## Using the tronco package

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July 21, 2015

**Abstract.** Genotype-level *cancer progression models* describe the temporal ordering in which genomic alterations such as somatic mutations and copy number alterations tend to fixate and accumulate during cancer formation and progression. These graphical models can describe trends of *natural selection* across a population of independent tumour samples (cross-sectional data), or reconstruct the clonal evolution in a single patient's tumour (multi-region or single-cell data). In terms of application, such models can be used to better elucidate genotype-phenotype relation, predict cancer hallmarks and outcome of personalised treatment as well as suggest novel targets for therapy design.

tronco (translational oncology) is a R package which collects algorithms to infer progression models from Bernoulli 0/1 profiles of genomic alterations across a tumor sample. Such profiles are usually visualised as a binary input matrix where each row represents a patient's sample (e.g., the result of a sequenced tumor biopsy), and each column an event relevant to the progression (a certain type of somatic mutation, a focal or higher-level chromosomal copy number alteration etc.); a 0/1 value models the absence/presence of that alteration in the sample. In this version of TRONCO such profiles can be readily imported by boolean matrices and MAF/GISTIC files. The package provides various functions to editing, visualise and subset such data, as well as functions to query the Cbio portal for cancer genomics. This version of TRONCO comes with the parallel implementations the CAPRESE [PLoS ONE 9(12): e115570] and CAPRI [Bioinformatics, doi:10.1093/bioinformatics/btv296] algorithms to infer possible progression models arranged as trees, or general direct acyclic graphs. Bootstrap functions to assess the parametric, non-prametric and statistical confidence of every inferred model are also provided. The package comes with some data available as well, which include the dataset of Atypical Chronic Myeloid Leukemia samples provided by Piazza et al., Nat. Genet., 45 (2013), and examples.

Requirements: You must have rgraphviz installed to use the package, see Bioconductor.org.

### **Event selection**

First, load tronco in your R console and the example "dataset".

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```
> library(TRONCO)
```

- > data(aCML)
- > hide.progress.bar <<- TRUE

### We use show function to get a short summary of the aCML dataset

```
> show(data)
Description: CAPRI - Bionformatics aCML data.
Dataset: n=64, m=31, |G|=23.
Events (types): Ins/Del, Missense point, Nonsense Ins/Del, Nonsense point.
Colors (plot): darkgoldenrod1, forestgreen, cornflowerblue, coral.
Events (10 shown):
         gene 4 : Ins/Del TET2
         gene 5 : Ins/Del EZH2
         gene 6 : Ins/Del CBL
         gene 7 : Ins/Del ASXL1
         gene 29 : Missense point SETBP1
         gene 30 : Missense point NRAS
         gene 31 : Missense point KRAS
         gene 32 : Missense point TET2
         gene 33 : Missense point EZH2
         gene 34 : Missense point CBL
Genotypes (10 shown):
          gene 4 gene 5 gene 6 gene 7 gene 29 gene 30 gene 31 gene 32 gene 33 gene 34
patient 1
               0
                      0
                                     0
                                                             0
                             0
                                             1
                                                     0
                                                                      0
                                                                              0
patient 2
               0
                      0
                             0
                                     0
                                             1
                                                     0
                                                             0
                                                                      0
                                                                              0
patient 3
               0
                      0
                             0
                                     0
                                             1
                                                     1
                                                             0
                                                                      0
                                                                              0
```

0

0

0

1

1

1

0

0

0

0

0

0

0

0

0

0

0

0

0

1

0

1

0

0

### These are all the events it contains

0

0

0

0

0

0

0

0

> as.events(data)

patient 4

patient 5

patient 6

```
type
                             event
gene 4
         "Ins/Del"
                             "TET2"
         "Ins/Del"
                             "EZH2"
gene 5
         "Ins/Del"
                             "CBL"
gene 6
gene 7
         "Ins/Del"
                             "ASXL1"
gene 29
         "Missense point"
                             "SETBP1"
         "Missense point"
                             "NRAS"
gene 30
gene 31
         "Missense point"
                             "KRAS"
         "Missense point"
                             "TET2"
gene 32
         "Missense point"
                             "EZH2"
gene 33
         "Missense point"
                             "CBL"
gene 34
gene 36
         "Missense point"
                             "IDH2"
gene 39
         "Missense point"
                             "SUZ12"
                             "SF3B1"
gene 40
         "Missense point"
gene 44
         "Missense point"
                             "JARID2"
gene 47
         "Missense point"
                             "EED"
gene 48
         "Missense point"
                             "DNMT3A"
         "Missense point"
                             "CEBPA"
gene 49
         "Missense point"
                             "EPHB3"
gene 50
gene 51 "Missense point"
                             "ETNK1"
```

```
"GATA2"
gene 52 "Missense point"
gene 53 "Missense point"
                           "IRAK4"
gene 54 "Missense point"
                           "MTA2"
gene 55 "Missense point"
                           "CSF3R"
gene 56 "Missense point"
                           "KIT"
gene 66 "Nonsense Ins/Del" "WT1"
gene 69 "Nonsense Ins/Del" "RUNX1"
gene 77
        "Nonsense Ins/Del" "CEBPA"
gene 88 "Nonsense point"
                           "TET2"
gene 89 "Nonsense point"
                           "EZH2"
gene 91 "Nonsense point"
                           "ASXL1"
gene 111 "Nonsense point"
                           "CSF3R"
```

### Which account for alterations in the following genes

> as.genes(data)

```
[1] "TET2"
              "EZH2"
                       "CBL"
                                "ASXL1"
                                         "SETBP1" "NRAS"
                                                            "KRAS"
                                                                     "IDH2"
                                                                              "SUZ12"
[10] "SF3B1"
              "JARID2" "EED"
                                "DNMT3A" "CEBPA"
                                                  "EPHB3"
                                                            "ETNK1"
                                                                     "GATA2"
                                                                              "IRAK4"
[19] "MTA2"
                                         "RUNX1"
              "CSF3R" "KIT"
                                "WT1"
```

### These are SETBP1 alterations across input samples

> as.gene(data, genes='SETBP1')

		Missense	point	SETBP1
patient	1		_	1
patient	2			1
patient	3			1
patient	4			1
patient	5			1
patient	6			1
patient	7			1
patient	8			1
patient	9			1
patient	10			1
patient	11			1
patient	12			1
patient	13			1
patient	14			1
patient	15			0
patient	16			0
patient	17			0
patient	18			0
patient	19			0
patient	20			0
patient	21			0
patient	22			0
patient	23			0
patient	24			0
patient	25			0
${\tt patient}$	26			0
${\tt patient}$	27			0
${\tt patient}$	28			0
${\tt patient}$	29			0
${\tt patient}$	30			0

patient	31	0
patient	32	0
${\tt patient}$	33	0
${\tt patient}$	34	0
${\tt patient}$	35	0
${\tt patient}$	36	0
${\tt patient}$	37	0
${\tt patient}$	38	0
${\tt patient}$	39	0
${\tt patient}$	40	0
${\tt patient}$	41	0
${\tt patient}$	42	0
${\tt patient}$	43	0
${\tt patient}$	44	0
${\tt patient}$	45	0
${\tt patient}$	46	0
${\tt patient}$	47	0
${\tt patient}$	48	0
${\tt patient}$	49	0
${\tt patient}$	50	0
${\tt patient}$	51	0
patient	52	0
patient	53	0
${\tt patient}$	54	0
patient	55	0
patient	56	0
patient	57	0
patient	58	0
${\tt patient}$	59	0
${\tt patient}$	60	0
patient	61	0
${\tt patient}$	62	0
${\tt patient}$	63	0
${\tt patient}$	64	0

These are the genes for which we found a literature supporting the patterns that we include below. References are in the main CAPRI paper.

```
> gene.hypotheses = c('KRAS', 'NRAS', 'IDH1', 'IDH2', 'TET2', 'SF3B1', 'ASXL1')
```

Regardless the distinct types of mutations that we included, we want to select only genes altered in 5% of the cases. Thus we first transform data in "Alteration" (collapsing all event types for the same gene), and then we use select only those events

```
> alterations = events.selection(as.alterations(data), filter.freq = .05)

*** Aggregating events of type(s) {Ins/Del, Missense point, Nonsense Ins/Del, Nonsense point}
in a unique event with label "Alteration".
Dropping event types Ins/Del, Missense point, Nonsense Ins/Del, Nonsense point for 23 genes.

*** Binding events for 2 datasets.

*** Events selection: #events=23, #types=1 Filters freq|in|out = {TRUE, FALSE, FALSE}
Minimum event frequency: 0.05 (3 alterations out of 64 samples).
Selected 7 events.
```

Selected 7 events, returning.

We visualize the selected genes. This plot has no title since name annotation is not copied by events.selection

```
> dummy = oncoprint(alterations)
```

\*\*\* Oncoprint for ""

with attributes: stage=FALSE, hits=TRUE

Sorting samples ordering to enhance exclusivity patterns.

Setting automatic row font (exponential scaling): 13

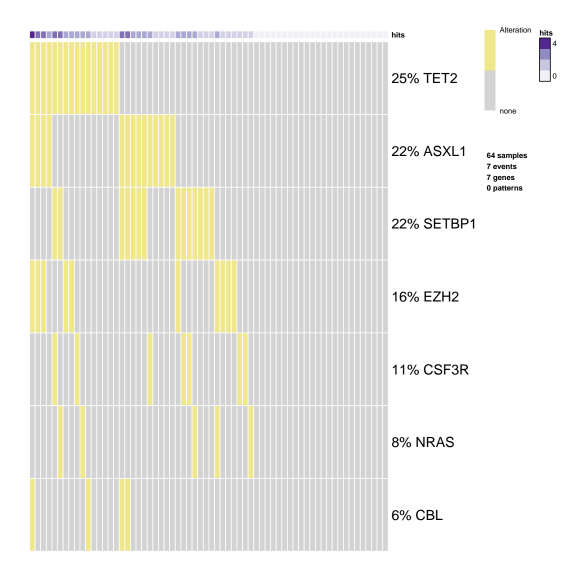


Figure 1: Oncoprint output

### **Adding Hypotheses**

Then to reconstruct the aCML model we select from data which have been selected in alteration - via as.genes(alterations) or that are part of the prior in gene.hypotheses. We use filter.in.names to force selection of all the events involving those genes from data

> hypo = events.selection(data, filter.in.names=c(as.genes(alterations), gene.hypotheses))

\*\*\* Events selection: #events=31, #types=4 Filters freq|in|out = {FALSE, TRUE, FALSE} [filter.in] Genes hold: TET2, EZH2, CBL, ASXL1, SETBP1 ... [10/14 found]. Selected 17 events, returning.

> hypo = annotate.description(hypo, 'CAPRI - Bionformatics aCML data (selected events)')

# We show selected data and we annotate genes in <code>gene.hypotheses</code> to identify them. Samples names are also shown

> dummy = oncoprint(hypo, gene.annot = list(priors= gene.hypotheses), sample.id = T)
\*\*\* Oncoprint for "CAPRI - Bionformatics aCML data (selected events)"
with attributes: stage=FALSE, hits=TRUE
Sorting samples ordering to enhance exclusivity patterns.
Annotating genes with RColorBrewer color palette Set1 .
Setting automatic row font (exponential scaling): 10.7
Setting automatic samples font half of row font: 5.3

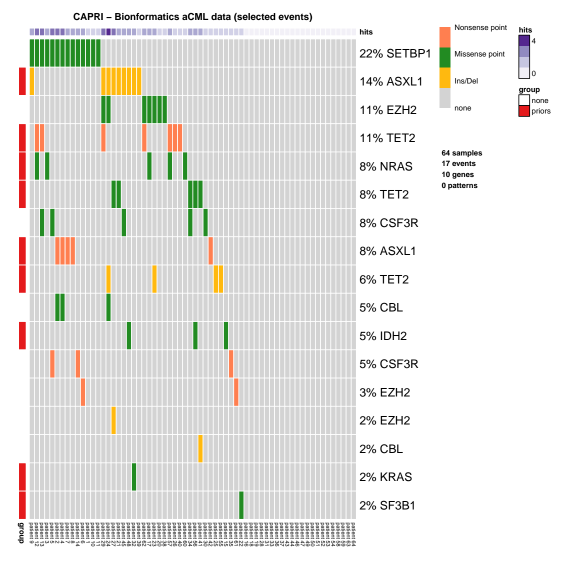


Figure 2: **Oncoprint output** 

We now add the hypotheses that are described in CAPRI's manuscript

Add hypotheses of hard exclusivity (XOR) for NRAS/KRAS events (Mutation). The hypothesis is tested against all other dataset events

```
> hypo = hypothesis.add(hypo, 'NRAS xor KRAS', XOR('NRAS', 'KRAS'))
```

Here we try to include also a soft exclusivity (OR) pattern but, since its "signature" is the same of the hard one, it will not be included. The code below is commented because it gives errors.

```
> hypo = hypothesis.add(hypo, 'NRAS or KRAS', OR('NRAS', 'KRAS'))
```

For the sake to better highlight the perfect (hard) exclusivity between NRAS/KRAS mutations one can visualize their alterations

```
> dummy = oncoprint(events.selection(hypo, filter.in.names = c('KRAS', 'NRAS')))
*** Events selection: #events=18, #types=4 Filters freq|in|out = {FALSE, TRUE, FALSE}
[filter.in] Genes hold: KRAS, NRAS ... [2/2 found].
Selected 2 events, returning.
*** Oncoprint for ""
with attributes: stage=FALSE, hits=TRUE
Sorting samples ordering to enhance exclusivity patterns.
Setting automatic row font (exponential scaling): 14.4
```

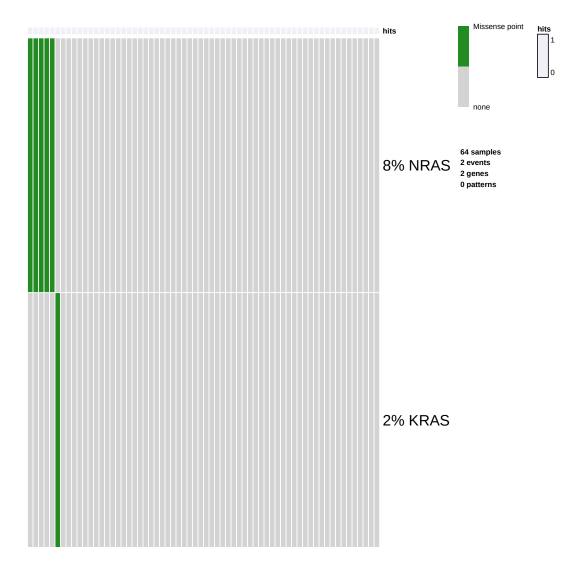


Figure 3: Oncoprint output

This is as above, but includes other events. Again, we can include only the hard exclusivity pattern

```
> hypo = hypothesis.add(hypo, 'SF3B1 xor ASXL1', XOR('SF3B1', OR('ASXL1')), '*')
> hypo = hypothesis.add(hypo, 'SF3B1 or ASXL1', OR('SF3B1', OR('ASXL1')), '*')
```

We now do the same for TET2 and IDH2. In this case 3 events for TET2 are present, which are "Ins/Del", "Missense point" and "Nonsense point". For this reason, since we are not specifying a subset of such events all TET2 alterations are used. Since these show a perfect hard exclusivity trend these will be included in XOR.

```
> hypo = hypothesis.add(hypo, 'TET2 or IDH2', OR('TET2', 'IDH2'), '*')
> dummy = oncoprint(events.selection(hypo, filter.in.names = c('TET2', 'IDH2')))
*** Events selection: #events=20, #types=4 Filters freq|in|out = {FALSE, TRUE, FALSE}
[filter.in] Genes hold: TET2, IDH2 ... [2/2 found].
Selected 4 events, returning.
*** Oncoprint for ""
with attributes: stage=FALSE, hits=TRUE
Sorting samples ordering to enhance exclusivity patterns.
Setting automatic row font (exponential scaling): 13.8
```

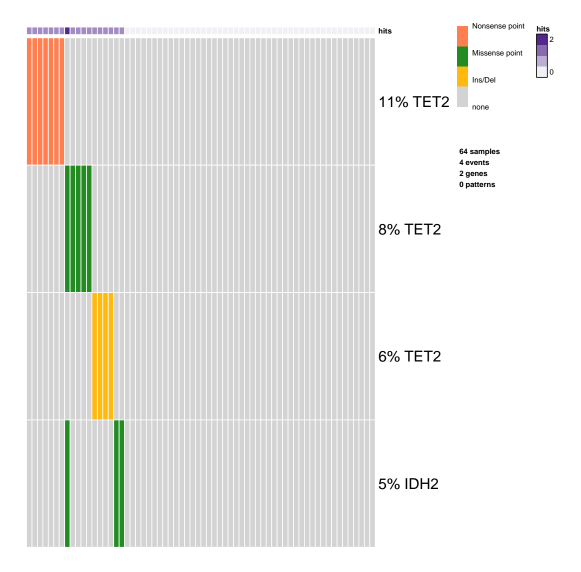


Figure 4: Oncoprint output

For every gene that has more than one event associated we also add a soft exclusivity pattern for its events

> hypo = hypothesis.add.homologous(hypo)

\*\*\* Adding hypotheses for Homologous Patterns

Genes: TET2, EZH2, CBL, ASXL1, CSF3R

Function: OR

Cause: \*
Effect: \*

Hypothesis created for all possible gene patterns.

#### The dataset input to CAPRI is shown

```
> dummy = oncoprint(hypo, gene.annot = list(priors= gene.hypotheses), sample.id = T)
*** Oncoprint for "CAPRI - Bionformatics aCML data (selected events)"
with attributes: stage=FALSE, hits=TRUE
Sorting samples ordering to enhance exclusivity patterns.
Annotating genes with RColorBrewer color palette Set1 .
Setting automatic row font (exponential scaling): 9.1
Setting automatic samples font half of row font: 4.5
```

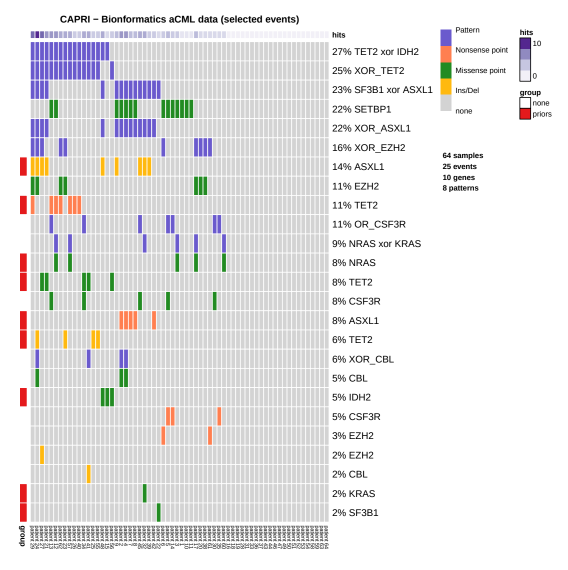


Figure 5: Oncoprint output

### Model reconstruction

We execute CAPRI with its default parameter: we use both AIC/BIC regularizators, Hill-climbing exhaustive bootstrap (100 replicates for Wilcoxon testing), p-value 0.05 and we set seed

```
> model = tronco.capri(hypo, boot.seed = 12345, regularization='bic', nboot=6)
*** Checking input events.
*** Inferring a progression model with the following settings.
        Dataset size: n = 64, m = 25.
        Algorithm: CAPRI with "bic" regularization and "hc" likelihood-fit strategy.
        Random seed: 12345.
        Bootstrap iterations (Wilcoxon): 6.
                exhaustive bootstrap: TRUE.
                p-value: 0.05.
               minimum bootstrapped scores: 3.
*** Bootstraping selective advantage scores (prima facie).
        Evaluating "temporal priority" (Wilcoxon, p-value 0.05)
        Evaluating "probability raising" (Wilcoxon, p-value 0.05)
*** Loop detection found loops to break.
        Removed 32 edges out of 64 (50%)
*** Performing likelihood-fit with regularization bic.
The reconstruction has been successfully completed in 00h:00m:02s
```

We can plot the reconstructed model. We set some parameters to get a fancy plot; confidence is shown as temporal priority and probability raising (selective advantage scores) and hypergeometric testing (goodness of input data).

```
> tronco.plot(model,
   fontsize = 13,
   scale.nodes = .6,
   confidence = c('tp', 'pr', 'hg'),
+ height.logic = 0.25,
  legend.cex = .5,
   pathways = list(priors= gene.hypotheses))
*** Expanding hypotheses syntax as graph nodes:
*** Rendering graphics
Nodes with no incoming/outgoing edges will not be displayed.
Annotating nodes with pathway information.
Annotating pathways with RColorBrewer color palette Set1 .
Set automatic fontsize for edge labels: 6.5
Adding confidence information: tp, pr, hg
RGraphviz object prepared.
Plotting graph and adding legends.
```

### **CAPRI – Bionformatics aCML data (selected events)**

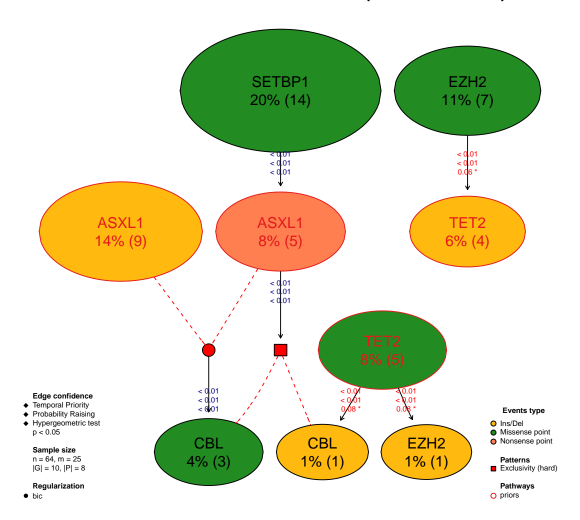


Figure 6: aCML Reconstructed model Pre bootstrap.

### **Bootstrapping data**

## + legend.cex = .5)

\*\*\* Expanding hypotheses syntax as graph nodes:

\*\*\* Rendering graphics

Nodes with no incoming/outgoing edges will not be displayed.

Set automatic fontsize for edge labels: 6.5

Adding confidence information: npb

RGraphviz object prepared.

Plotting graph and adding legends.

## **CAPRI – Bionformatics aCML data (selected events)**

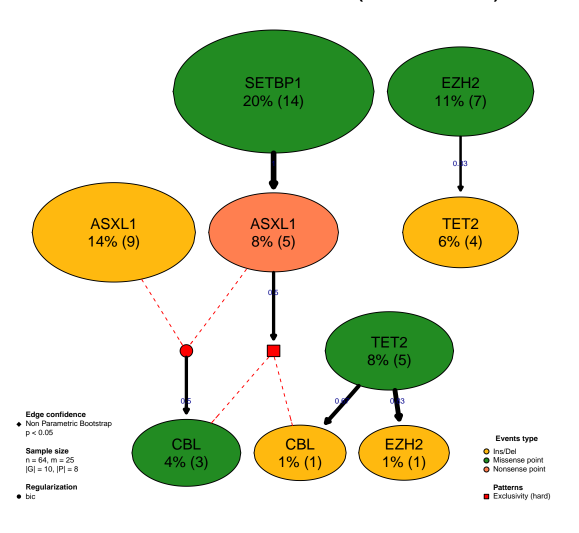


Figure 7: aCML Reconstructed model After bootstrap.