# tugHall version 3.0: USER-GUIDE-tugHall

#### Requirements for tugHall simulation:

R version 3.6.0 or later

libraries: stringr, actuar, tidyr

Note that the program has two different procedures in general: the first is the simulation and the second is the analysis of the simulation results. Please, pay attention that the requirements for these procedures are **different**. This User-Guide pertains to the **simulation procedure** alone.

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# 1. Quick start guide

The simplest way to run tugHall:

- Save the /tugHall 2 2 CNA/ directory to the working folder;
- Run tugHall\_3.0.R.

The code has its initial input parameters and input files in the /Input/ folder. After the simulation the user can see results of the simulation (please, see **User-Guide-Analysis\_3** for details) in the dialogue box, which will save to the /Output/ and /Figures/ folders. Note that the analysis procedure requires additional libraries and a higher version of R - 3.6.0.

# 2. Structure of directories

#### **Documentation directory:**

**User-Guide-tugHall\_v\_3.0.XXX** - user guide for a simulation in the XXX = Rmd, html or pdf formats.

**User-Guide-Analysis\_v3.0.XXX** - user guide for the generation of an analysis and a report in the XXX = Rmd, html or pdf formats.

dir /tugHall\_3\_0\_CNA/ - the directory that contains the software tugHall version 3.0.

#### /tugHall\_3\_0\_CNA/ directory:

tugHall\_3.0.R - R script to run a simulation and to define the parameters.

dir /Code/ - the folder with a code and a function library.

dir /Input/ - the folder with the input files.

dir /Output/ - the folder with the output files.

dir /Figures/ - the folder with the plot figures.

#### /Code/ directory:

pic\_lic.jpg - the necessary file for the user guide.

tugHall\_3.0\_functions.R - the file that contains the functions for the simulation / core of program.

**read\_maps.R** - the file to read chromosomal locations got genes of interest from CCDS.current.txt file in the *Input/* folder

Analysis\_clones.R - the file to analyze the results of a simulation and to plot figures.

Functions\_clones.R - the file with the functions for the analysis of results.

#### /Input/ directory:

cloneinit.txt - file with a list of initial cells with/without destroyed genes.

gene\_cds2.txt - file with hallmark variables and weights.

CCDS.current.txt - file with information about chromosomal locations that was getting from CCDS database.

#### /Output/ directory:

cloneout.txt - the file with simulation output.

geneout.txt - the file with information about hallmark variables and the weights.

**log.txt** - the file with information about all parameters.

Weights.txt - the file with information about weights between hallmarks and genes.

point\_mutations.txt - the file contains information about point mutations in genome of clones.

**CNA\_mutations.txt** - the file contains information about copy number alterations in genome of clones.

gene\_map.txt - file with information about chromosomal locations for genes of interest only.

Order\_of\_malfunction.txt - see USER-GUIDE-Analysis.

VAF.txt - see USER-GUIDE-Analysis.

#### /Figures/ directory

In the /Figures/ directory there are figures in \*.jpg format, which appear after the analysis of the simulation results. See USER-GUIDE-Analysis\_3.

# 3. Inputs

## Input of hallmark variables and gene weights

The file tugHall/Input/gene\_hallmarks.txt defines the hallmark variables and weights:

Table 1. Input file for genes. Example of input file for hallmarks and weights in the file tugHall\_2\_2\_CNA/Input

/gene\_hallmarks.txt.

	<u> </u>		
Genes	Suppressor or Oncogene	Hallmark	Weights
APC	S	apoptosis	0.2616483
APC	S	growth	0.3285351
APC	S	invasion	0.3746081
KRAS	0	apoptosis	0.2099736
KRAS	0	growth	0.2881968
KRAS	0	immortalization	0.4735684
KRAS	0	angiogenesis	0.3525394
KRAS	0	invasion	0.0446472
TP53	S	apoptosis	0.2543523
TP53	S	growth	0.3076387
TP53	S	angiogenesis	0.4012288
TP53	S	immortalization	0.5264316
TP53	S	invasion	0.0645107
PIK3CA	0	invasion	0.3588945
PIK3CA	0	growth	0.2879753
PIK3CA	0	angiogenesis	0.3261495
PIK3CA	0	apoptosis	0.2938981

<sup>1.</sup> **Genes** - name of gene, e.g., TP53, KRAS. The names must be typed carefully. The program detects all the unique gene names.

- 2. Suppressor or oncogene. Distinction of oncogene/suppressor:
- o: oncogene
- o s: suppressor
- ?: unknown (will be randomly assigned)
- 3. Hallmark hallmark name, e.g., "apoptosis". Available names:
- apoptosis
- o immortalization
- growth
- o anti-growth
- angiogenesis
- invasion

Note that "growth" and "anti-growth" are related to the single hallmark "growth/anti-growth". Note that "invasion" is related to "invasion/metastasis" hallmark.

4. **Weights** - Hallmark weights for genes, e.g., 0.333 and 0.5. For each hallmark, the program checks the summation of all the weights. If it is not equal to 1, then the program normalizes it to reach unity. Note that, if the gene belongs to more than one hallmark type, it must be separated into separate lines.

After that, the program defines all the weights. **Unspecified weights** are set to 0. Program performs normalization so that the sum of all weights should be equal to 1 for each column. The **tugHall/Output** /**Weights.txt** file saves these final input weights for the simulation. Only the first 10 lines are presented here:

**Table 2. Weights for hallmarks.** Example of weights for hallmarks and genes from *tugHall/Output*/Weights.txt file. Unspecified values equal 0.

Genes	Apoptosis, $H_a$	Angiogenesis, $H_b$	$ \begin{array}{c} \textbf{Growth / Anti-} \\ \textbf{growth, } H_d \end{array} $	Immortalization, $H_i$	Invasion / Metastasis, $H_{im}$
APC	0.2565501	0.0000000	0.2709912	0.0000000	0.4445540
KRAS	0.2058822	0.3264502	0.2377183	0.4735684	0.0529836
TP53	0.2493962	0.3715365	0.2537549	0.5264316	0.0765560
PIK3CA	0.2881715	0.3020133	0.2375356	0.0000000	0.4259064

- 1. Genes name of genes.
- 2. **Apoptosis,**  $H_a$  weights of hallmark "Apoptosis".
- 3. **Angiogenesis,**  $H_b$  weights of hallmark "Angiogenesis".
- 4. **Growth / Anti-growth,**  $H_d$  weights of hallmark "Growth / Anti-growth".
- 5. **Immortalization,**  $H_i$  weights of hallmark "Immortalization".
- 6. **Invasion / Metastasis,**  $H_{im}$  weights of hallmark "Invasion / Metastasis".

## Input the probabilities

The input of the probabilities used in the model is possible in the code for parameter value settings, "tugHall\_3\_0.R":

Probability variable and value	Description
E0 <- 2E-4	Parameter $E_0$ related to environmental resource limitation
F0 <- 1E0	Parameter $F_0$ related angiogenesis
m <- 1E-6	Point mutation probability $m^\prime$
uo <- 0.5	Gene malfunction probability by point mutation for oncogene $u_{o}$
us <- 0.5	Gene malfunction probability by point mutation for suppressor $u_s$
s <- 10	Parameter in the sigmoid function $s$
k <- 0.1	Environmental death probability $k^\prime$
m_dup <- 0.01	CNA duplication probability $m_{dup}$
m_del <- 0.01	CNA deletion probability $m_{del}$
lambda_dup <- 7000	CNA duplication average length $\lambda_{dup}$ (of the geometrical distribution for the length)
lambda_del <- 5000	CNA deletion average length $\lambda_{del}$ (of the geometrica distribution for the length)
uo,dup <- 0.8	Gene malfunction probability by CNA duplication for oncogene $u_{o,dup}$
us,dup <- 0	Gene malfunction probability by CNA duplication for suppressor, $u_{s,dup}$ . Currently, 0 is assumed.
uo,del <- 0	Gene malfunction probability by CNA deletion for oncogene $u_{o,del}$ . Currently, 0 is assumed.
us,del <- 0.8	Gene malfunction probability by CNA deletion for suppressor, $u_{s,del}$ .

## Filename input

Also in the code "tugHall\_3\_3.R" user can define names of input and output files, and additional parameters of simulation:

Variables and file names	Description
genefile <- 'gene_hallmarks.txt'	File with information about gene-hallmarks weights
mapfile <- 'gene_map.txt'	File with information about genes' map
clonefile <- 'cloneinit.txt'	Initial Cells
geneoutfile <- 'geneout.txt'	Gene Out file with hallmarks
cloneoutfile <- 'cloneout.txt'	Output information of simulation
logoutfile <- 'log.txt'	Log file to save the input information of simulation
censore_n <- 30000	Max cell number where the program forcibly stops
censore t <- 200	Max time where the program forcibly stops

### Input of the initial clones

Pay attention, it does not work yet, and it will be working.

The initial states of cells are defined in "tugHall\_3\_0\_CNA/Input/cloneinit.txt" file:

Clone ID	List of malfunctioned genes	Number of cells
1	11 11	1000
2	"APC"	10
3	"APC, KRAS"	100
4	"KRAS"	1
5	"TP53, KRAS"	1
		100
1000	11 11	10

- 1. Clone ID ID of clone, e.g., 1, 324.
- 2. **List of malfunctioned genes** list of malfunctioned genes for each clone, e.g. "","KRAS, APC". The values are comma separated. The double quotes ("") without gene names indicate a clone without malfunctioned genes.
- 3. Number of cells number of cells in each clone, e.g., 1, 1000.

#### Input of the genes' maps

This new version of **tugHall** allows to calculate CNAs in the genome. The breakpoints of CNAs may fall on genic regions consisting of exons and introns. That's why it's needed to enter information about gene's map. In the **/Input/** directory you can find **CCDS.current.txt**, which was getting from <u>CCDS database</u> at the National Center for Biotechnology Information and has information about genes. At the beginning of simulation, the program reads this file and extracts genes' map, which is put into "**tugHall\_2\_clones/Input/gene\_map.txt**". For example, the map is shown as follow:

Chr	CCDS_ID	Gene	Start	End	Len
5	CCDS4107.1	APC	112754890	112755024	135
5	CCDS4107.1	APC	112766325	112766409	85
5	CCDS4107.1	APC	112767188	112767389	202
5	CCDS4107.1	APC	112775628	112775736	109
5	CCDS4107.1	APC	112780789	112780902	114
5	CCDS4107.1	APC	112792445	112792528	84
5	CCDS4107.1	APC	112801278	112801382	105
5	CCDS4107.1	APC	112815494	112815592	99
5	CCDS4107.1	APC	112818965	112819343	379
5	CCDS4107.1	APC	112821895	112821990	96

- 1. Chr Name of the chromosome, e.g., 1, 12, X, Y.
- 2. CCDS\_ID ID of the gene in the CCDS database.
- 3. **Gene** the name of the gene.
- 4. Start the start position of each exon of the gene.
- 5. **End** the final position of each exon of the gene.
- 6. **Len** the length of gene's location Len = End Start + 1

# 4. Outputs

The output data consists of several files after the simulation.

### "log.txt" file

The file "log.txt" contains information about probabilities and file names. These variables are explained in the "Inputs".

**Table 3. log.txt file.** Example of log.txt file.

Variable	Value
genefile	Input/gene_hallmarks.txt
clonefile	Input/cloneinit.txt
geneoutfile	Output/geneout.txt
cloneoutfile	Output/cloneout.txt
logoutfile	Output/log.txt
E	1e-04
F	10
m0	1e-06
uo	0.5
us	0.5
S	10
k	0.2
m_dup	8e-08
m_del	1e-09
lambda_dup	5000
lambda_del	7000
uo_dup	0.8
us_dup	0
uo_del	0
us_del	0.8
censore_n	1e+05
censore_t	100
d0	0.35

## "geneout.txt" file

The file "geneout.txt" contains input information about the weights that connect the hallmarks and genes, which are defined by the user. These variables also are explained in the "Inputs".

**Table 4. geneout.txt file.** Given below is an example of the geneout.txt file.

Gene_name	Hallmark_name	Weight	Suppressor_or_oncogene
APC	apoptosis	0.2565501	s

Gene_name	Hallmark_name	Weight	Suppressor_or_oncogene
KRAS	apoptosis	0.2058822	0
TP53	apoptosis	0.2493962	s
PIK3CA	apoptosis	0.2881715	0
KRAS	immortalization	0.4735684	0
TP53	immortalization	0.5264316	s
APC	growth anti-growth	0.2709912	S
KRAS	growth anti-growth	0.2377183	0
TP53	growth anti-growth	0.2537549	S
PIK3CA	growth anti-growth	0.2375356	0
KRAS	angiogenesis	0.3264502	0
TP53	angiogenesis	0.3715365	S
PIK3CA	angiogenesis	0.3020133	0
APC	invasion	0.4445540	S
KRAS	invasion	0.0529836	0
TP53	invasion	0.0765560	s
PIK3CA	invasion	0.4259064	0

#### "cloneout.txt" file

The file "cloneout.txt" contains the results of the simulation and includes the evolution data: all the output data for each clone at each time step (only the first 10 lines are presented):

**Table 5. Output data.** Example of output data for all clones. The names of columns are related to the description in the Tables 1,2 and *USER-GUIDE-Analysis\_3*'s figures. Columns are from 1 to 16.

Time	N_cells	AvgOrIndx	ID	ParentID	Birth_time	С	d	i	im	а	k	E	N	Nmax	M
0	-	avg	-	-	-	0	0.25	1	0	0.0066	0.2	1e-04	1000	10000	0
0	1000	1	1	0	0	0	0.25	1	0	0.0066	0.2	1e-04	1000	10000	0
1	-	avg	-	-	-	0.2496	0.2502	1	0.0004	0.0066	0.2	9.9924	1001	10030.	0
1	989	1	1	0	0	0.2496	0.2499	1	0	0.0066	0.2	1e-04	1001	10000	0
1	1	2	2	1	0	0.2496	0.2499	1	0	0.0066	0.2	1e-04	1001	10000	0
1	1	3	3	1	0	0.2496	0.2499	1	0	0.0066	0.2	1e-04	1001	10000	0
1	1	4	4	1	0	0.2496	0.5626	1	0.4259	0	0.2	2.4874	1001	40201.	0
1	1	5	5	1	0	0.2496	0.2499	1	0	0.0066	0.2	1e-04	1001	10000	0

Time	N_cells	AvgOrIndx	ID	ParentID	Birth_time	С	d	i	im	а	k	E	N	Nmax	М
1	1	6	6	1	0	0.2496	0.2499	1	0	0.0066	0.2	1e-04	1001	10000	0
1	1	7	7	1	0	0.2496	0.2499	1	0	0.0066	0.2	1e-04	1001	10000	0

- 1. **Time** the time step, e.g., 1, 50.
- 2. N\_cells the number of cells in this clone, e.g. 1000, 2.
- 3. **AvgOrIndx** "avg" or "index": "avg" is for a line with averaged values across different (index) lines at the same time step; "index" shows the cell's index at the current time step, e.g., avg, 4,7.
- 4. **ID** the unique ID of clone, e.g., 1, 50.
- 5. Parent\_ID the parent index, e.g., 0, 45.
- 6. Birth\_time the time step of the clone's birth, e.g., 0, 5.
- 7. **c** the counter of cell divisions for the clone.
- 8. d the probability of division for the cell, e.g., 0.1, 0.8.
- 9. i the probability of immortalization for the cell, e.g., 0.1, 0.8.
- 10. im the probability of invasion/metastasis for the cell, e.g., 0.1, 0.8.
- 11. a the probability of apoptosis for the cell, e.g., 0.1, 0.8.
- 12. k the probability of death due to the environment, e.g., 0.1, 0.8.
- 13. **E** the E coefficient for the function of the division probability, e.g., 10<sup>4</sup>, 10<sup>5</sup>.
- 14. **N** the number of primary tumor cells at this time step, e.g., 134, 5432.
- 15. Nmax the theoretically maximal number of primary tumor cells, e.g., 10000, 5000.
- 16. **M** the number of metastasis cells at this time step, e.g., 16, 15439.

#### Continuation of Table 5. Columns are from 17 to 28.

Time	AvgOrIndx	На	Him	Hi	Hd	Hb	type	mut_den	driver_genes	passenger_genes	PointMut_ID	CNA_ID
0	avg	0	0	0	0	0	0	0	-	-	-	-
0	1	0	0	0	0	0	0	0	0000	0000	0	0
1	avg	0.000287	0.000425	0	0.000237	0.000301	0	0.000249	-	-	-	-
1	1	0	0	0	0	0	0	0	0000	0000	0	0
1	2	0	0	0	0	0	0	0	0000	1000	1	0
1	3	0	0	0	0	0	0	0	0000	0001	3	0
1	4	0.288171	0.425906	0	0.237535	0.302013	0	0.25	0001	0000	0	1
1	5	0	0	0	0	0	0	0	0000	1000	0	2
1	6	0	0	0	0	0	0	0	0000	1000	0	3
1	7	0	0	0	0	0	0	0	0000	1000	0	4
1	8	0	0	0	0	0	0	0	0000	1000	0	5
1	9	0	0	0	0	0	0	0	0000	1000	0	6
1	10	0	0	0	0	0	0	0	0000	1000	0	7
1	11	0	0	0	0	0	0	0	0000	0010	0	8
1	12	0	0	0	0	0	0	0	0000	1000	5	0

Time	AvgOrIndx	На	Him	Hi	Hd	Hb	type	mut_den	driver_genes	passenger_genes	PointMut_ID	CNA_ID
1	13	0	0	0	0	0	0	0	0000	1000	7	0
2	avg	0.000518	0.000898	0	0.000548	0	0	0.000505	-	-	-	-
2	1	0	0	0	0	0	0	0	0000	0000	0	0

- 17. **Ha** the value of the hallmark "Apoptosis" for the cell, e.g., 0.1, 0.4444.
- 18. Him the value of the hallmark "Invasion / Metastasis" for the cell, e.g., 0.1, 0.4444.
- 19. Hi the value of the hallmark "Immortalization" for the cell, e.g., 0.1, 0.4444.
- 20. **Hd** the value of the hallmark "Growth / Anti-growth" for the cell, e.g., 0.1, 0.4444 .
- 21.  $\mathbf{Hb}$  the value of the hallmark "Angiogenesis" for the cell, e.g., 0.1, 0.4444 .
- 22. type the type of the cell: "0" is primary tumor cell, "1" is the metastatic cell, e.g., 0, 1.
- 23. mut\_den the density of mutations (tumor mutation burden) for the cell, e.g., 0, 0.32.
- 24. **driver\_genes** the binary numbers indicate the driver mutation at the gene related to order of genes in onco as well as order of the next columns with genes' names.
- 25. **passenger\_genes** the binary numbers indicate the passenger mutation at the gene related to order of genes in onco as well as order of the next columns with genes' names.
- 26. **PointMut\_ID** the index of data row for point mutation data frame saved at the end of simulation in the file **Point mutations.txt**, e.g., 23, 32.
- 27. CNA\_ID the index of data row for CNA data frame saved at the end of simulation in the file CNA.txt, e.g., 44, 21.

There are two columns (24th and 25th) with the indexes of point mutations and CNAs in Table 5. Each index corresponds to index in the related data frames for point mutations and for CNAs (Tables 6 and 7 respectively).

#### Continuation of Table 5. Columns are from 29 to end.

onco_ID	CDS_APC	CDS_KRAS	CDS_TP53	CDS_PIK3CA	Len_APC	Len_KRAS	Len_TP53	Len_PIK3CA	p0	prob_point_mut	prob_del	prob_du
-	-	-	-	-	-	-	-	-	-	-	-	_
1	8532	567	1182	3207	89236	35590	6986	35539	0.9733	0.4987	0.0061	0.4950
-	-	-	-	-	-	-	-	-	-	-	-	_
1	8532	567	1182	3207	89236	35590	6986	35539	0.9733	0.4987	0.0061	0.4950
2	8532	567	1182	3207	89236	35590	6986	35539	0.9733	0.4987	0.0061	0.4950
3	8532	567	1182	3207	89236	35590	6986	35539	0.9733	0.4987	0.0061	0.4950
4	8532	567	1182	3207	89236	35590	6986	35539	0.9733	0.4987	0.0061	0.4950
5	8994	567	1182	3207	89698	35590	6986	35539	0.9728	0.5064	0.0060	0.4874
6	8607	567	1182	3207	89311	35590	6986	35539	0.9732	0.5000	0.0061	0.4937
7	8947	567	1182	3207	89651	35590	6986	35539	0.9728	0.5057	0.0061	0.4881
8	13648	567	1182	3207	94352	35590	6986	35539	0.9679	0.5711	0.0052	0.4235
9	8553	567	1182	3207	89257	35590	6986	35539	0.9732	0.4991	0.0061	0.4947
10	8532	567	1182	3207	89236	35590	6986	35539	0.9733	0.4987	0.0061	0.4950
11	8532	567	1930	3207	89236	35590	9366	35539	0.9724	0.5087	0.0060	0.4852

onco_ID	CDS_APC	CDS_KRAS	CDS_TP53	CDS_PIK3CA	Len_APC	Len_KRAS	Len_TP53	Len_PIK3CA	p0	prob_point_mut	prob_del	prob_du
12	8532	567	1182	3207	89236	35590	6986	35539	0.9733	0.4987	0.0061	0.4950
13	8532	567	1182	3207	89236	35590	6986	35539	0.9733	0.4987	0.0061	0.4950
-	-	-	-	-	-	-	-	-	-	-	-	-
1	8532	567	1182	3207	89236	35590	6986	35539	0.9733	0.4987	0.0061	0.4950

<sup>28.</sup> **onco\_ID** - the index of the data related to onco object at simulation that has info about lengths of genes and genes' CDS for each chromosome.

29-32. **CDS\_(gene's name)**, for example **CDS\_APC** - the length of CDS for each gene in the order of names of genes for ONLY FIRST chromosome of a clone.

33-36. **Len\_(gene's name)**, for example **Len\_APC** - the length of gene in the order of names of genes for ONLY FIRST chromosome of a clone. The length of genes for second chromosome can be different.

- 37. p0 the probability that during a trial, a cell of the clone has NO mutation. Applied to all cells in the clone.
- 38. **prob\_point\_mut** the conditional probability that if cell will have a mutation it should be a **point mutation**.
- 39. prob\_del the conditional probability that if cell will have a mutation it should be a deletion.
- 40. prob\_dup the conditional probability that if cell will have a mutation it should be a duplication.

The information of the columns from 28 to 40 is related to *onco* object in simulation for a clone. Please, pay attention that probability of mutations depend on length of CDS and gene of all chromosome but the table has information only for first chromosome.

Table 6. Point mutation data frame which will be saved to the file Point mutations.txt at the end of simulation.

	PointMut_ID	Parental_1or2	Chr	Ref_pos	Phys_pos	Delta	Copy_number	Gene_name	MalfunctionedByPointMut	mut_order
179	179	2	17	7676150	[7676150]	[0]	1	TP53	TRUE	163
180	179	1	17	7676150	[NA]	[NA]	1	TP53	NA	163
181	181	2	5	112841874	[112841874, 112843388]	[0, 1514]	2	APC	FALSE	164
182	181	1	5	112841874	[NA]	[NA]	1	APC	NA	164
183	183	2	5	112842532	[112842532]	[0]	1	APC	TRUE	168
184	183	1	5	112842532	[NA]	[NA]	1	APC	NA	168
185	185	2	5	112840836	[112840836]	[0]	1	APC	FALSE	170
186	185	1	5	112840836	[NA]	[NA]	1	APC	NA	170
187	187	2	5	112838108	[112838108]	[0]	1	APC	FALSE	171
188	187	1	5	112838108	[NA]	[NA]	1	APC	NA	171
189	189	1	5	112838634	[112838634]	[0]	1	APC	FALSE	172
190	189	2	5	112838634	[NA]	[NA]	1	APC	NA	172

	PointMut_ID	Parental_1or2	Chr	Ref_pos	Phys_pos	Delta	Copy_number	Gene_name	MalfunctionedByPointMut	mut_order
191	191	2	5	112841590	[112841590]	[0]	1	APC	TRUE	181
192	191	1	5	112841590	[NA]	[NA]	1	APC	NA	181
193	193	2	5	112841756	[112841756]	[0]	1	APC	FALSE	182
194	193	1	5	112841756	[NA]	[NA]	1	APC	NA	182
195	195	2	5	112840487	[112840487]	[0]	1	APC	FALSE	183
196	195	1	5	112840487	[NA]	[NA]	1	APC	NA	183
197	197	1	3	179201398	[179201398]	[0]	1	PIK3CA	TRUE	184
198	197	2	3	179201398	[NA]	[NA]	1	PIK3CA	NA	184
199	199	1	5	112792458	[112792458]	[0]	1	APC	TRUE	185
200	199	2	5	112792458	[NA]	[NA]	1	APC	NA	185

- PointMut\_ID ID of point mutation, first ID is related to variant allele 'B' and same second ID to the original allele A.
- 2. Parental\_1or2 indicates either of the two parental chromosomes.
- 3. Chr name of a chromosome.
- 4. **Ref\_pos** the reference position of an allele. The reference position is on the coordinate system of the human reference genome.
- 5. Phys\_pos the physical position of an allele. The physical length of a (parental) chromosome is extended or shrunk by CNA duplications or deletions, respectively. When a duplication happens, the reference position is divided into two or more physical positions, which are represented by multiple elements in a vector. When a deletion happens and the allele is lost, the lost is represented by "-" on the coordinate system of physical positions.
- 6. **Delta** difference between the reference and physical positions.
- 7. Copy\_number the copy number of an allele.
- 8. **Gene\_name** the name of a gene.
- MalfunctionedByPointMut logical indicator of whether or not the gene is malfunctioned by the point
  mutation. 10. mut\_order indicator of mutation order in the simulation, it's used to detect order of
  mutations in the clone at each chromosome.

Table 7. CNA mutation data frame which will be saved to the file CNA.txt at the end of simulation.

	CNA_ID	Parental_1or2	dupOrdel	Chr	Ref_start	Ref_end	Gene_names	MalfunctionedByCNA	mut_order
200	200	2	dup	5	112842255	112844456	APC	FALSE	426
201	201	2	dup	5	112835091	112835164	APC	FALSE	428
202	202	1	dup	5	112839406	112844125	APC	FALSE	429
203	203	1	dup	5	112843606	112844125	APC	FALSE	430
204	204	1	dup	3	179219684	179225408	PIK3CA	TRUE	434
205	205	1	dup	5	112841043	112844125	APC	FALSE	435
206	206	1	dup	5	112819033	112821331	APC	FALSE	439

	CNA_ID	Parental_1or2	dupOrdel	Chr	Ref_start	Ref_end	Gene_names	MalfunctionedByCNA	mut_order
207	207	1	dup	5	112842671	112843715	APC	FALSE	440
208	208	1	dup	3	179225970	179228295	PIK3CA	TRUE	442
209	209	1	dup	5	112840539	112840575	APC	FALSE	446
210	210	1	dup	5	112838089	112838810	APC	FALSE	449
211	211	1	dup	5	112827128	112827316	APC	FALSE	450
212	212	2	dup	5	112842435	112844125	APC	FALSE	451
213	213	1	dup	5	112840306	112844125	APC	FALSE	452
214	214	1	dup	5	112841721	112844125	APC	FALSE	456
215	215	1	dup	5	112841873	112843314	APC	FALSE	457
216	216	1	dup	5	112843153	112844125	APC	FALSE	458
217	217	2	dup	5	112841809	112844125	APC	FALSE	459
218	218	1	dup	5	112839366	112844125	APC	FALSE	461
219	219	2	dup	5	112839636	112844125	APC	FALSE	462
220	220	2	dup	3	179224173	179225375	PIK3CA	TRUE	465

- 1. CNA\_ID ID of CNA.
- 2. Parental\_1or2 indicates either of the two parental chromosomes.
- 3. **DuplicationOrDeletion** indicator of duplication or deletion for CNA.
- 4. Chr name of a chromosome.
- 5. **Reference\_start** the reference position of the CNA start.
- 6. **Reference\_end** the reference position of the CNA end.
- 7. **Gene\_name** the name(s) of a gene(s).
- 8. MalfunctionedByCNA logical indicator of whether or not the gene(s) is malfunctioned by the CNA.
- 9. **mut\_order** indicator of mutation order in the simulation, it's used to detect order of mutations in the clone at each chromosome.

# 5. How to run

In order to make the simulation, please follow the procedure:

- 1. Copy /tugHall\_3\_0\_CNA/ directory into the working directory.
- 2. CD to the /tugHall\_3\_0\_CNA/ directory.
- 3. Run the tugHall\_3.0.R file, using the command line like

R --vanilla < tugHall\_3\_0.R

or using the line by line procedure in R Studio. In this case we have:

- load library(stringr) and source(file = "Code/tugHall\_2.2\_functions.R");
- o create the Output and Figures directories, if needed;
- o define the simulation parameters;
- o make the input file for initial cells, if needed;
- run the *model()* function to simulate;
- run source("Code/Amalysis\_clome\_R") in order to analyze the results and plot the figures in the dialogue box (see User-Guide-Analysis\_v3.0).
- 4. To obtain analysis reports of the simulation, please refer to User-Guide-Analysis\_v3.0.RMD. In User-Guide-Analysis\_v3.0.RMD, commands are embedded to include files under Output/ and Figure/. So, after analysis with tugHall, you can generate analysis reports automatically from User-Guide-Analysis\_v3.0.RMD. For more details, please refer to "Writing reproducible reports in R" on the github (https://nicercode.github.io/guides/reports/).

# 6. Differences with cell-based code and version 2.0

#### 6.1. Reason to develop clone-based code

- Clone-based code was designed to accelerate calculation and increase number of tumor cell. Advantage
  of clone-based algorithm is making trial for all cells at 1 clone with one application of trial() function. In
  cell-based algorithm trial() applies to each cell. But if number of cells equal number of clones, then speed
  up is 1. That's why clone-based code works faster for any cases.
- Another reason is a case, when we need to simulate huge number of cells like 10<sup>7</sup> or 10<sup>9</sup>, but mutation
  rate is very low. Cell-based algorithm takes a huge computational cost, and vice versa clone-based
  algorithm will work very fast, if mutated cells will appear slowly.

#### 6.2. Usage of trial() function

In trial() function program applies several trials like environmental death, apoptosis death, division
process, etc. We changed the trials with probability p (for some death process) for each cell in the clone
with for 1 trial with procedure:

$$N_{cells} = N_{cells} - Binom(p, N_{cells}),$$

where  $Binom(p, N_{cells})$  is random number from the binomial distribution with probability  $p, N_{cells}$  is a number of cells in a clone. Probability p is one of probability of death process, for example p=a' or p=k etc.

 $\circ$  For cell division with probability d' the new number of cells will be:

$$N_{cells} = N_{cells} + Binom(d', N_{cells})$$

• Check at the end of **trial()** function: if  $N_{cells} = 0$ , then the clone has died.

#### 6.3. Usage of mutation function

 In mutation function we have changed the mutation to birth of a new clone (one mutation is a birth of one clone):

 $N_{new\_clones} = Binom(m, N_{new\_cells}),$  $N_{new\_cells} = Binom(d', N_{cells}).$ 

> Passenger or Driver mutations do not matter for new clone's generation. Only during analysis, we will distinguish Passengers or Drivers clones.

#### 6.4. Average function

 $\circ$  The average values  $\overline{x}$  of probabilities or hallmarks are found by summation on the  $x_i$  with multiplication by cells number  $N_{cells,i}$  of this clone:

$$\bar{x} = \sum_i x_i \times w_i,$$

where  $w_i = \frac{N_{cells,i}}{N_{cells,tot}}$  is *i*th clone's occupancy in whole cell population  $N_{cells,tot} = \sum_i N_{cells,i}$ ,  $x_i$  is the value for *i*th clone, summation applies for all clones  $i = 1 \dots N_{clones}$ .

 For this purpose, we added the calculation of cells number (primary and metastasis) before average and hallmarks update.

#### 6.5. Difference with version 2.0

In the current version we use library *actuar* to make non-zero-binom calculation faster, and we use the approximation for big numbers of cells in **trial()** function, because **rbinom()** function in R has restriction for big numbers like  $n \times p > 10^{12}$ .

# 7. Differences with clone-based code and version 2.1

### 7.1. Reason to develop CNA-based code

New version of tugHall with copy number alteration (CNA) was designed for correct calculation of VAF influenced by CNA and tumor purity. It's expected that this design should improve comparison between observation VAF  $\in$  [0; 1] and calculated VAF. The previous versions of tugHall have VAF in the range [0; 0.5] because of the neglect of CNA and tumor purity.

#### 7.2. Calculation of point and CNA mutations

Probabilities of CNA mutations are calculated in the same way as point mutations:

•  $m_{point} = m_0 \times l_{CDS}$  - for point mutation, where  $l_{CDS}$  is the length of all exons of a gene and  $m_0$  is a constant per base pairs defined by users;

- $m_{0,dup}$  and  $m_{0,del}$ , or we collectively call  $m_{0,CNA}$ , indicate the first breakpoint event of a CNA and is a constant per base pairs defined by users.  $m_{CNA} = m_{0,CNA} \times l_{genes}$ , where  $l_{genes}$  is the total region size of all genes of interest which consists of exons as well as introns.
- a length of CNA is calculated using geometrical distribution:  $l_{CNA} = rgeom(1, 1/\lambda_{CNA}) + 1$ , where  $\lambda_{CNA}$  is average base-pair length defined by users.
- o probability of malfunctioning a gene  $u = u_{s,CNA}$  for suppressor and  $u = u_{o,CNA}$  for oncogene.

So, the algorithm of CNA is as follow:

```
if ( runif(1) < m_dup + m_del ) then 'Generate CNA':
    - define which event should occur - duplication or deletion using ratio m_dup/m_del like:
        event <- sample(c('dup', 'del'), 1, prob = c( m_dup, m_del )/sum(m_dup, m_del) )</pre>
```

- find randomly first position within the regions of genes of interest;
- find the length of CNA from geometrical distribution
- define with probability 0.5 is it the parental chromosome 1 or 2;
- define the list of genes in CNA;
- define with probabilty  $u = \{u_o \text{ or } u_s\}$  is it malfunction for each gene;
- check overlap of position for other mutations (and if it's necessesary change their positions).

The calculation of probabilities and hallmarks variables is not changed.

At the end of a simulation the VAF frequencies are calculated in accordance with formulation:

$$VAF^{i} = \frac{(1-\rho)n_{B,N}^{i} + \sum_{s=1}^{\# sp} \tau_{s}n_{B,S}^{i}}{(1-\rho)(n_{A,N}^{i} + n_{B,N}^{i}) + \sum_{s=1}^{\# sp} \tau_{s}(n_{A,S}^{i} + n_{B,S}^{i})},$$

where:

i is position (site) index,

s is subpopulation index,

 $\tau$  is subpopulation fraction,

 $\rho$  is tumor purity,

n is copy number,

A denotes an original allele A, B - variant B, and N - normal.

In usual application and in program we used for normal cells  $n_{A,N}^i=2$  and  $n_{B,N}^i=0$ , so VAF is calculated as follow:

$$VAF^{i} = \frac{\sum_{s=1}^{\# sp} \tau_{s} n_{B,S}^{i}}{2(1-\rho) + \sum_{s=1}^{\# sp} \tau_{s} (n_{A,S}^{i} + n_{B,S}^{i})}$$