CancerMutationAnalysis package

Giovanni Parmigiani Dana-Farber Cancer Institute and Harvard School of Public Health email: gp@jimmy.harvard.edu,

Simina M. Boca Georgetown University Medical Center email: smb310@georgetown.edu

May 2, 2019

Contents

L	Overview	1
2	Data objects	2
3	Calculating gene scores	3
1	Estimating passenger probabilities	3
5	Performing gene-set analyses	4
3	Simulating data for gene-sets 6.1 Manipulating the SetMethodsSims objects	8
7	Acknowledgments	9

1 Overview

This package contains software which implements a variety of methods for gene and gene-set level analysis in somatic mutation studies of cancer. The methods implemented are described in [12] and [8], for the gene level analysis, and [1], for the gene-set level analysis. The gene level methods considered were developed to distinguish between "driver genes" and "passenger genes;" the former play an active role in tumorigenesis, having mutations which are selected for in cancer, while the latter are mutated in tumor samples, but have no role in tumorigenesis. They incorporate the study design used in [12, 13, 4, 9, 10]: The complete collection of coding exons (the exome) was sequenced in several "discovery samples," with a smaller number of genes being further sequenced in a larger number of "prevalence samples," based on the results of the discovery screen, and possibly, on prior knowledge about their role in cancer.

The gene-set methods implemented carry out the "patient-oriented" approach, which calculates gene-set scores for each sample, then combines them across samples. By comparison, gene-oriented

methods calculate a score for each gene across all samples, then combine them into gene-set scores. The four patient-oriented methods described in [1] are included, as well as the gene-oriented method described in [7] (see also [11]), which is in the limma package and performs a Wilcoxon test. Data from several comprehensive studies are also distributed with this package [12, 13, 4, 9, 10].

This vignette represents an introduction to the CancerMutationAnalysis package. The primary functions are: cma.scores, which calculates a variety of gene scores; cma.fdr, which estimates passenger probabilities and false discovery rates; and cma.set.stat, which implements a number of gene-set level methods. The function cma.set.sim performs gene-set simulations using either the permutation null or the passenger null (see [1]). The functions extract.sims.method and combine.sims are used to manipulate the objects resulting from cma.set.sim.

2 Data objects

We first load the data and look at them:

- > library(CancerMutationAnalysis)
- > data(WoodBreast07)
- > data(WoodColon07)
- > data(JonesPancreas08)
- > data(ParsonsGBM08)
- > data(ParsonsMB11)

The datasets are labelled according to the publication year of the study and the tumor type. Thus, data is available for studies of breast cancer, colon cancer, pancreatic cancer, glioblastoma multiforme (GBM), and medulloblastoma (MB).

The objects containing information on the somatic mutations are labelled GeneAlter*. As an example, consider the GeneAlterGBM object:

> head(GeneAlterGBM)

	Gene	Туре	${\tt Sample}$	${\tt Screen}$	${\tt WTNuc}$	${\tt Context}$	${\tt MutNuc}$
1	MRPL55@@15731	Mut	BR11P	Disc	C	CpG	T
2	USH2A@@45	Mut	BR20P	Disc	G	${\tt GpA}$	T
3	KCNH1@@6084	Mut	BR09P	Disc	G		T
4	LAMB3@@1338	Mut	BR17X	Disc	G		A
5	PIGR@@7060	Mut	BR08X	Disc	G	CpG	A
6	REN@@7303	Mut	BR20P	Disc	G	CpG	Α

The columns represent the transcript, mutation type (Mut, Amp, Del, for point mutations, amplifications, and deletions, respectively), sample ID, screen (Disc or Prev for "Discovery" or "Prevalence"), wild type nucleotype, context, and nucleotide after mutation. Only point mutations are used to calculate gene scores, estimate passenger probabilities, and perform gene-set level analyses. Only discovery samples are used to perform the set analyses.

The GeneCov* objects contain data on the number of nucleotides of each type successfully sequenced for each transcript:

> head(GeneCovGBM)

```
Gene Screen WTNuc Context Coverage 1 15E1.2@013849 Disc C CpG 387.0147
```

```
2 15E1.2@@13849
                   Prev
                             C
                                    CpG
                                          0.0000
3 15E1.2@@13849
                   Disc
                             G
                                    CpG 465.2525
4 15E1.2@@13849
                   Prev
                             G
                                    CpG
                                          0.0000
5 15E1.2@013849
                   Disc
                             G
                                    GpA 470.0150
6 15E1.2@@13849
                             G
                   Prev
                                    GpA
                                          0.0000
```

The GeneSamp* objects store the number of discovery and prevalence samples for each transcript:

> head(GeneSampGBM)

	Gene	Screen	NrSamp
1	15E1.20013849	Disc	21
2	15E1.20013849	Prev	0
3	2'-PDE@@16564	Disc	21
4	2'-PDE@@16564	Prev	0
5	3'HEXO@@9816	Disc	21
6	3'HEXO@@9816	Prev	0

The BackRates* objects contain estimated passenger probabilities used in the original studies.

3 Calculating gene scores

Gene scores are obtained using the cma.scores function. These scores include the Cancer Mutation Prevalence (CaMP) score [8] and the log Likelihood Ratio (logLRT) score [3]. They take into account the mutation profile and the total number of nucleotides successfully sequenced for each gene, as well as an estimated "background passenger rate" of mutation:

```
> ScoresGBM <- cma.scores(cma.alter = GeneAlterGBM,
                           cma.cov = GeneCovGBM,
+
                           cma.samp = GeneSampGBM,
                           passenger.rates = BackRatesGBM["MedianRates",])
> head(ScoresGBM)
                   CaMP
                           logLRT
A2M@@408
              0.9028709 2.158645
A4GALT@@21470 0.5706862 1.678895
ABCA10@0388
              1.0423011 2.575103
              0.7044877 1.853172
ABCA4@@140
ABCA7@@194
              0.1076564 1.157538
ABCB6@2071
              0.5499152 1.655159
```

4 Estimating passenger probabilities

Passenger probabilities for individual genes are estimated using the cma.fdr function, which performs an empirical Bayes (EB) analysis. The output of this function is a list, each score having an entry which, for each gene, gives the corresponding score, the number of genes with scores greater or equal in the dataset (F), the average number of genes with scores greater or equal in the simulated datasets (F0), the estimated false discovery rate (Fdr), the estimated local false discovery

rate (fdr), and the estimated value of the prior probability of the gene being null (p0). The estimated passenger probabilities for individual genes are the estimate local false discovery rates. If showFigure=TRUE, a plot is also generated for each score, displaying the right tail of the density of null scores and a 1-D "rug plot" histogram of the observed scores. A cutoff is chosen so that the false discovery rate has the largest possible value below it. See Figure 1 for an example.

```
> set.seed(188310)
> FdrGBM <- cma.fdr(cma.alter = GeneAlterGBM,
                      cma.cov = GeneCovGBM,
                      cma.samp = GeneSampGBM,
                      scores = "logLRT",
                      passenger.rates = BackRatesGBM["MedianRates",],
                      showFigure=TRUE,
                     cutoffFdr=0.1,
                     M = 5
Score: logLRT.
> head(FdrGBM[["logLRT"]])
                               Score
                                              FO Fdr fdr p0
PCOTH@@18196
                       5.837195e-05 20671 20671
                                                        1
PMS2L3@@9470
                       9.693291e-05 20670 20670
                                                        1
                                                           1
ENST00000360216@@16327 1.011182e-04 20669 20669
CCL4L2@@14948
                       1.029200e-04 20668 20668
                                                        1
                                                           1
Q6PDB4_HUMAN@@17521
                       1.036599e-04 20667 20667
                                                        1
                                                           1
L0C149643@@15875
                        1.085358e-04 20666 20666
```

5 Performing gene-set analyses

The function cma.set.stat returns a data-frame with p-values and q-values for all the methods selected. By default, all the methods are implemented; however this takes several minutes. Setting the gene.method parameter to TRUE implements the gene-oriented method. The other method parameters refer to the patient-oriented methods: perm.null.method refers to the permutation null without heterogeneity, perm.null.het.method to the permutation null with heterogeneity, pass.null.method to the passenger null without heterogeneity, and pass.null.het.method to the passenger null with heterogeneity. See [1] for more details on all these methods and [6] for a related method.

We use gene-sets from KEGG [5] in this example through the KEGG.db package [?]:

```
> library(KEGG.db)
> KEGGPATHID2EXTID

PATHID2EXTID map for KEGG (object of class "AnnDbBimap")
```

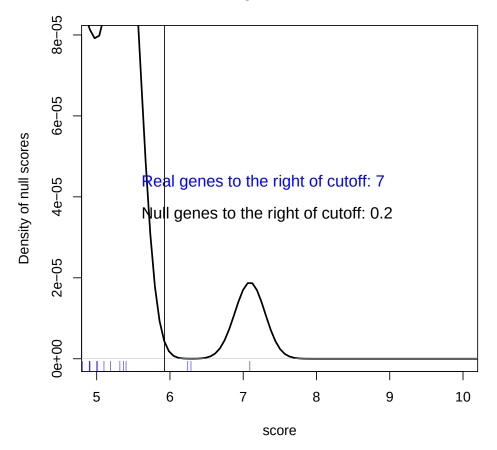
Since the genes in the GBM datasets are annotated by gene names, while KEGGPATHID2EXTID uses EntrezGene identifiers, we also need to load the vector mapping the identifiers to the names, which we obtained by using the biomaRt package [2].

Figure 1: Example of figure obtained by setting ${\tt showFigure=TRUE}$ in the ${\tt cma.fdr}$ function.

Score: logLRT.

	Score	F	FO	Fdr	fdr	p0
PCOTH@@18196	5.837195e-05	20671	20671	1	1	1
PMS2L3@09470	9.693291e-05	20670	20670	1	1	1
ENST00000360216@@16327	1.011182e-04	20669	20669	1	1	1
CCL4L2@@14948	1.029200e-04	20668	20668	1	1	1
Q6PDB4_HUMAN@@17521	1.036599e-04	20667	20667	1	1	1
LOC149643@@15875	1.085358e-04	20666	20666	1	1	1

logLRT score



> data(EntrezID2Name)

We now use the function cma.set.stat to perform the gene-set analyses on three KEGG sets:

```
> as.character(KEGGPATHNAME2ID[c("Endometrial cancer",
                                  "Non-small cell lung cancer",
                                  "Alanine, aspartate and glutamate metabolism")])
Alanine, aspartate and glutamate metabolism
                         Endometrial cancer
                                     "05213"
                 Non-small cell lung cancer
                                     "05223"
> SetResultsGBM <-
    cma.set.stat(cma.alter = GeneAlterGBM,
+
                 cma.cov = GeneCovGBM,
+
                 cma.samp = GeneSampGBM,
                 GeneSets = KEGGPATHID2EXTID[c("hsa05213",
                   "hsa05223",
                                 "hsa00250")],
                 ID2name = EntrezID2Name,
                 gene.method = FALSE,
                 perm.null.method = TRUE,
                 perm.null.het.method = FALSE,
                 pass.null.method = TRUE,
                 pass.null.het.method = FALSE)
```

The resulting object is a data-frame containing p-values and q-values for the methods which were implemented:

> SetResultsGBM

```
p.values.perm.null q.values.perm.null p.values.pass.null
hsa05213
               9.198666e-16
                                   2.759600e-15
                                                       2.108377e-10
hsa05223
               1.678783e-15
                                   2.518175e-15
                                                       3.403831e-11
               3.109024e-01
                                   3.109024e-01
                                                       6.340137e-01
hsa00250
         q.values.pass.null
hsa05213
               3.162565e-10
hsa05223
               1.021149e-10
hsa00250
               6.340137e-01
```

6 Simulating data for gene-sets

We now demonstrate the use of the cma.set.sim function, which simulates datasets under either the permutation or passenger null (see [1]), depending on whether pass.null is set to TRUE or FALSE, and calculates the p-values and q-values for those datasets for the selected methods. The simulations may also include spiked-in gene-sets, by using the perc.samples and spiked.set.sizes options. For example, if one desires to have two spiked-in gene-sets, both having 50 genes, but one having the

probability of being altered in any given sample of 0.75 and the other of 0.95, then these parameters should be set to perc.samples = c(75, 90) and spiked.set.sizes = 50. The spiked-in genesets are artificially created with hypothetical genes (for more details on how they are generated, see [1]). To simulate the data without spiked-in sets, under the permutation or passenger null hypotheses, the parameters should be set as following: perc.samples = NULL, spiked.set.sizes = NULL. The object outputted by cma.set.sim is of the class SetMethodsSims. Note that this code takes several minutes to run:

```
> set.seed(831984)
> resultsSim <-
      cma.set.sim(cma.alter = GeneAlterGBM,
                  cma.cov = GeneCovGBM,
                  cma.samp = GeneSampGBM,
                  GeneSets = KEGGPATHID2EXTID[c("hsa05213",
                     "hsa05223", "hsa00250")],
                  ID2name = EntrezID2Name,
                  nr.iter = 2,
                  pass.null = TRUE,
                  perc.samples = c(75, 95),
                  spiked.set.sizes = c(50),
                  show.iter = TRUE,
                  gene.method = FALSE,
                  perm.null.method = TRUE,
                  perm.null.het.method = FALSE,
                  pass.null.method = TRUE,
                  pass.null.het.method = FALSE)
> resultsSim
Simulation results for gene-set analysis of mutations
  Data-generating mechanism : Passenger null
  Number of simulations
  Number of gene-sets
                             : 5
     Original : 3
     Spiked-in: 2
  Spiked-in sets
     Probability of being mutated in a set : 0.75 0.95
     Length (as number of genes)
                                            : 50
> slotNames(resultsSim)
                                           "spiked.set.sizes" "GeneSets"
[1] "null.dist"
                        "perc.samples"
[5] "cma.alter"
                        "cma.cov"
                                           "cma.samp"
                                                               "Scores"
[9] "results"
> resultsSim@null.dist
[1] "Passenger null"
```

6.1 Manipulating the SetMethodsSims objects

> extract.sims.method(resultsSim,

We provide 2 functions to help manipulate the SetMethodsSims objects which result from the cma.set.sim function: extract.sims.method and combine.sims. The extract.sims.method function is used to obtain a single data frame with the p-values or q-values from one of the specific methods. For instance, the command to get the p-values for the permutation null with no heterogeneity method is:

```
"p.values.perm.null")
                        [,1]
                                     [,2]
               5.672050e-01 3.026478e-01
hsa05213
hsa05223
               5.925646e-01 3.264250e-01
hsa00250
               5.593495e-01 8.459466e-01
gene.set.50.75 1.688373e-06 1.217415e-06
gene.set.50.95 3.680435e-14 4.736465e-18
The function combine.sims may be used to combine 2 simulations:
> combine.sims(resultsSim, resultsSim)
Simulation results for gene-set analysis of mutations
  Data-generating mechanism : Passenger null
  Number of simulations
  Number of gene-sets
                             : 5
     Original: 3
     Spiked-in: 2
  Spiked-in sets
     Probability of being mutated in a set: 0.75 0.95
     Length (as number of genes)
                                            : 50
```

References

- [1] S.M. Boca, K.W. Kinzler, V.E. Velculescu, B. Vogelstein, and G. Parmigiani. Patient-oriented gene set analysis for cancer mutation data. *Genome Biology*, 11(11):R112, 2010.
- [2] Steffen Durinck and Wolfgang Huber. biomaRt: Interface to BioMart databases (e.g. Ensembl, COSMIC, Wormbase and Gramene). R package version 2.8.1.
- [3] G. Getz, H. Höfling, J. P. Mesirov, T. R. Golub, M. Meyerson, R. Tibshirani, and E. S. Lander. Comment on "the consensus coding sequences of human breast and colorectal cancers". *Science*, 317(5844):1500, 2007.
- [4] S. Jones, X. Zhang, D.W. Parsons, J.C.H. Lin, R.J. Leary, P. Angenendt, P. Mankoo, H. Carter, H. Kamiyama, A. Jimeno, et al. Core signaling pathways in human pancreatic cancers revealed by global genomic analyses. *Science*, 321(5897):1801, 2008.
- [5] M. Kanehisa and S. Goto. KEGG: Kyoto encyclopedia of genes and genomes. Nucleic Acids Research, 28(1):27, 2000.

- [6] J. Lin, C.M. Gan, X. Zhang, S. Jones, T. Sjöblom, L.D. Wood, W. Parsons, N. Papadopoulos, K.W. Kinzler, B. Vogelstein, G. Parmigiani, and V.V. Velculescu. A multidimensional analysis of genes mutated in breast and colorectal and colorectal cancers. *Genome Research*, 17(9):1304– 1318, 2007.
- [7] J. Michaud, K.M. Simpson, R. Escher, K. Buchet-Poyau, T. Beissbarth, C. Carmichael, M.E. Ritchie, F. Schütz, P. Cannon, M. Liu, et al. Integrative analysis of RUNX 1 downstream pathways and target genes. *BMC Genomics*, 9(1):363, 2008.
- [8] G. Parmigiani, J. Lin, S. Boca, T. Sjöblom, KW Kinzler, VE Velculescu, and B. Vogelstein. Statistical methods for the analysis of cancer genome sequencing data. *Johns Hopkins University*, Dept. of Biostatistics Working Paper 126, 2007.
- [9] D.W. Parsons, S. Jones, X. Zhang, J.C.H. Lin, R.J. Leary, P. Angenendt, P. Mankoo, H. Carter, I. Siu, et al. An integrated genomic analysis of human glioblastoma multiforme. *Science*, 321(5897):1807, 2008.
- [10] D.W. Parsons, M. Li, X. Zhang, S. Jones, R.J. Leary, J.C.H. Lin, S.M. Boca, H. Carter, J. Samayoa, C. Bettegowda, et al. The genetic landscape of the childhood cancer medulloblastoma. *Science*, 331(6016):435, 2011.
- [11] EM Schaeffer, L. Marchionni, Z. Huang, B. Simons, A. Blackman, W. Yu, G. Parmigiani, and DM Berman. Androgen-induced programs for prostate epithelial growth and invasion arise in embryogenesis and are reactivated in cancer. *Oncogene*, 27(57):7180–7191, 2008.
- [12] T. Sjöblom, S. Jones, L.D. Wood, D.W. Parsons, J. Lin, T.D. Barber, D. Mandelker, R.J. Leary, J. Ptak, N. Silliman, et al. The consensus coding sequences of human breast and colorectal cancers. *Science*, 314(5797):268–274, 2006.
- [13] L.D. Wood, D.W. Parsons, S. Jones, J. Lin, T. Sjöblom, R.J. Leary, D. Shen, S.M. Boca, T. Barber, J. Ptak, et al. The genomic landscapes of human breast and colorectal cancers. *Science*, 318(5853):1108, 2007.

7 Acknowledgments

We would like to thank Michael A. Newton and Luigi Marchionni for generously sharing their code and Benjamin Langmead, Fernando Pineda, and Michael Ochs for some very helpful discussions.