# Assessing Tumor Microsatellite Instability from Tumor Exome Somatic Mutations—The MSIseq Package

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The MSIseq package provides a mechanism for detecting microsatellite instability (MSI) in somatic mutation data from whole exome sequencing. The package provides both a classifier (detector), as well as a means to train new classifiers, which may be necessary depending on changes in variant-calling algorithms.

This package contains two main functions. The function MSIseq.train() generates a detector for MSI status from training data that consists of somatic mutation information and MSI status. The function MSIseq.classify() uses the generated detector to classify the MSI status of new tumors. The package also provides a helper function, Compute.input.variables(), to generate the input needed by these two functions.

# 1 Input data

As input, the MSIseq package requires somatic mutation information (i.e. from The Cancer Genome Atlas (TCGA) website, https://tcga-data.nci.nih.gov/tcga/) in the format of a "mutation annotation file" (https://wiki.nci.nih.gov/display/tcga/File+Format+Specifications). For example, NGStrain-data and NGStestdata are in this format.

- > library('MSIseq')
- > data(NGStraindata)
- > data(NGStestdata)
- > head(NGStraindata)

	Chrom	Start_Position	End_Position	Variant_Type	Tumor_Sample_Barcode
1	chr19	58862932	58862932	SNP	TCGA-D1-A15Z
3	chr10	52575855	52575856	INS	TCGA-BG-AOMO
4	chr10	52575855	52575856	INS	TCGA-BG-AOM3
5	chr10	52575855	52575856	INS	TCGA-BG-AOM9
6	chr12	9229467	9229467	SNP	TCGA-D1-A16B
7	chr12	9229527	9229527	SNP	TCGA-BG-AOM4

Usually the TCGA mutation annotation file contains 37 columns. The NGStraindata only contains the 5 columns that are necessary for this package and the Compute.input.variables() function. Any other columns are ignored. The names of the 5 columns must be exactly as shown.

In the 5 columns, Chrom indicates the chromosome identifier. Start\_Position and End\_Position are the start and end positions of the mutation in the chromosome. Variant\_Type indicates the type of variant, for which the legal values are "SNP", "INS" and "DEL". Other values will cause an error. Tumor\_Sample\_Barcode is the sample ID.

To obtain such a somatic mutation information table for your own data, you will need to create it from your sequence alignments and suitable annotation.

Another information that MSIseq package needs is the sequence length, which is the total length of the genomic regions from a DNA sample captured by sequencing techniques. For example, NGStrainseqLen and NGStestseqLen contain the information.

- > data(NGStrainseqLen)
- > data(NGStestseqLen)
- > head(NGStrainseqLen)

1 chr10

Tumor\_Sample\_Barcode Sequence\_Length

1	1CGA-CM-6677	44
2	TCGA-CA-6717	44
3	TCGA-AZ-4315	44
4	TCGA-D5-6531	44
5	TCGA-CM-6162	44
6	TCGA-AA-3663	44

The sequence length table contains 2 columns, Tumor\_Sample\_Barcode and Sequence\_Length, which indicate the sample IDs and their corresponding sequence lengths in megabases.

The genomic coordinates of simple sequence repeats in the reference genome are also required by MSIseq. The sequence repeats table contains 3 columns. Chrom indicates the chromosome identifier. Start\_Position and End\_Position are the start and end positions.

For example, we can get the simple sequence repeats in human genome (version Hg19) from the following link:

```
> url <-
+ "http://steverozen.net/data/Hg19repeats.rda"
> file <- basename(url)
> download.file(url, file)
> load("Hg19repeats.rda")
> head(Hg19repeats)

Chrom Start_Position End_Position
```

60213

60217

2	chr10	60287	60291
3	chr10	60379	60383
4	chr10	60518	60523
5	chr10	60741	60745
6	chr10	60898	60902

Hg19repeats is a static dataset which can be used for sequencing data generated with the same reference genome. You can also get it from http://hgdownload.cse.ucsc.edu/goldenpath/hg19/database/ for other reference genome.

Hg19repeats is a bed format data containing all the simple repeat regions (mono-, di-, tri-, tetra-nucleotide repeats) in the human genome version hg19. In this data, di-nucleotide, tri-nucleotide and tetra-nucleotide repeats are from the table in UCSC Genome Bioinformatics Site: ftp://hgdownload.cse.ucsc. edu/goldenPath/hg19/database/simpleRepeat.txt.gz. Mono-nucleotide repeats with a length  $\geq 5$  are generated with the following two functions, find.mono.repeats and find.mono.repeats against the human hg19 genome.

Here is an example of how one can generate the mono-nucleotide repeats in one chromosome:

```
> ## download the chromosome 20 sequence from UCSC
> url2 <-
+ "ftp://hgdownload.cse.ucsc.edu/goldenPath/hg19/chromosomes/chr20.fa.gz"
> file <- basename(url2)
> download.file(url2, file)
> gunzip(file)
> file <- 'chr20.fa'
> ## generate mono-nucleotide repeats regions from chromosome 20
> data.chr20 = read.fasta(file)
> mono.repeats.chr20 = find.mono.repeats(data.chr20)
> names(mono.repeats.chr20)<-c('Chrom', 'Start_Position', 'End_Position')</pre>
```

Users can easily apply these functions to their own fastq file and generate their own repeats file.

The MSI status information is required by MSIseq.train() function specifically. If your want to train a classifier with your own data, you need to have a classification table showing the clinical test result of MSI status of your samples.

- > data(NGStrainclass)
- > head(NGStrainclass)

```
Tumor_Sample_Barcode MSI_status
           TCGA-A6-5661
1
                              MSI-H
2
           TCGA-A6-5665
                              MSI-H
3
           TCGA-A6-6653
                              MSI-H
4
           TCGA-A6-6781
                              MSI-H
           TCGA-AA-3492
5
                              MSI-H
10
           TCGA-AA-3663
                              MSI-H
```

In the classification table, Tumor\_Sample\_Barcode represents the sample ID. MSI\_status is a factor with two levels, "MSI-H" and "Non-MSI-H". Other values will cause an error.

The cancer type information is optional. If you would like to use cancer type as a candidate input for your classifier, you need to have a cancer type table.

```
> data(NGStraintype)
> data(NGStesttype)
> head(NGStraintype)
  Tumor_Sample_Barcode cancer_type
1
          TCGA-D1-A15Z endometrial
2
          TCGA-D1-A17D endometrial
3
          TCGA-BG-AOMQ endometrial
4
          TCGA-BS-A0U9 endometrial
5
          TCGA-D1-A16D endometrial
6
          TCGA-BG-AOM7 endometrial
```

In the cancer type table, Tumor\_Sample\_Barcode represents the sample ID. cancer\_type is a factor which gives the corresponding cancer types.

## 2 Functions

First, the helper function Compute.input.variables() is used to generate the input needed by the two main functions.

```
> train.mutationNum<-Compute.input.variables(NGStraindata,
+ repeats = Hg19repeats, seq.len = NGStrainseqLen)</pre>
```

This function takes four arguments: Compute.input.variables(data, repeats, uniform.seq.len = 38, seq.len = NULL). The formats for data, repeats, and seq.len are explained in the Input data section. And the default seq.len argument is 38. This argument is used when sequences for all samples have the same length.

This function computes and extracts mutation count information from the argument data. The variable sequence length is used as a denominator to generate mutation count per megabase. The mutation can be either a single nucleotide substitution (SNS) or a short insertion/deletion (indel).

The returned value is a data frame containing the following 9 variables:

- T.sns: total count of SNSs/Mb
- $\bullet$  S.sns: count of SNSs in simple sequence repeats/Mb
- T.ind: total count of indels/Mb
- S.ind: count of indels in simple sequence repeats/Mb

- T: total mutation count/Mb
- S: mutation count in simple sequence repeats/Mb

• Ratio.sns: S.sns/T.sns

• Ratio.ind: S.ind/T.ind

• Ratio: S/T

Now let's look at the two main functions.

- > sampleclassifier<-MSIseq.train(mutationNum = train.mutationNum,
- + classification=NGStrainclass, cancerType = NGStraintype)

#### 5 fold cross validation result: 98.61496

The function MSIseq.train() takes three arguments: MSIseq.train(mutationNum, classification, cancerType = NULL). The format of mutationNum should be the same as the returned value of the helper function Compute.input.variables(). The format for classification and cancerType are explained in the Input data section. Again, the cancerType argument is optional. It depends on whether you want to train your classifier with cancer type information.

This function uses the 'RWeka' package to build and evaluate a J48 decision tree with the 9 variables (or 10 variables including 'cancer type'). The function will also give a five-fold cross validation result for the classification accuracy of the model.

The return value for MSIseq.train() is a Weka\_classifier object, a J48 decision tree classifier.

## > sampleclassifier

```
J48 pruned tree
```

 $S.ind \le 0.394737: Non-MSI-H (295.0/3.0)$ 

S.ind > 0.394737: MSI-H (66.0)

Number of Leaves : 2

Size of the tree: 3

The two output classes of the decision tree classifier are MSI-H and Non-MSI-H. 3 variables (S.ind, T.sns, S) are chosen to build the decision tree. When training with other data, you will get a different decision tree.

In this sample classifier, the desicion tree is based on a single variable, S.ind. If S.ind > 0.395, the tumor is classified as MSI-H. Otherwise, the classification is non-MSI-H. This classifier is also provided by MSIseq named as NGSclassifier. And it is offered as the default classifier for the second function, MSIseq.classify().

The function  ${\tt MSIseq.classify()}$  classifies tumors with unknown MSI status.

This function takes three arguments: MSIseq.classify(mutationNum, classifier, cancerType = NULL). The format of mutationNum should be the same as the returned value of the helper function Compute.input.variables(). The default classifier is a built-in classifier, NGSclassifier. You can also use your own classifier, which should be a returned value from the function MSIseq.train(). Remember if the input classifier is trained with cancerType argument, you should also give cancer type information in this function. And the format of cancerType should be the same as mentioned before.

## > head(result)

```
Tumor_Sample_Barcode MSI_status
1
          TCGA-A5-AOGE Non-MSI-H
2
          TCGA-D1-A176
                             MSI-H
3
          TCGA-BG-AOW1
                        Non-MSI-H
4
          TCGA-EO-A1Y5
                        Non-MSI-H
5
          TCGA-D1-A17U
                             MSI-H
6
          TCGA-AX-A064
                             MSI-H
```

The return value for MSIseq.classify() is a data frame with two columns, Tumor\_Sample\_Barcode and the corresponding classified MSI\_status.