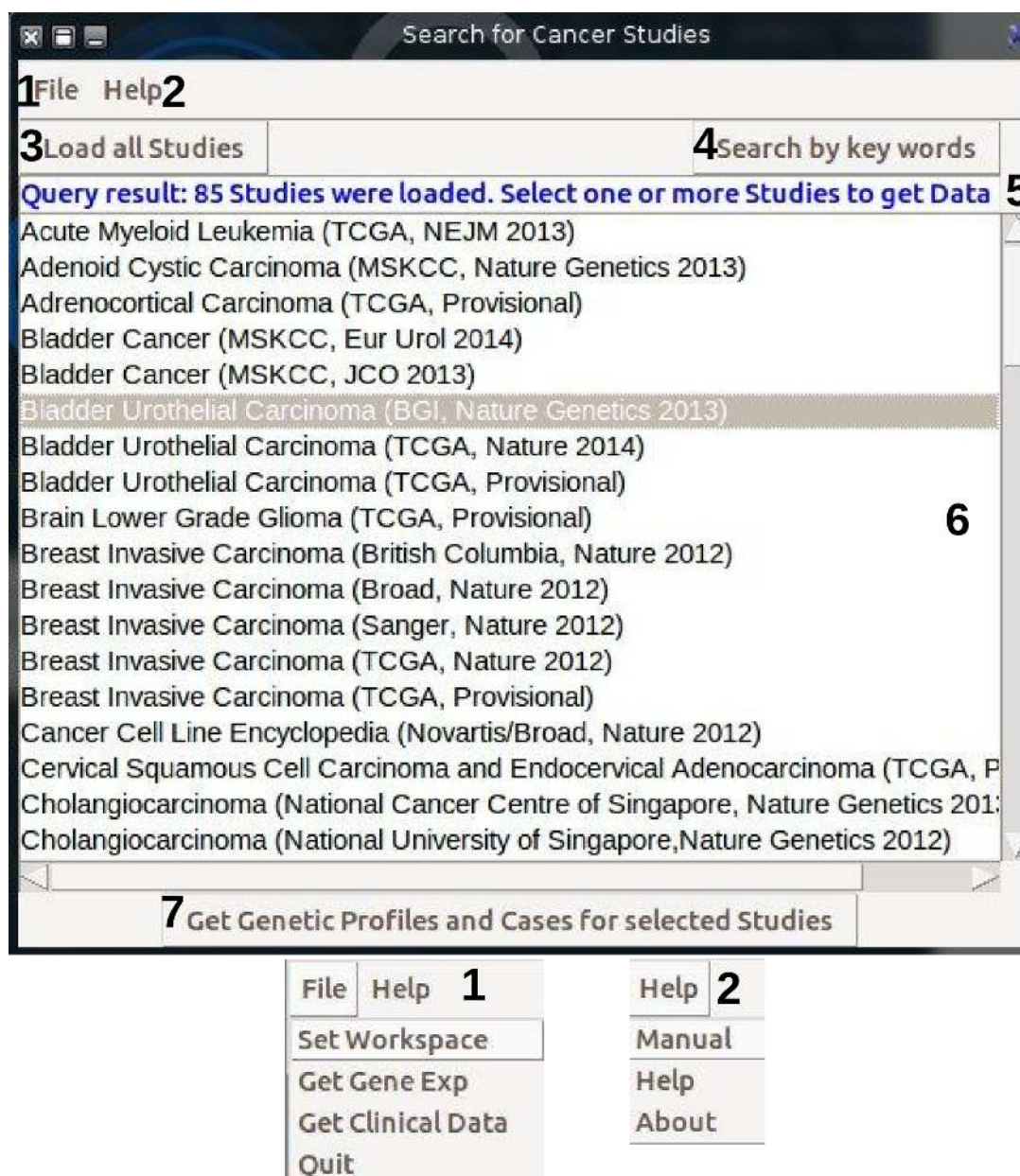


Figure 1: Starting Windows. 1, File Menu; 2, Help Menu; 3, Button to get all available studies; 4, Button to get only matched studies using key words; 5, list box that displays the number of studies listed in 6; 6, list box that displays the result of querying the Cancer Genomics Data Server. User can select one or multiple Studies; 7, Button to get Genetic Profiles and Clinical data for selected Studies.

3.1 Setting Workspace

canceR package uses input files to compute models and generates output files for biological knowledges. It is important to set workspace and know the location of used files and results. The Button "Set Workspace" allows user to set easily workspace (Figure 2). User needs just to browse workspace folder or creates a new one. The others necessary folders would be created by simple pressing "Set" buttons.

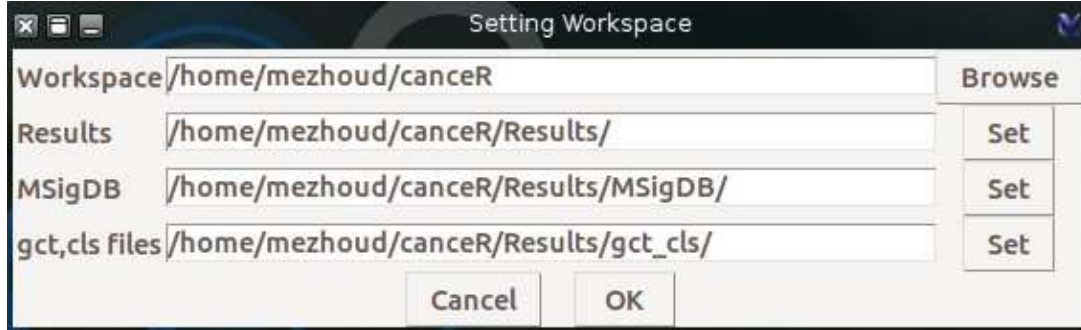


Figure 2: Setting Workspace

4 Main Window

After selecting studies and pressing on "Get Cases and Genetic Profiles" Button, the main window appears (Figure 3) and displays the progress of loading data of selected studies. The Main Window has a **Toolbar** with Menus (see following paragraphs). It is subdivided in two columns. The first column lists Cases for all selected studies. The first line of every study indicates its Index and its short description. The remain lines enumerate Cases with short description of data type and the number of samples. The second list box shows selected Cases. Similarly, the second column displays informations of Genetic Profiles. User can select a single or multiple lines with attention to correspond the Case with appropriate Genetic Profile.

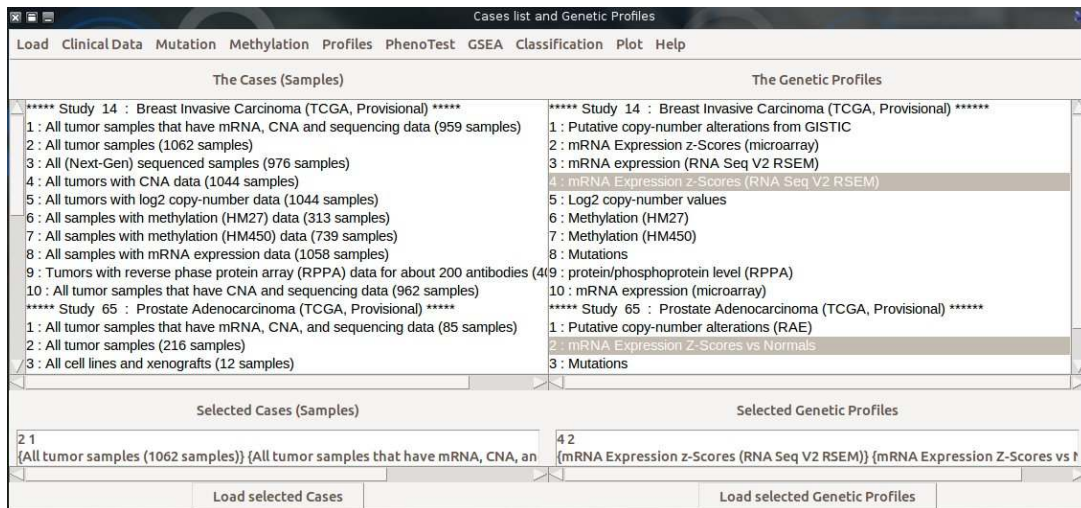


Figure 3: Main Window of **canceR** package. 1, Toolbar, 2, list box of loaded Cases; 3, list box of selected Cases; 4, list box of loaded Genetic Profiles; 5, list box of selected Genetic Profiles

4.1 Gene List

The first step to get genomics data is to specify what are interesting genes for user. The **Gene List** button browses folders to load Gene list file or displays examples of genes list. The genes could be in text file (.txt) with one gene by line using HUGO gene Symbol. The function removes automatically duplicate genes.

4.2 Clinical Data

The **Multiple Cases** button displays successively selected Cases. Results are returned in a table with row for each case and a column for each clinical attribute (Figure 4B). User could select all or some clinical data by checking dialog box (Figure 4A). For example, we select clinical attributes:

- Overall Survival months: Overall survival, in months.
- Overall Survival Status: Overall survival status, usually indicated as "LIVING" or "DECEASED".
- Disease Free Survival months: Disease free survival, in months.
- Disease Free Survival status: Disease free survival status, usually indicated as "DiseaseFree" or "Recurred/Progressed".
- Age at diagnosis: Age at diagnosis.

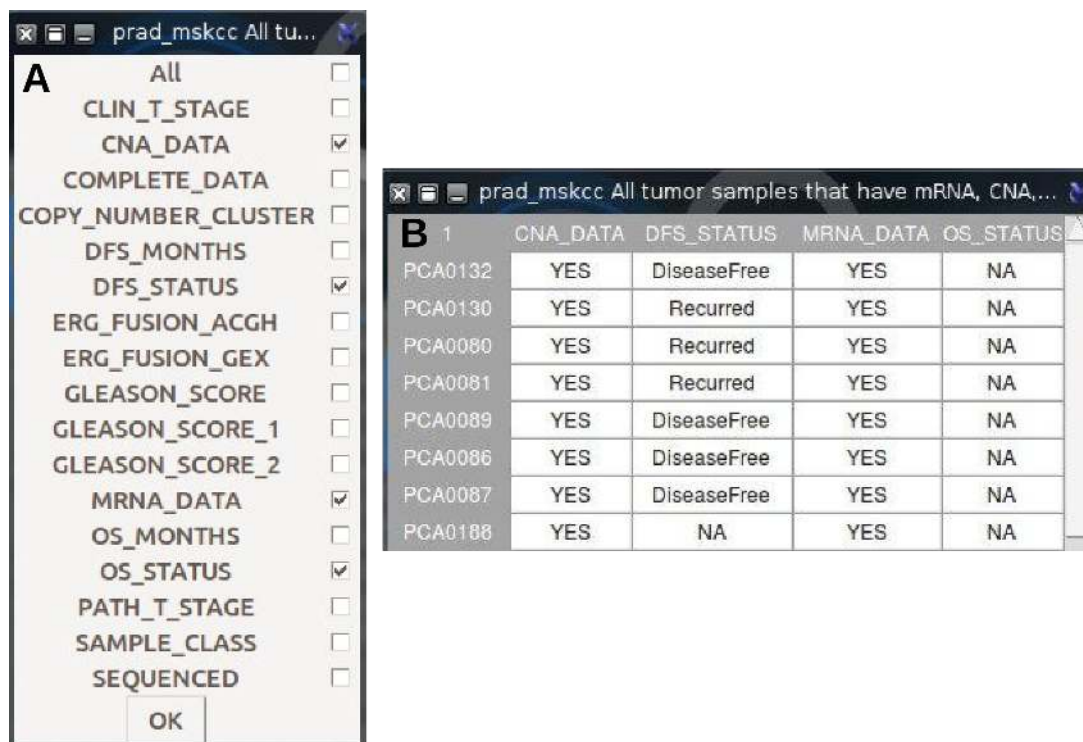


Figure 4: Getting clinical data for Breast Invasive Carcinoma. A, Dialog Check Box to select clinical data; B, Results of querying clinical data of Breast Invasive Carcinoma (TCGA, Nature 2012).

4.3 Mutation

User can search all mutation in gene list of all selected studies. He needs to select **All tumors samples** in **Cases** and **Mutations** in **genetics profiles** to get mutations (Figure 5).

Mutation function allows user to select about 15 informations corresponding to mutations (Figure 6A). The results is a table with rows for each sample/case, and columns corresponding to the informations checked in dialog mutation check box (Figure 6B).

User can filter mutation result only for specific amino acid change (Figure 7).

4.4 Methylation

User can search gene methylation and its correlation with mRNA expression. User needs to select **Cases** and **Genetic Profiles** with same methylation assay (HM450 or HM27) for the same study. Multiple Cases selection is allowed for one gene list (Figure 8).

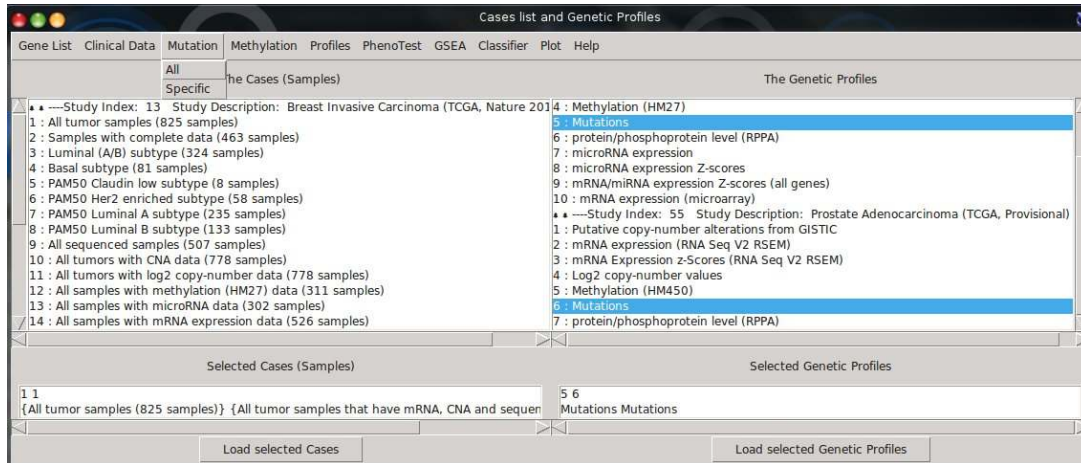


Figure 5: Get all/specific mutations for selected Cases

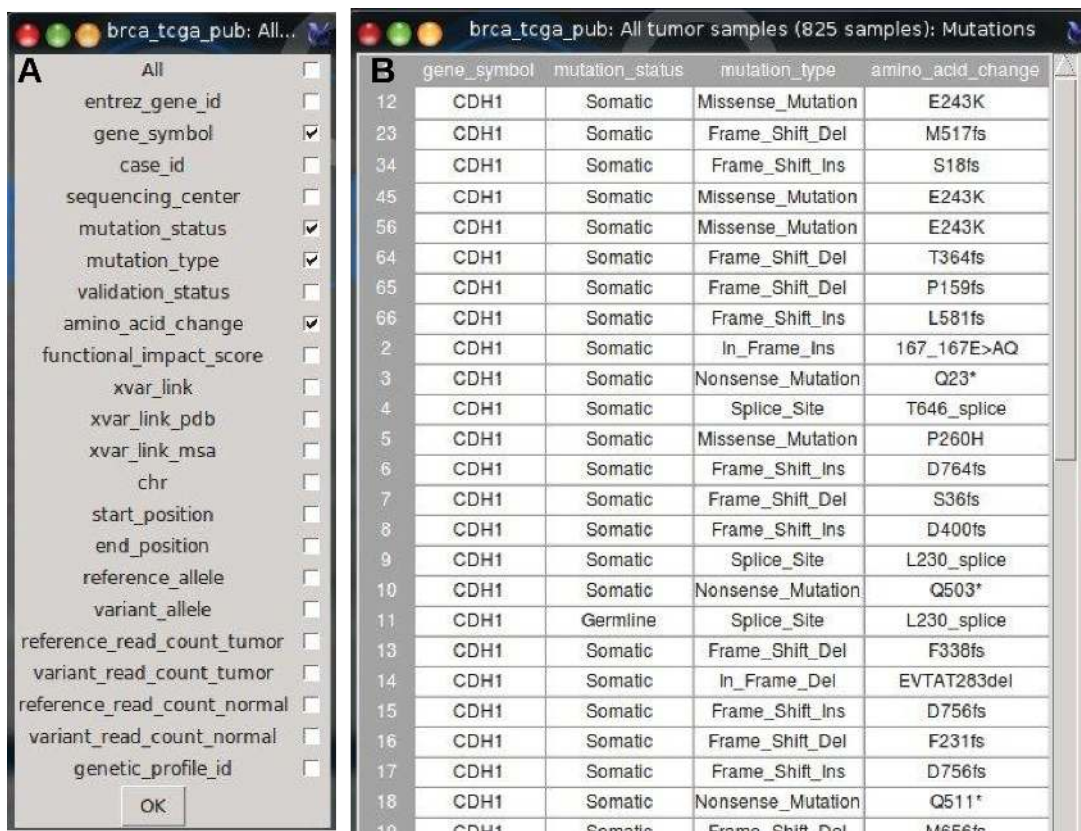


Figure 6: A, mutation dialog; B, mutation data

The dialog box of **methylation** function allows user to specify the threshold of the correlation rate (Figure 9A). cBioportal [2, 4] includes only methylation data from the probe with the strongest negative correlation between the methylation signal and the gene's expression. The result table (Figure 9B) lists genes with median of rate upper than 0.8.

4.5 Profiles

The function get Profile Data depends on gene list, cases, and genetic profiles. If a **Single gene** option is done, dialog box appears to specify gene symbol (Figure 10A). The returned dialog check box allows user to choose some/all profiles data (Figure 10B). The result (table) lists some/all genetic profiles data in columns (CNA, Met, Mut, mRNA, RPPA) and all available samples in rows (Figure 10C). Oppositely,

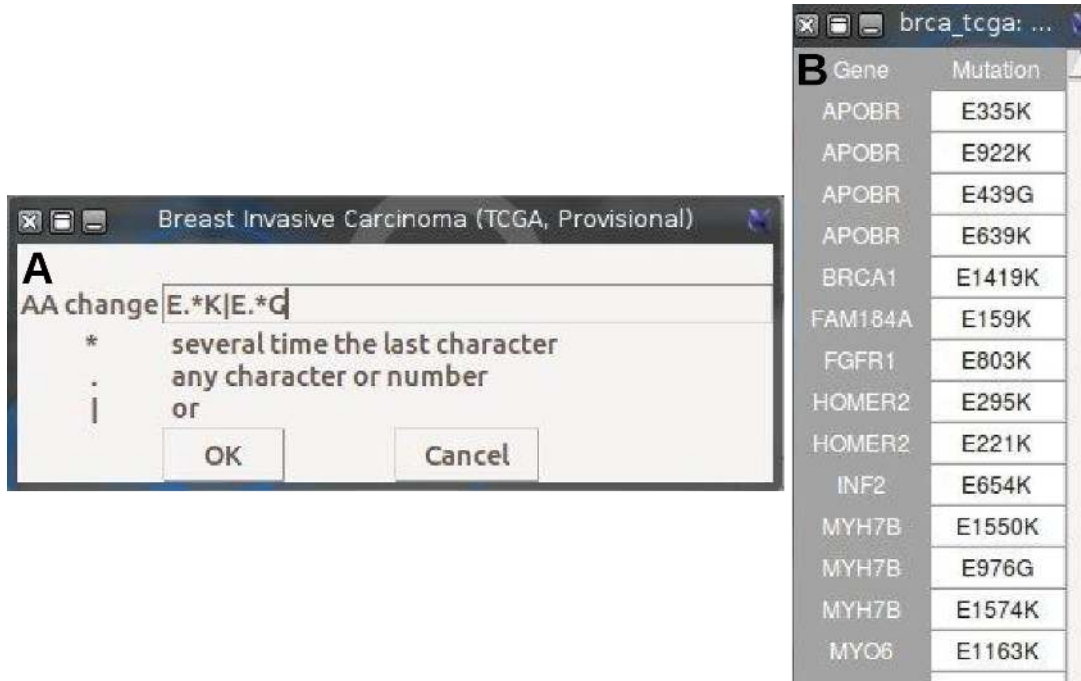


Figure 7: Filtering mutation by specific amino acid change. A, the search is done using regular express language. In this case the query specify 3 options: D438., search all mutation of D Amino acid at position 438 or D.*K search the position of amino acid change of D to K or D.*. search for all D amino acid changes; B, Results of the merge of the 3 options filter of D amino acid changes.

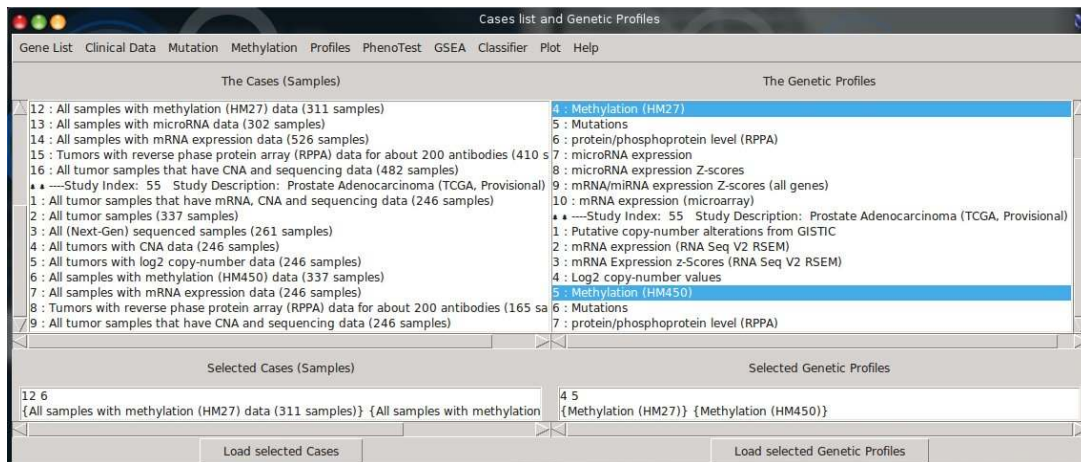


Figure 8: Selecting methylation data

if **Multiple genes** option is done, the returned table displays genes expression for gene list (column) for all samples (rows). In the case of multiple genes, the tables are saved in **Results/ProfilesData** folder.

4.6 PhenoTest

The function was implemented from package **PhenoTest** [8]. The object of this function is to predict the association between a list of phenotype variables (Survival, DFS Status, OS Status) and the gene expression. There are two possible formula to get associations:

1. Three variables: *Survival* status (event/time as Dead-Living / 30 Months), *Categorical* or *ordinal* description (DSF STATUS or Tumor stage), and *Continuous* value (DFS MONTHS, Tumor size).
2. Two variables: *Categorical* or *ordinal* description (DSF STATUS or Tumor stage), and *Continuous* value (DFS MONTHS, Tumor size). In this case user does not need to select any variables for

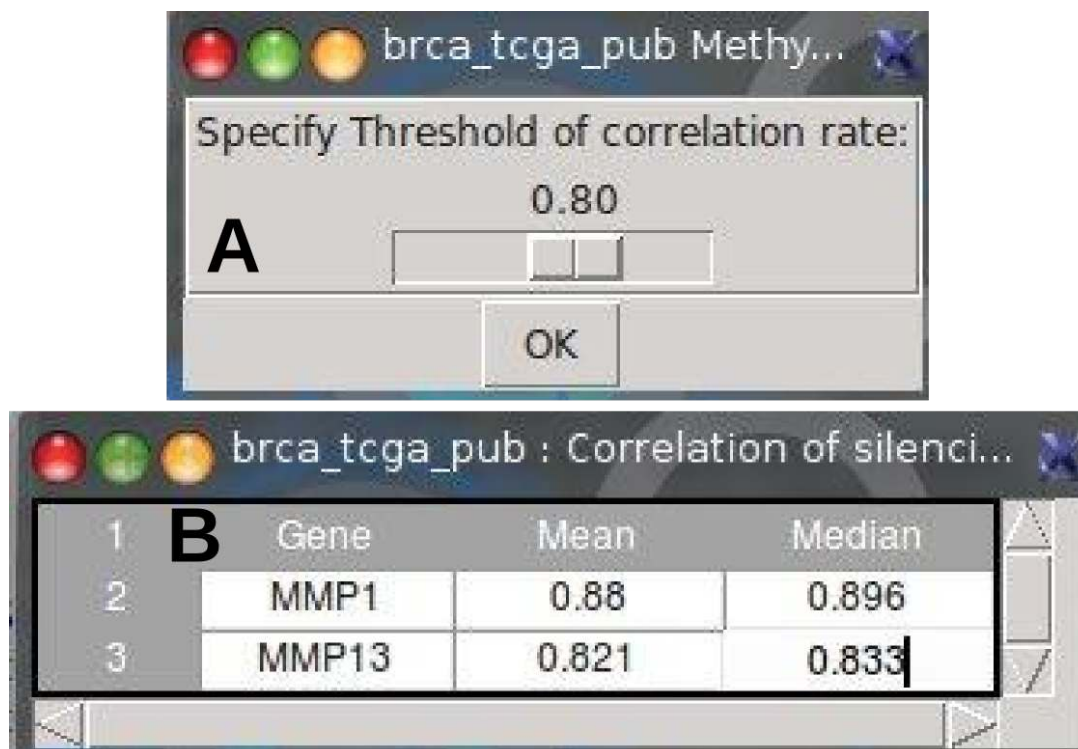


Figure 9: A, Methylation dialog box; B, Methylation results. Correlation of silencing gene expression by methylation (HM27) with correlation rate $r > 0.8$ for gene list in Breast Invasive Carcinoma (TCGA, Nature 2012).

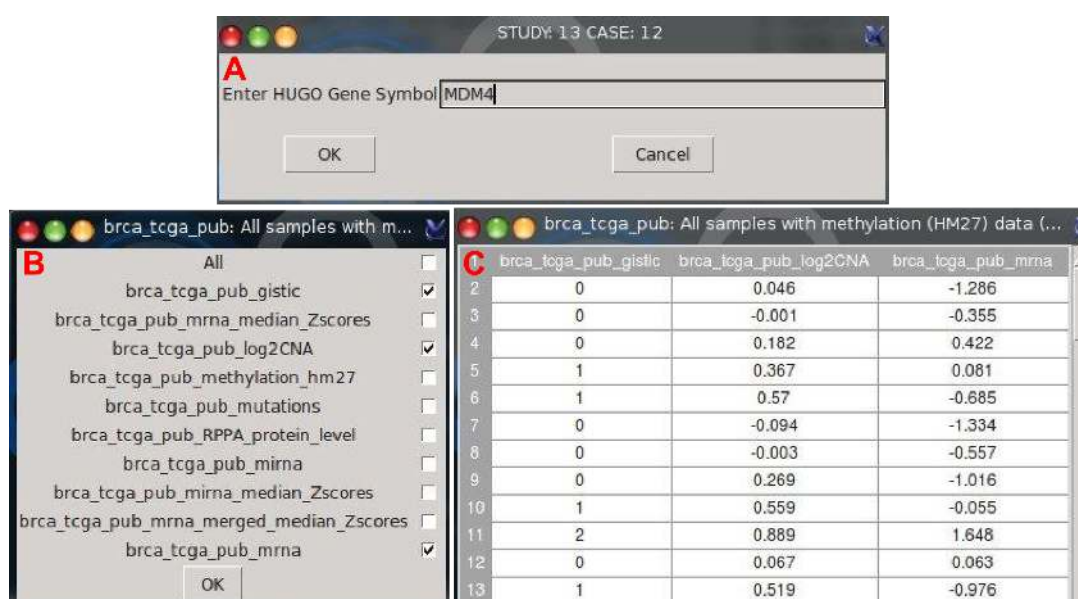


Figure 10: Getting profile data of a single gene

survival variable in the phenoTest dialog box (Figure 12A).

The output of this function does not expect to give systematically a relevant association between all formula of the chosen variables, although in some cases it is possible to cluster a list of genes significantly regulated (gene expression) at a range of tumor size (continuous) or tumoral stage (ordinal) for recurred or DiseaseFree cases (survival and categorical). The type of variables could be explored with Clinical Data tables and selected in the phenoTest dialog box (Figure 12A). The effect of both *continuous*, *categorical* and *ordinal* phenotype variables on gene expression levels are tested via *lmFit* from *limma* package [12].

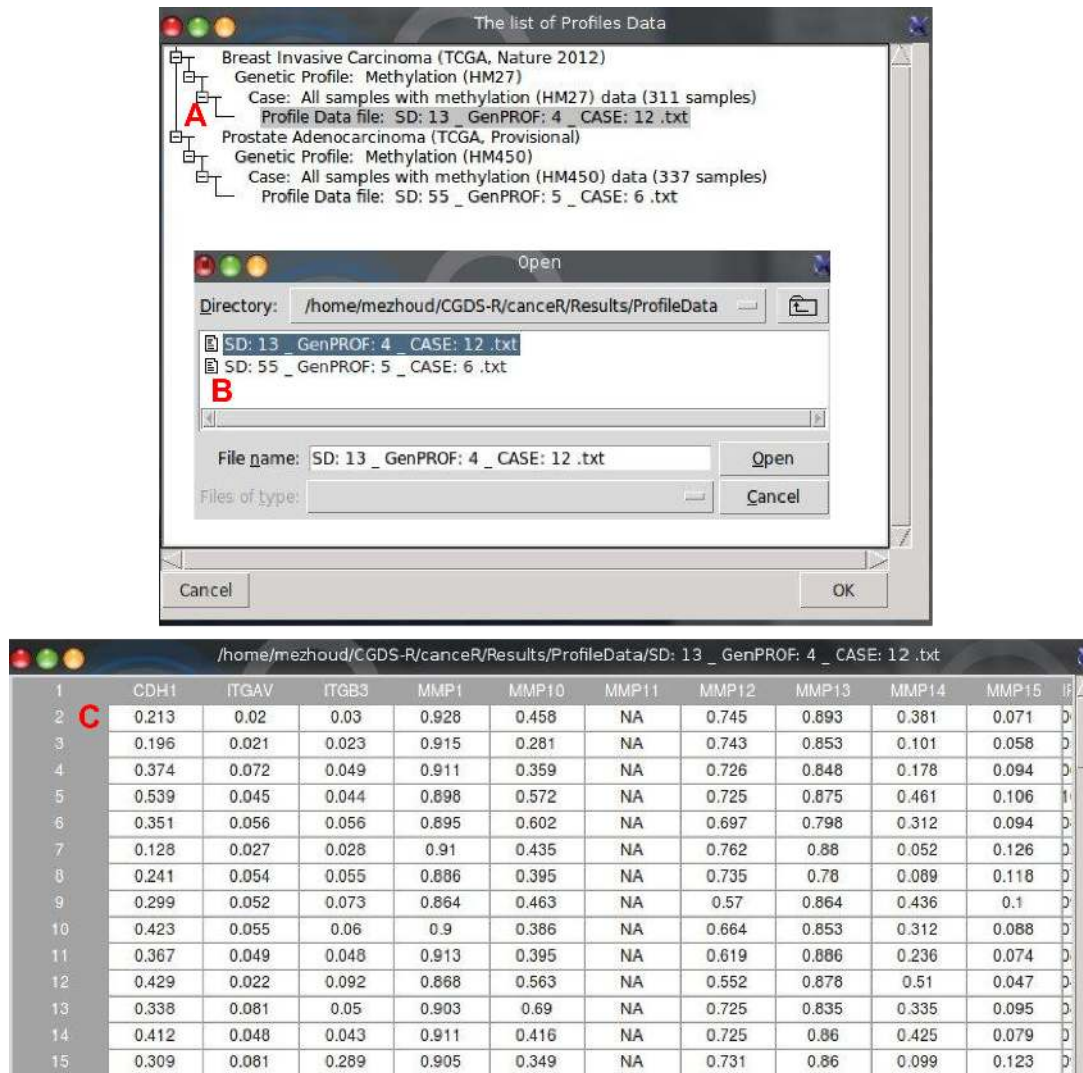


Figure 11: Profile data of a multiple genes

Gene expression effects on *survival* are tested via Cox proportional hazards model [3], as implemented in function `coxph` from `survival` package.

NB: Continuous or Categories can not have more than 4 classes.

Examples

Study: Prostate Adenocarcinoma (Broad/Cornal, Call 2013)

- Cases: All tumor samples (57 samples),
- Genetic Profiles: mRNA expression,
- Gene list: 1021.txt file
- Survival variable: empty
- Categorical variable: Pathology Tumor Stage
- Numeric variable: Serum PSA level
- pVal adjust method: BH
- PhenoTest with Two variables (Figure 12)

After running **Pheno/Exp**, **PhenoTest** function returns two tables. The first table ranks gene list by **pval** (Figure 12B). The first part (red square) displays pValues of the association between gene expression and Tumor stage. The second part (blue square) displays the fold change (fc) by PSA level rang.

Interpretation: Notice that a single pValue is reported for each phenotype variable. For categorical variables these corresponds to the overall null hypothesis that there are no differences between groups. In the second table, **PhenoTest** function filters only gene that has significant pval (pval <0.05, Figure 12C red). Here we see that tumor stage has been categorized into 2 groups (pT2c, pT3c) and PSA level has been ranged into 2 groups (7.3-12.9, 12.9-16.7). This results shows that **AN03** gene is significantly down regulated (negative fold change) for the two pathology tumor stages (pT2c, pT3c).

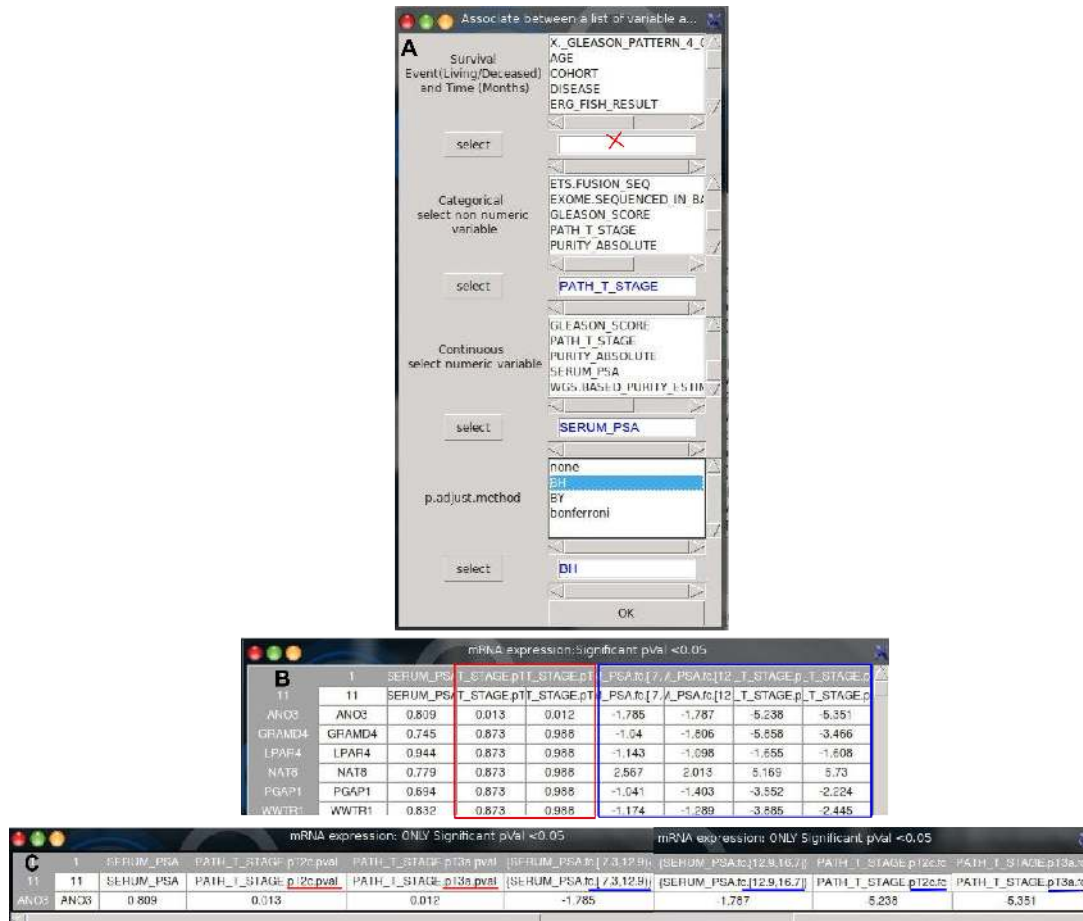


Figure 12: PhenoTest; A, Dialog Box used to select variables; B, Results; C, Only significant pValues.

Heteroneous Clinical Data In some cases it is possible to have digital (0-9) and character (a-z) data in the same variable. in this case phenoTest function considers it as Categorical variable (Figure 13).

Study: Prostate Adenocarcinoma, Metastatic (Michigan, Nature 2012)

- Cases: All tumor samples (61 samples),
- Genetic Profiles: mRNA expression
- Gene list: 1021.txt file
- Survival variable: OS MONTHS, OS STATUS
- Categorical variable: OS STATUS
- Numeric variable: Serum PSA level
- pVal adjust method: BH

prad_broad_2013: mRNA expression									
1	AGE	COHORT	DISEASE	_IN_B/_SON_S/_ATH_T/_STA/	PURITY_ABSOLUTE	ERUM_PSD_PURITY_E			
PR.08.2153	55	Uropath	primary adenocarcinom	NO	4+3	pT3a	0.55	7.8	0.52
PR.08.4154	58	Uropath	primary adenocarcinom	NO	3+4	{pT4 N1}	0.72	36	0.71
P07.837	69	Uropath	primary adenocarcinom	YES	4+3	pT3a	{not determined}	7.8	0.59
R.STID000000268	71	WCMC	primary adenocarcinom	YES	4+3	pT2c	1	7.6	0.76
PR.2741	73	WCMC	primary adenocarcinom	NO	4+4	pT3a	0.71	5	0.72
P05.1657	47	Uropath	primary adenocarcinom	NO	3+4	pT2b	0.61	9.3	0.6
P08.5852	51	Uropath	primary adenocarcinom	NO	3+3	pT2c	{not determined}	12.6	0.47
P03.2345	56	Uropath	primary adenocarcinom	YES	3+4	pT2c	0.55	7	0.56
PR.3042	67	WCMC	primary adenocarcinom	NO	3+4	pT2c	{not determined}	3.3	0.74
PR.03.728	54	Uropath	primary adenocarcinom	NO	3+4	pT3a	non-aneuploid	12.9	not determinec

Figure 13: Clinical Data heterogeneity. In PURITY ABSOLUTE column is there digital and character values. in this cas PhenoTest consider this variable as Categorical.

- PhenoTest with three variables (Figure 14)

In This test, Overall Survival (OS.STATUS) was used in survival and caterogical variables. The Clinical Data does not have enough categorical variables. Figure 14 B and C shows significant association between 7 genes and Living Status (OS.STATUS.Living.pval column). The two last columns show opposite regulation of the 7 genes expression in living patient with serum PSA level. The Cox proportion hazard model does not give results with survival variables (OS.STATUS column).

Study:Lung Adenocarcinoma (TCGA, Nature, in press)

- Cases: All Samples with mRNA expression data (230 samples),
- Genetic Profiles: mRNA expression z-Scores (RNA Seq V2 RSEM)
- Gene list: 1021.txt file
- Survival variable: empty
- Categorical variable: OS STATUS
- Numeric variable: OS_MONTHS
- pVal adjust method: BH
- PhenoTest with two variables.

In this Lung cancer Study, the test shows significant association between living patient and 10 genes expression (Figure 15).

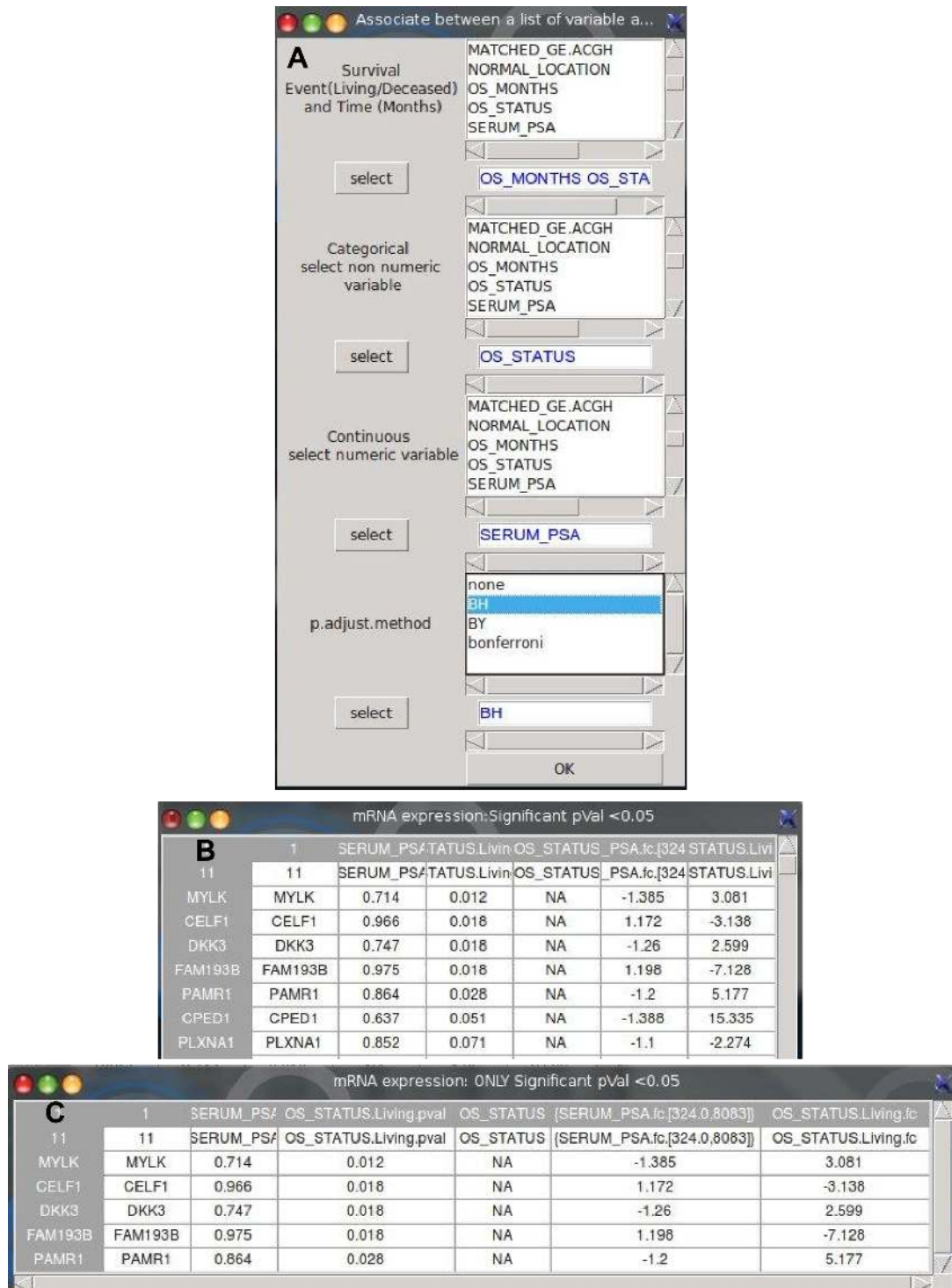


Figure 14: PhenoTest: Prostate Adenocarcinoma, Metastatic (Michigan, Nature 2012)

4.7 GSEA-R

Gene Set Enrichment Analysis (GSEA) is computational method that uses expression matrix of thousands of genes with phenotypes data (two biological states) and Molecular Signatures DataBase (MSigDB) to define which biological process or pathway or immune system are significantly different under the phenotypes and which genes are associated [10].

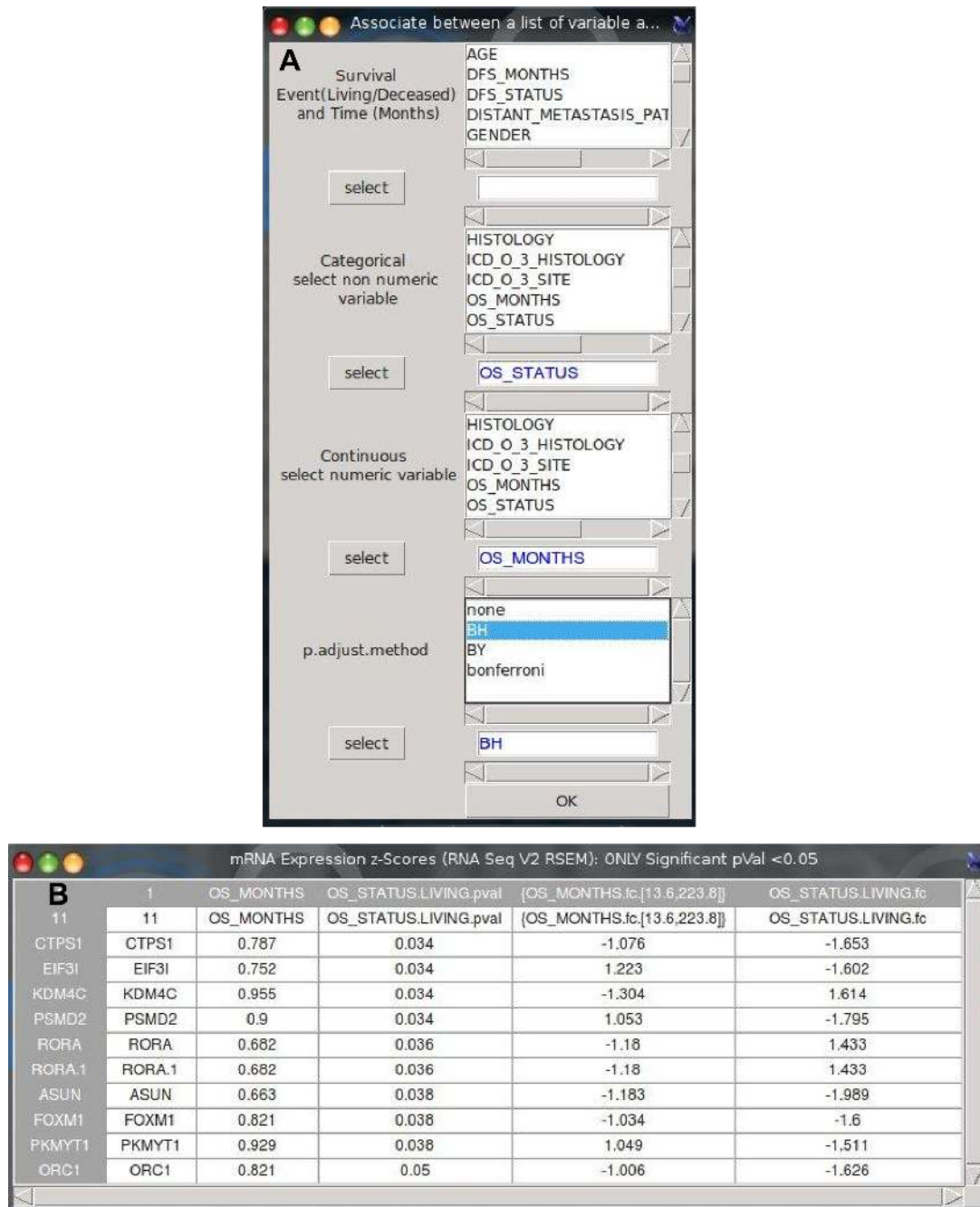


Figure 15: PhenoTest: Lung Adenocarcinoma (TCGA, Nature, in press)

4.7.1 Preprocessing of Experimental Data

getGCT_CLS function loads Profile and Clinical data of selected study and saves two files into "gct_cls" folder (Figure 16C).

The **GCT** file contents genes expression values with genes in the rows and samples in the columns.

The **CLS** file contents the two biological phenotypes selected from Clinical data. User needs to select clinical phenotype only with two classes.

4.7.2 Molecular Signatures DataBase

The Molecular Signatures DataBase (MSigDB) is a collection of annotated gene sets for use with GSEA computational method. The MSigDB gene sets are divided into 7 collections (positional gene sets,

curated gene sets, motif gene sets, computational gene sets, GO gene sets, oncogenic signatures, and immunological signatures). All these collections are available at (<http://www.broadinstitute.org/gsea/msigdb/index.jsp>). Every collections consists in a tab delimited file format (.GMT file) that describes gene sets. Each row shows annotation terme with associated genes. User needs to download .gmt file with genes Symbols and saves them into "workspace/MSigDB/" folder. The MSigDB folder is created with the file menu in the starting windows (Figure 1).

For more detail about GCT, CLS, GMT files, see this link (http://www.broadinstitute.org/cancer/software/gsea/wiki/index.php/Data_formats).

MSigDB Collection

C1: Positional Gene Sets Gene sets corresponding to each human chromosome and each cytogenetic band that has at least one gene. These gene sets are helpful in identifying effects related to chromosomal deletions or amplification, epigenetic silencing, and other region effects.

C2: Curated Gene Sets into Pathways Gene sets collected from various sources such as online pathway databases.

CGP: Chemical and Genetic Perturbation Gene sets represent expression signatures of genetic and chemical perturbations.

CP: Reactome gene sets Gene sets derived from the Reactome pathway database.

C3: Motifs Gene Sets Gene sets that contain genes that share:

MIR: microRNA targets A 3'-UTR microRNA binding motif.

TFT: transcription factor targets A transcription factor binding site defined in the TRANSFAC (version 7.4, <http://www.gene-regulation.com/>) database.

C4: Computational Gene Sets Computational gene sets defined by mining large collections of cancer-oriented microarray data.

C5: GO Gene Sets Gene sets are named by GO term (<http://geneontology.org/>) and contain genes annotated by that term: Biological Process, Cellular Component, and Molecular Function.

C6: Oncogenic Signatures Gene sets represent signatures of cellular pathways which are often dysregulated in cancer. The majority of signatures were generated directly from microarray data from NCBI GEO.

C7: Immunologic Signatures Gene sets that represent cell states and perturbations within the immune system. This resource is generated as part of the Human Immunology Project Consortium (HIPC; <http://www.immuneprofiling.org/>).

4.7.3 Examples

Study: Uterine Corpus Endometrioid Carcinoma (TCGA, Nature 2013)

- Cases: All Samples with mRNA, CNA, and sequencing data (232 samples),
- Genetic Profiles: mRNA expression (RNA Seq V2 RSEM)
- MSigDB: c5.bp.v4.0.symbols
- Gene list: 1021.txt file
- Nbr of Samples: 100
- Phenotype: DFS_STATUS

Based only on Gene list the function `getGCT,CLS` files builds the .gct and .cls files and save them under the folder "/gct_cls/". The Figure 16 shows the preprocessing steps to get gct and cls files.

For enrichment, GSEA function needs three files. The gct file with gene expression, the cls with phenotypes and gmt file with Molecular signature of Gene Sets. There are two options to load gmt file, from examples (Figure 17A, MSigDB.gmt button) available into `cancel` package or from "workspace/MSigDB/" folder (Figure 17A, browse button) . In the two ways the gmt files must be from Broad Institute (<http://www.broadinstitute.org/gsea/index.jsp>) and has gene Symbols.

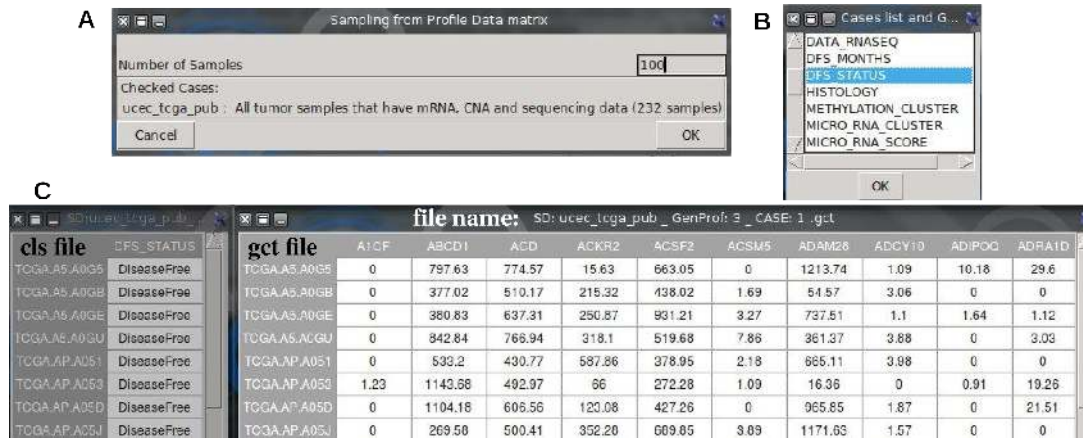


Figure 16: Preprocessing of **gct** and **cls** files. A, Sampling the Cases from Uterine Corpus Endometrioid Carcinoma study; B, selecting phenotype variable with only two classes; C, Displaying **cls** and **gct** files. The names of the two files are indicated in the title of the windows.

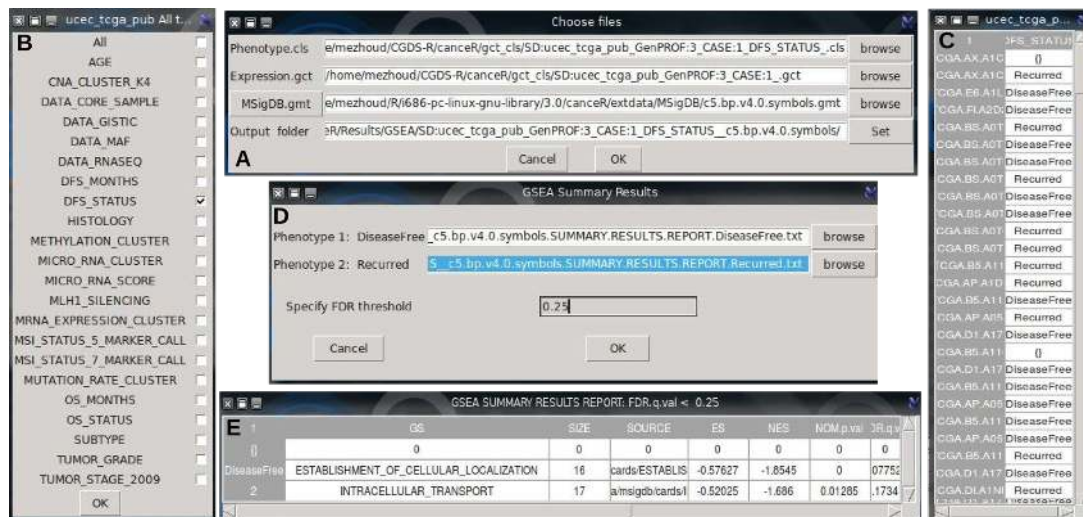


Figure 17: Gene Set Enrichment Analysis of Uterine Corpus Endometrioid Carcinoma study using "DiseaseFree/Recurred" phenotypes. A, Selecting **cls**, **gmt**, **gmt** files and setting output folder; B, Specifying the phenotype; C, displaying the classes of the selected phenotype; D, Selecting Summary results files of the output and setting FDR; E, displaying specific (FDR=0.25) Gene Sets (GS) involved in DiseaseFree phenotype. In this GSEA there is not significant GS involved specifically for Recurred phenotype.

Study: Breast Invasive Carcinoma (TCGA, Provosional)

- Cases: All Samples with mRNA expression data (562 samples),
- Genetic Profiles: mRNA expression (RNA Seq V2 RSEM)
- MSigDB: c5.bp.v4.0.symbols
- Gene list: 1021.txt file
- Nbr of Samples: 100
- Phenotype: OS_STATUS

The Figure 18 summarizes the gene sets enrichment analysis of Breast Invasive Carcinoma study using "Deceased/living" phenotypes. The Figure 18C shows 4 *vs* 1 biological process involved respectively into Deceased/Living phenotypes. The size and the genes of appropriate GS are indicated in the second (size) and third (source) columns. The report of significant GS are saved into "/Results/GSEA/name_of_folder/". Every GS has a report (.txt file) indicating the gene list and which

genes (CORE_ENRICHMENT column: YES) are involved in the specific phenotype (DECEASED for RESPONSE_To_STRESS). The plots of significant GS are save with .pdf format file. In a heat map, expression values are represented as colors for every patient, where the range of colors (red, pink, light blue, dark blue) shows the range of expression values (high, moderate, low, lowest). For more details about result interpretation see user guide of GSEA (<http://www.broadinstitute.org/gsea/doc/GSEAUserGuideFrame.html>).

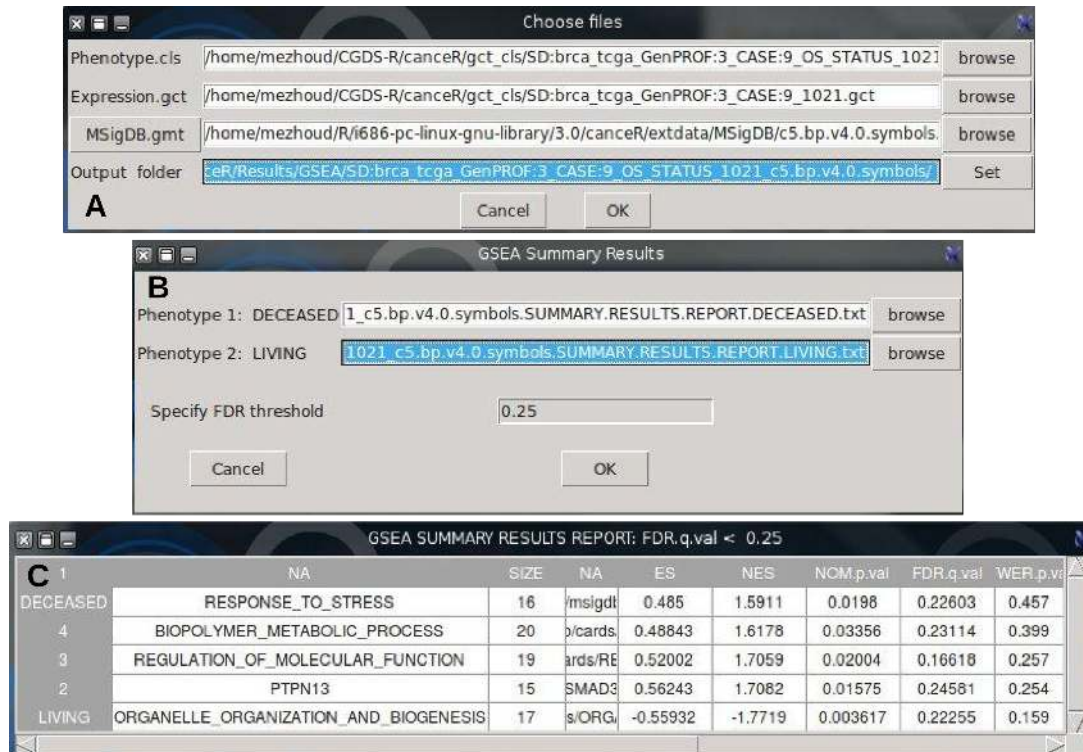


Figure 18: Gene Sets Enrichment Analysis of Breast Invasive Carcinoma (TCGA, Provosional) study using "Deceased/Living" phenotypes. A, Selecting cls, gmt, gmt files ans setting output folder; B, Selecting Summary results files of the output and setting FDR; C, displying specific (FDR=0.25) Gene Sets (GS) involved in Deceased and Living phenotypes.

4.7.4 GSEA-R Result Interpretation

The primary result of the gene set enrichment analysis is the enrichment score (ES), which reflects the degree to which a gene set is overrepresented at the top or bottom of a ranked list of genes. A positive value indicates correlation with the first phenotype and a negative value indicates correlation with the second phenotype. For continuous phenotypes (time series or PSA level), a positive value indicates correlation with the phenotype profile and a negative value indicates no correlation or inverse correlation with the profile [10].

The number of enriched gene sets that are significant, as indicated by a false discovery rate (FDR) of less than 25%. Typically, these are the gene sets most likely to generate interesting hypotheses and drive further research.

The number of enriched gene sets with a nominal p-value of less than 1% and of less than 5%. The nominal p-value is not adjusted for gene set size or multiple hypothesis testing; therefore, it is of limited value for comparing gene sets.

The false discovery rate (FDR) is the estimated probability that a gene set with a given NES represents a false positive finding. For example, an FDR of 25% indicates that the result is likely to be valid 3 out of 4 times. The GSEA analysis report highlights enrichment gene sets with an FDR of less than 25% as those most likely to generate interesting hypotheses and drive further research, but provides analysis results for all analyzed gene sets. In general, given the lack of coherence in most expression datasets and the relatively small number of gene sets being analyzed, an FDR cutoff of 25% is appropriate. However, if you have a small number of samples and use geneset permutation (rather than phenotype permutation)

for your analysis, you are using a less stringent assessment of significance and would then want to use a more stringent FDR cutoff, such as 5% [10].

Resolved limits

- GSEA does not accept negative values from Profile data. In this case the adding of absolute of less negative value to all the matrix is done.
- GSEA does not accept missing value in `c1s` file. The sampling is done only on existing phenotype information.
- GSEA needs more than 10000 to be robust but `url/gcds-r` package accept less than 1000 genes.
- The size of samples is between 50 and 100
- Removing and cleaning the heterogeneity of the data frames of gene expression: space, character, empty boxes and convert them to readable form by GSEA-R function.

4.8 Linear Modeling of GSEA (GSEAlm)

GSEAlm is a function implemented from **GSEAlm** package [7]. **GSEAlm** function is a Linear Model inference and diagnostics for Gene Set Enrichment Analysis. It uses mRNA expression matrix of gene list, variable(s) from clinical data (phenotype) and Molecular Signature Data Base (MSigDB) that groups genes sharing common biological function, chromosomal location, pathway or regulation [10]. The result is a prediction of the most up/down regulated gene sets belonging to one of MSigDB collections [10, 6].

The linear model assumes that the mean of the response variable has a linear relationship with the explanatory variable(s). The data and the model are used to calculate a fitted value for each observation (gene expression). It is strongly recommended to use crude gene expression data and avoid z-score or standardized preprocessing data. we recommend to use the mRNA expression (RNA Seq V2 RSEM).

Two options are available to get linear model. In the first option (phenotypes into disease) the model predicts which gene sets are modulated between patients having the same disease and different phenotypes (OS_STATUS, DFS_STATUS). In the second option (disease *vs* disease) the model predicts which gene sets are modulated between patient having different disease.

The output is saved in `/Results/GSEAlm/selected-disease-name` folder. The names of output files describe their content. Three files are saved by model. The `pVal_MSigDB_GeneList_Disease.txt` file lists full results. The `down/upRegulated_MSigDB_GeneList_Disease.txt` files filter only significant gene sets. The filtered result is merged in one table and displayed in the screen.

Results interpretation:

- NA NA: The gene set is unrepresentative in the used gene list
- 1 1: No significant different of mRNA expression between the two phenotypes

Limitations:

- Use categorical phenotype (not numeric) only with TWO classes
- Use crude and not normalised mRNA expression data
- The accuracy of the results is better with high number of genes (1000 - 10000) and 1000 permutations. This request takes a while to run. User is recommended to reduce the permutation to 100 instead 1000 and/or reduce the size of gene list.
- If two or more phenotypes are checked, only the latter is taken.

4.8.1 Which Molecular Signature Data base (MSigDB) for gene list

This function matches genes list with MSigDB files selected from example or directory (Figure 19) and computes the mean of matched Genes by Gene Sets as:

$$Mean_{(Gene \in GeneSet)} = \frac{\sum_{i=1}^I \sum_{j=1}^J Gene_i \in GeneSet_j}{\sum_{i=1}^I Gene_i}$$

With I is the number of genes in the list and J is the number of Gene Sets in the MSigDB file. The returned table indicates the mean of gene number that matched for every Gene Set. For example in the first row (Figure 19), there are about 10 genes that mached for every Genes Set in c2.cp.reactome.v4.0.Symbol.gmt.

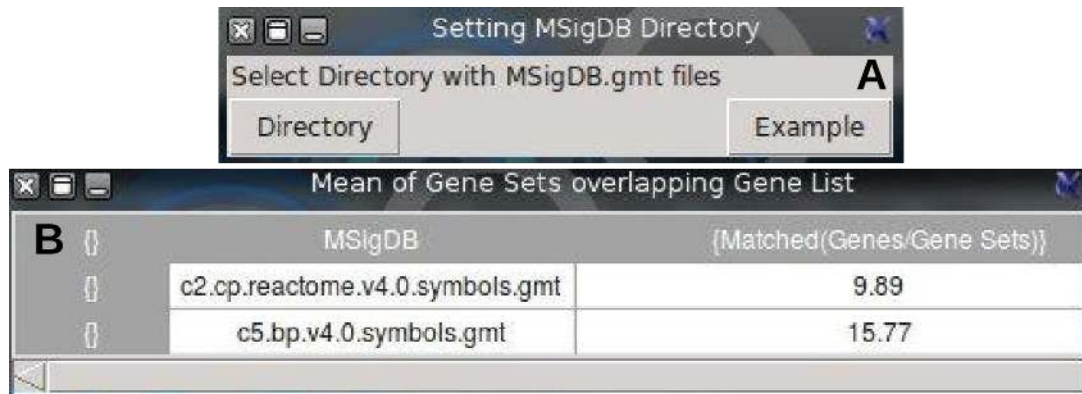


Figure 19: Match gene list with MSigDB folder.

4.8.2 get SubMSigDb for genes list

The SubMSigDb is a subset of MSigDB generated from gene list in the expression Set (eSet). This specific subsetting reduces the time of GSEA computing. User needs to run before eSet from phenoTest menu.

4.8.3 GSEA1m: Phenotypes into Disease

Disease Free Status (DFS_STATUS) into Prostate Cancer:

- Study: Prostate Adenocarcinoma (TCGA, Provisional)
- Cases: All Samples with mRNA expression data (246 samples),
- Genetic Profiles: mRNA expression (RNA Seq V2 RSEM)
- MSigDB: c2.cp.reactome.v4.0.symbols.gmt
- Gene list: 73.txt file
- Phenotype: DFS_STATUS
- Permutation: 1000
- pVal: 0.05

This example uses Reactome pathways gene sets (Figure 20A, <http://www.reactome.org/>) to compare the gene expression of patients with reccured prostate disease versus patient with free prostate disease. The run was done with 1000 permuatation and pVal 0.05 (Figure 20B). The result (Figure 20C) shows only up regulated gene sets from Reactome pathways were observed in disease free patients.

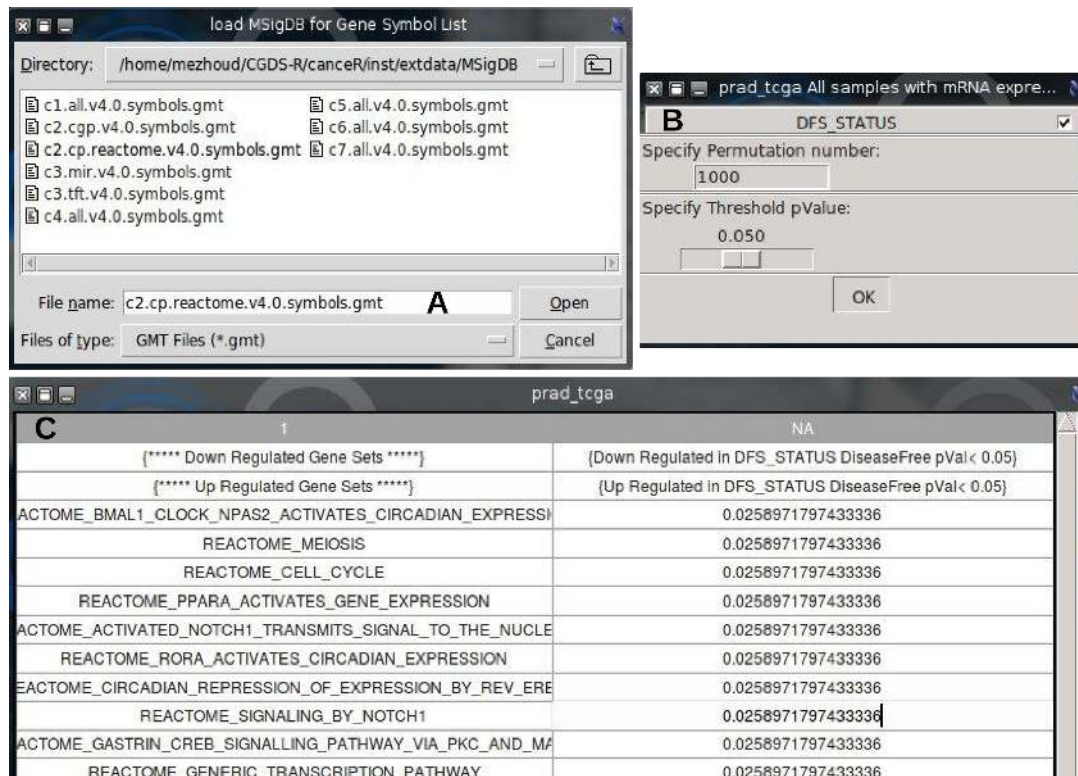


Figure 20: Linear model of GSEA of Prostate Adenocarcinoma (TCGA, Provosional) study using "DiseaseFree/Reccured" phenotypes. A, Selecting MSigDB file; B, Selecting Phenotypes, permutation, and pVal parameters; C, displaying down and up regulated gene sets in table.

Copy Number Cluster Level into Stomach Adenocarcinoma:

- Study: Stomach Adenocarcinoma (TCGA, Nature 2014)
- Cases: All Samples that have mRNA, CNA, and sequencing data (258 samples),
- Genetic Profiles: mRNA expression (RNA Seq V2 RSEM)
- MSigDB: c5.bp.reactome.v4.0.symbols.gmt
- Gene list: 73.txt file
- Phenotype: Copy_Number_Cluster (Low, High)
- Permutation: 1000
- pVal: 0.05

The losses and the gains of DNA can contribute to alterations in the expression of tumor suppressor genes and oncogenes. Therefore, the identification of DNA copy number alterations in tumor genomes may help to discover critical genes associated with cancers and, eventually, to improve therapeutic approaches. In this study we test GSEAlm algorithm with the level of the Copy Number cluster using Biological Process gene sets from Reactome. The Figure 21C shows significant down/up regulated gene sets in patients with low level compared to patients with high level of copy number cluster.

This finding suggests to repeat modeling with copy number alteration profile to predict which biological process (gene set) could be with low level of copy number cluster. The Figure 22D lists gene sets with low level of copy number cluster in the case of stomach adenocarcinoma.

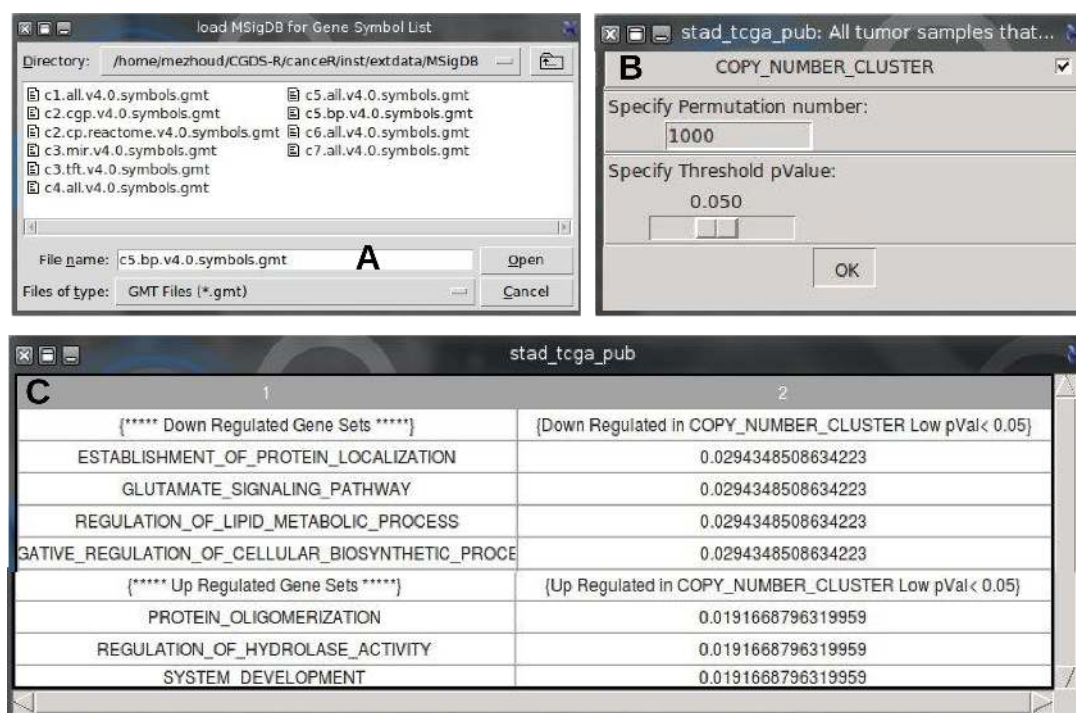


Figure 21: Linear model of GSEA of Stomach Adenocarcinoma (TCGA, Nature 2014) study using "Low/High" level of copy number cluster phenotypes. A, Selecting MSigDB file; B, Selecting Phenotypes, permutation, and pVal parameters; C, displaying down and up regulated gene sets in table.

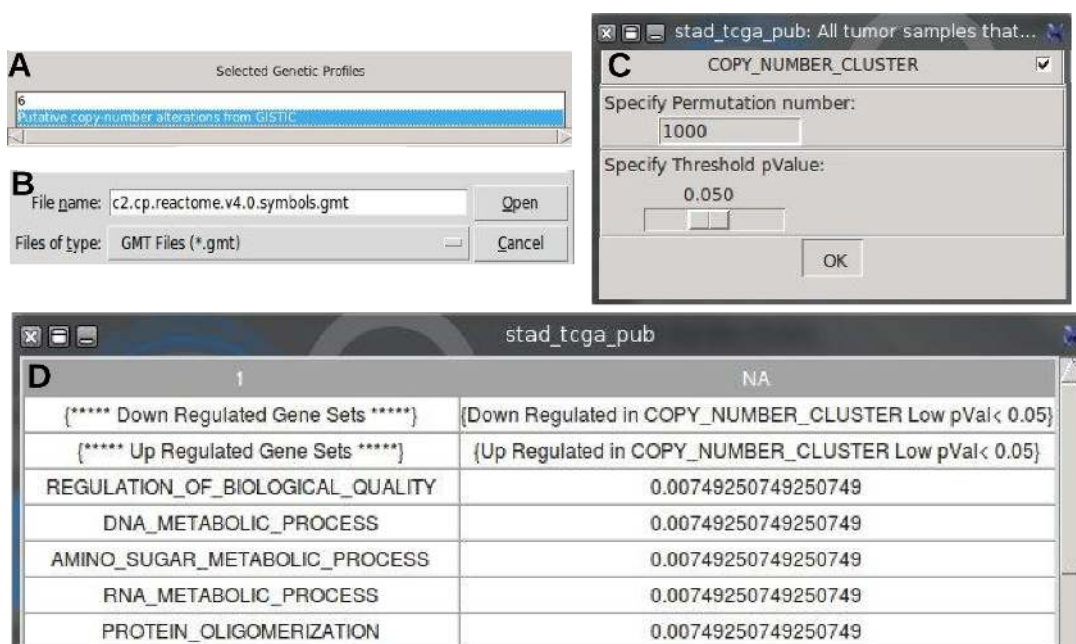


Figure 22: Linear model of GSEA of Stomach Adenocarcinoma (TCGA, Nature 2014) study using copy number alteration data (GISTIC algorithm) as matrix and copy number cluster level (Low/High) as phenotypes. A, Selecting genetic profile; B, Selecting MSigDB file; C, Selecting Phenotypes, permutation, and pVal parameters; D, displaying gene sets with low copy number cluster level.

4.8.4 GSEA1m: Disease vs Disease

Breast vs Prostate Cancers: In this function, the linear modeling uses the disease type as phenotype for the run. User needs to select two diseases and a gene list.

- Studies: Breast Invasive Carcinoma (TCGA, Provisional) versus Prostate Adenocarcinoma (TCGA, Provisional)
- Cases: All Samples with mRNA expression data (959/257 samples),
- Genetic Profiles: mRNA expression (RNA Seq V2 RSEM)
- MSigDB: c2.cp.reactome.v4.0.symbols.gmt
- Gene list: 73.txt file
- Phenotype: Diseases type
- Permutation: 1000
- pVal: 0.05
- Samples number: 50

The Figure 23 shows the results of the linear modeling.

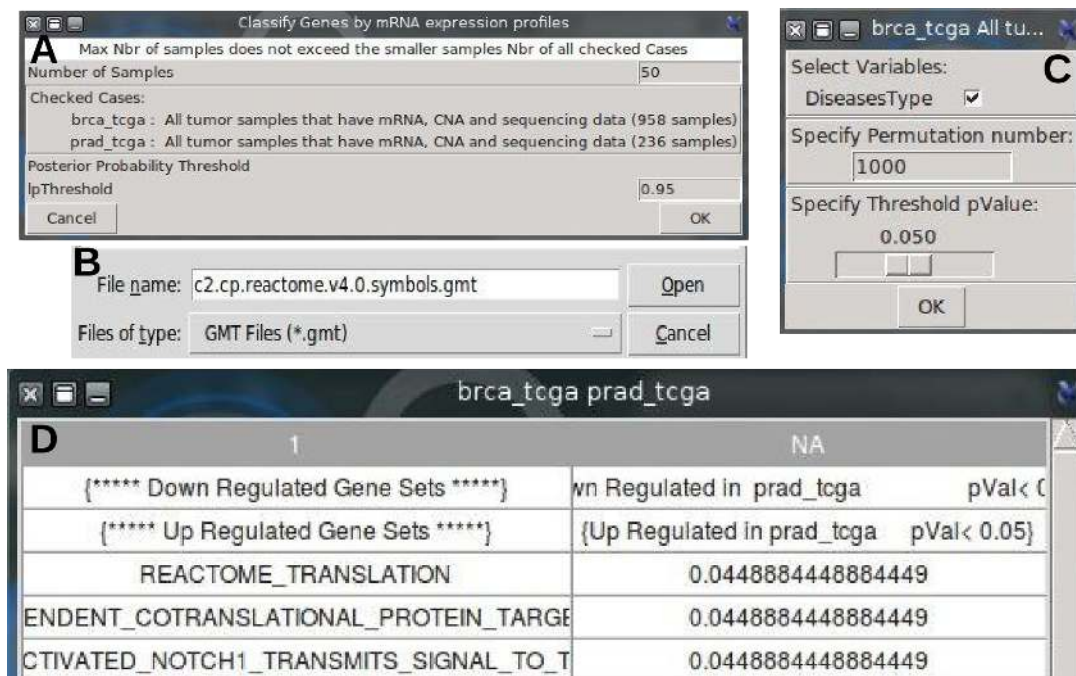


Figure 23: Linear model of GSEA of Breast Invasive carcinoma (TCGA, Provisional) and Prostate Adenocarcinoma (TCGA, Provisional) studies. A, Sampling; B, Selecting MSigDB file; c, Selecting Phenotypes, permutation, and pVal parameters; D, displaying gene sets upregulated in prostate cancer.

4.9 Genes Classification using mRNA expression (Classification)

The **Classification** menu displays two functions to rank genes by phenotypes into- and inter-diseases, depending on mRNA expression data.

4.9.1 Genes vs Diseases (inter-diseases)

The first classifier is implemented from `geNetClassifier` package [1]. It uses `calculateGenesRanking` function which based on Parametric Empirical Bayes method included in `EBarrays` package [5]. This method implements an expectation-maximization (EM) algorithm for gene expression mixture models, which compares the patterns of differential expression across multiple conditions and provides a *posterior probability*. The posterior probability is calculated for each gene-class pair, and represents how much each gene differentiates a class from the other classes; being 1 the best value, and 0 the worst. In this

way, the posterior probability allows to find the genes that show significant differential expression when comparing the samples of one class versus all the other samples (One-versus-Rest comparison) [1].

In the following examples, we would like to predict specific modulated genes by cancer type. The features of the runs are:

Example 1: Breast *vs* Glioblastoma *vs* Liver *vs* Lung Cancers (Figure 24A):

- Studies:
 - Breast Invasive Carcinoma (TCGA, Provisional)
 - Glioblastoma Multiforme (TCGA, Provisional)
 - Liver Hepatocellular Carcinoma (TCGA, Provisional)
 - Lung Squamous Cell Carcinoma (TCGA, Provisional)
- Cases: All Samples with mRNA expression data,
- Genetic Profiles: mRNA expression (RNA Seq V2 RSEM)
- Samples: 50
- Gene list: 223.txt file
- lpThreshold: 0.95

Example 2: Bladder *vs* Breast *vs* Glioblastoma *vs* Lung *vs* Ovarian *vs* Prostate Cancers (Figure 24A’):

- Studies:
 - Bladder Urothelial Carcinoma(TCGA, Provisional)
 - Breast Invasive Carcinoma (TCGA, Provisional)
 - Glioblastoma Multiforme (TCGA, Provisional)
 - Lung Adenocarcinoma (TCGA, Provisional)
 - Ovarian Serous Cystadenocarcinoma (TCGA, Provisional)
 - Prostate Adenocarcinoma (TCGA, Provisional)
- Cases: All Samples with mRNA expression data,
- Genetic Profiles: mRNA expression (RNA Seq V2 RSEM)
- Samples: 50
- Gene list: 73.txt file
- lpThreshold: 0.95

The significant genes (dots) are over the threshold of posterior probability (Figure 24B, B’). Their number are listed in the legend. The details are displayed in table (Figure 24C, C’) and saved in /Results/Classifier folder.

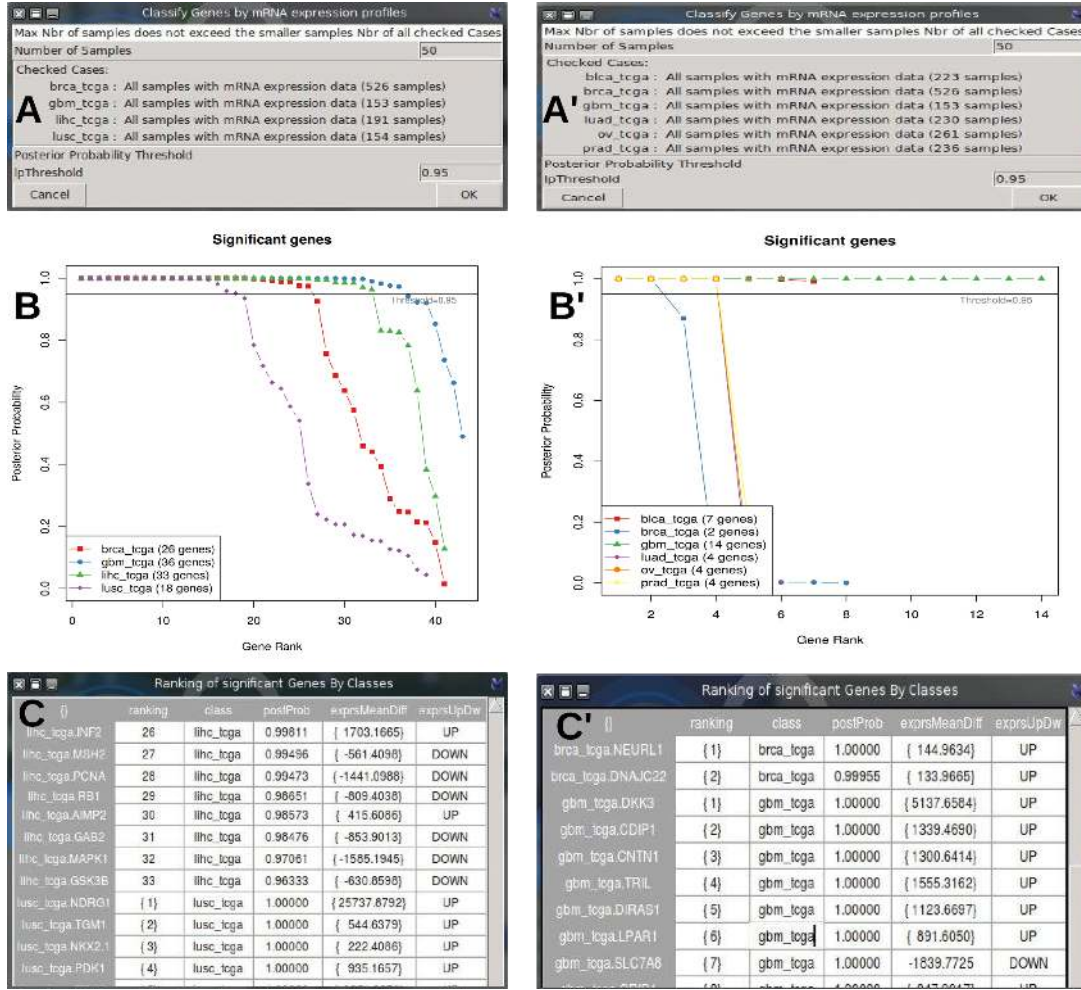


Figure 24: Genes/Disease Classification. A, A' Sampling and lpThreshold choice; B, B' Plot of the posterior probabilities of the genes of Disease classes, ordering the genes according to their rank. brca: breast; gbm: Glioblastoma; lihc: Liver; lusc: Lung; blca: Bladder; luad: Lung; ov: Ovary; prad: Prostate. C, C' The table lists the significant genes ranked by class (disease). The columns are ordered as Gene Symbols, Ranking, Class, PostProb, ExprsMeanDiff, ExprsUpDw.

4.9.2 Genes vs Phenotypes (intra-disease)

The second function of genes classification is implemented from **rpart** package [11]. The resulting models can be represented as binary trees with gene expression profile threshold ($P53 > 2.56$) is in node and classes (Living/Deceased) of selected variable in branch. The root of the tree is the best gene divisor of classes. The goal is to predict which genes combinaison ($P53 > 2.56 + CAD < -0.45 + MDM4 > 1.92$ lead to 80% Diseased) could be split classes in two or more groups. The tree is built by the following process: first the single variable is found which gene level splits the data into two groups. The data is separated, and then this process is applied separately to each sub-group, and so on recursively until the subgroups either reach a minimum size or until no improvement can be made [11].

There are 3 methods of splitting rule : **classification**, **anova**, and **Poisson**.

The **Classification** used either Gini or log-likelihood function rules. It is used when there are only two categories into the selected variable. The dependent variable is nominal (factor). At each splitting step, the rule tries to reduce the total impurity of the two son nodes relative to the parent node.

In the **anova**, the variable is numeric and it used to predict the closest value to the true one. The method uses splitting criteria $SS_T - (SS_L + SS_R)$, where $SS_T = \sum (y_i - \bar{y})^2$ is the sum of squares for the node, and SS_R, SS_L are the sums of squares for the right and left son respectively. This is equivalent to choosing the split to maximize the between-groups sum-of-squares in a simple analysis of variance [11].

The **poisson** splitting method attempts to extend **rpart** models to event rate data. The model in this case is $\lambda = f(x)$, where λ is an event rate and x is some set of predictors.

Example 1: Genes classification *vs* OS_STATUS (Living/Deceased)

- Studies: Breast Invasive Carcinoma (TCGA, Provisional)
- Cases: All Samples with mRNA expression data (526 samples),
- Genetic Profiles: mRNA expression z-score (RNA Seq V2 RSEM)
- Gene list: 223.txt file
- Variable: OS_STATUS
- Split method: `class`

In this example, the main clinical endpoint of interest is the Living/Deceased Status of patient having breast cancer. We run classification to predict which genes from 223 that make the difference between deceased and living patient having breast cancer. The tree in Figure 25B indicates that all deceased patients have the following z-score gene expression profile: $RAD51 < -1.091$, $HSPA1A > 1.424$, $JUN > -0.4016$. All remain patient are living.

Example 2: Genes regression *vs* OS_STATUS (Living/Deceased)

- Studies: Breast Invasive Carcinoma (TCGA, Provisional)
- Cases: All Samples with mRNA expression data (526 samples),
- Genetic Profiles: mRNA expression z-score (RNA Seq V2 RSEM)
- Gene list: 223.txt file
- Variable: OS_STATUS
- Split method: `anova`

When we split the same clinical data used in the first example (Living/Deceased) using regression method (`anova`), we see in the Figure 26C the same genes and split points with further splitting nodes. For example, the gene HSPA11 has 124 patients with 109 are living. In classification method, all patient have the same predicted value (0.846, Figure 25C) because the error (misclassification) with and without the split is identical. In the regression context the two predicted values of 16.08 and 1.84 (Figure 26B) are different. The split has identified a nearly pure subgroup of significant size.

Example 3: Genes classification *vs* tumor grade (grade1/2/3)

- Studies: Uterine Corpus Endometrioid Carcinoma (TCGA, Nature 2013)
- Cases: All Samples with mRNA expression data (333 samples),
- Genetic Profiles: mRNA expression z-score (RNA Seq V2 RSEM)
- Gene list: 223.txt file
- Variable: Tumor Grade
- Split method: `class`

Classification method could work with phenotype with more than two classes. In this case, the tumor grades are splitted following this order from left to right Grade1/Grade2/Grade3. The nodes are named by the must frequent grade (Figure 27).

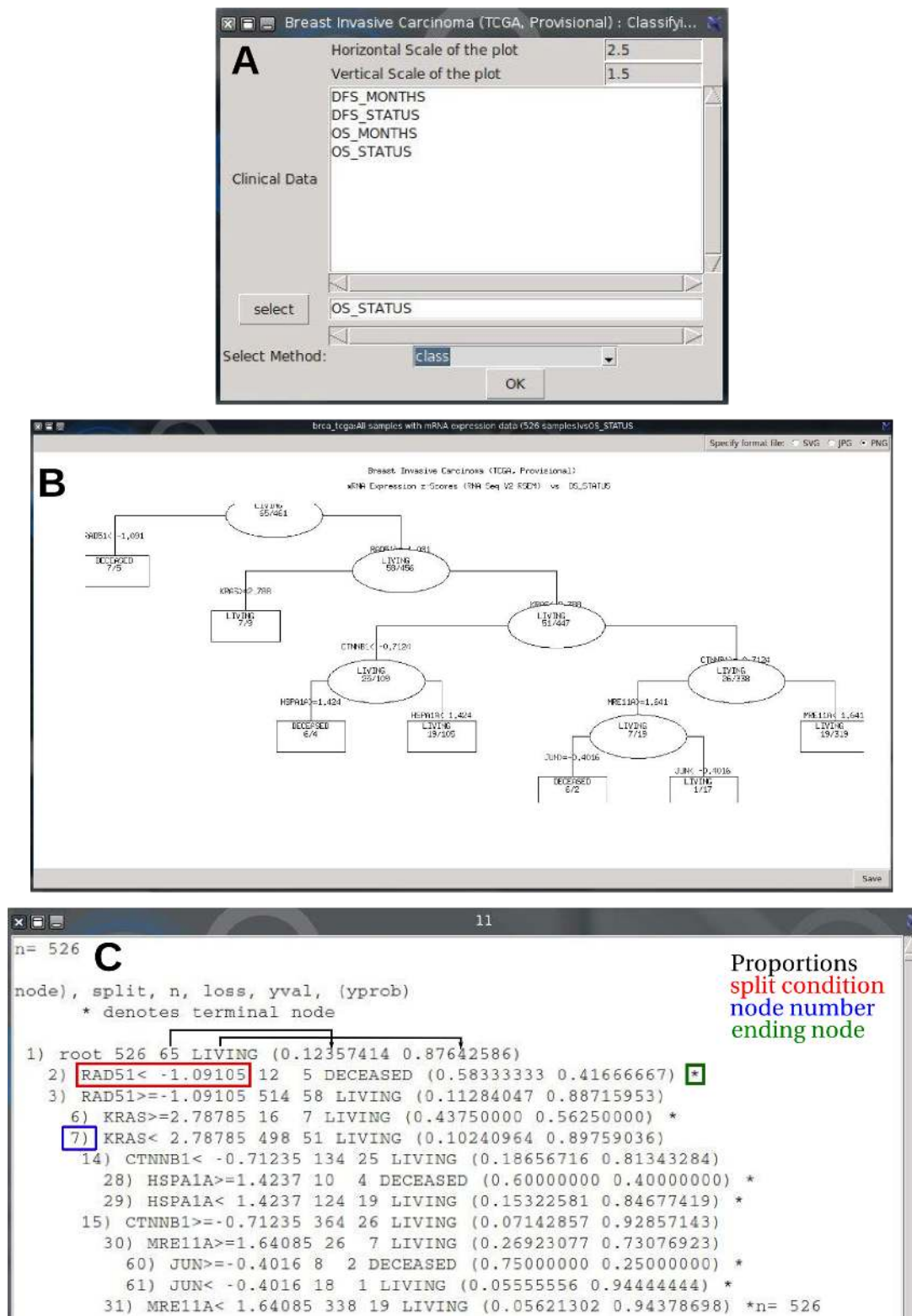


Figure 25: Genes/phenotype Classification. A, Selecting variable and the splitting method; B, Tree result, In any node (ellipse or rectangle) the ratio indicates in numerator/Deceased and in the denominator/Living patients. The condition RAD51 z-score < -1.091 discriminates 12 cases with 7 deceased and so on. The Deceased cases go to the left and the living cases go to the right. The plot could be saved as svg, png or jpg format and scaled in vertical and horizontal; C, The reading of the tree.

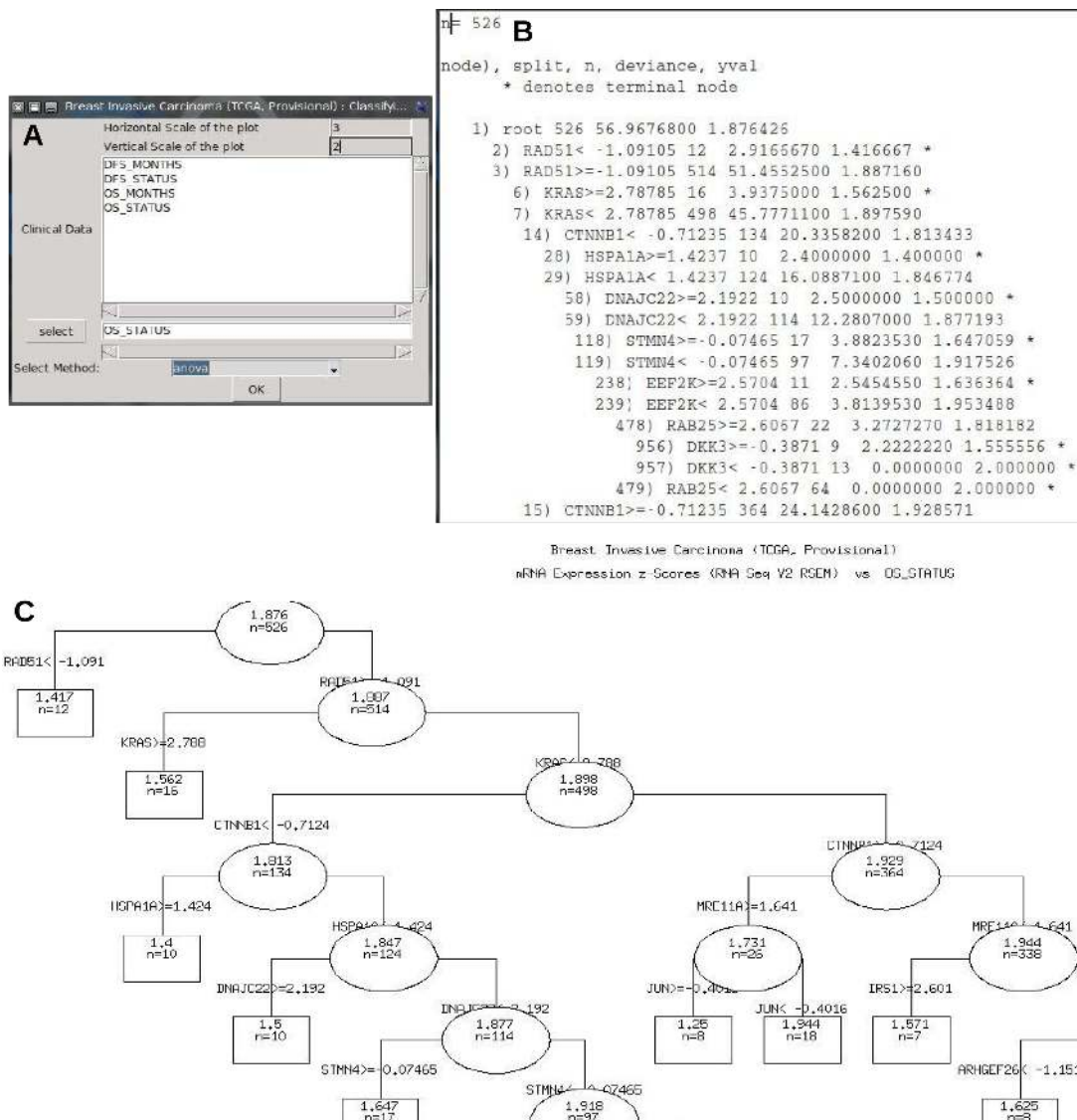


Figure 26: Genes/phenotype Regression. A, Selecting variable and the splitting method; B, The reading of the tree; C, Tree result, In any node (ellipse or rectangle) the ratio indicates in numerator/Deceased and in the denominator/Living patients.

4.10 Plots

There are two plotting functions. The first one (1 Gene/ 2 Gen. Profiles) plots gene data for specified case and two genetic profiles from the same study. It associates between the two selected genetic profiles. The dialog box allows user to specify:

- Layout Skin
 - cont: This is the default skin. It treats all data as being continuous
 - disc.cont: Requires two genetic profiles. The first dataset is handled as being discrete data, and the function generates a boxplot with distributions for each level of the discrete genetic profile.
 - cna_mrna_mut : This skin plots mRNA expression level as function of CNA or DNA methylation status for given gene. Data points are colored respectively by mutation status or CNA and mutation status.
- Correlation Method

No significant pearson correlation was observed between P53 Copy Number Variation (CNV) and P53 mRNA expression ($r=0.23$, Figure 28B). The CNV is ranged into 4 levels that derived from the copy-number analysis algorithms GISTIC or RAE, and indicate the copy-number level per gene. "-2" is a deep loss, possibly a homozygous deletion (Homdel), "-1" is a single-copy loss (heterozygous deletion: Hetloss), "0" is Diploid, "1" indicates a low-level gain, and "2" is a high-level amplification. Note that these calls are putative.

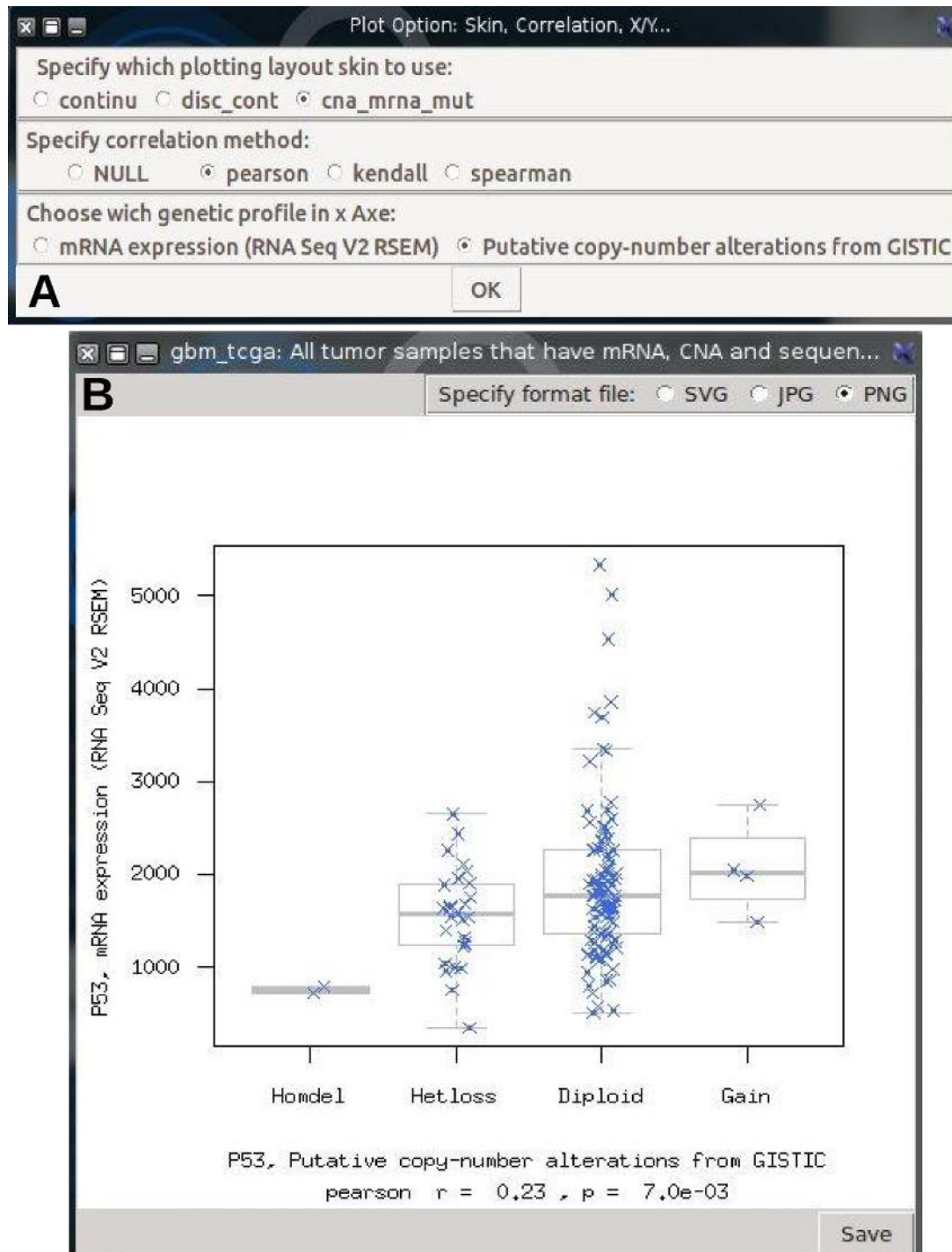


Figure 28: P53: CNA status vs mRNA expression. Homdel, homozygous deletion; Hetloss, heterozygous deletion; Diploid, No modification; Gain, ADN amplification.

The second plot function evaluates the relationship of two genes expression levels in the same study (Figure 29).

Example: MAP2K2 and ABHD17A mRNA expression (RNA Seq V2 RSEM) levels in Uterine Corpus Endometrioid Carcinoma (TCGA, Nature 2013)

- Study: Uterine Corpus Endometrioid Carcinoma (TCGA, Nature 2013)
- Cases: All Samples with mRNA expression data (333 samples),
- Genetic Profiles: mRNA expression (RNA Seq V2 RSEM)
- Genes: MAP2K2 and ABHD17A
- correlation method: pearson

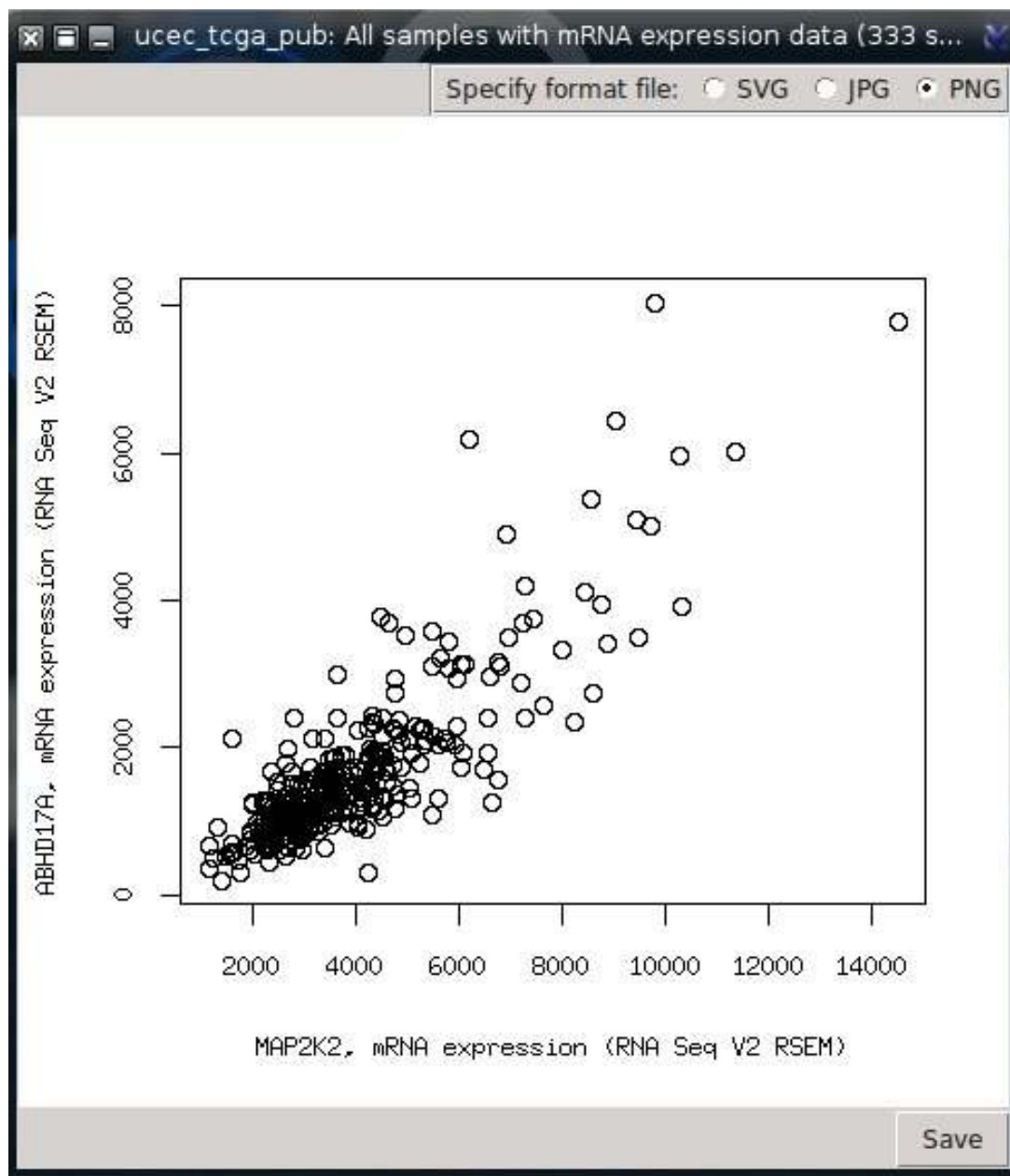


Figure 29: MAP2K2 and ABHD17A mRNA expression levels correlation in Uterine Corpus Endometrioid Carcinoma

5 Perspectives

1. Pairwise, Multiple comparative of Genetics Profiles

2. search nearest Cancer by Matching Genetic Profile for gene list.
3. Onco Query Language

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