

# Package ‘EthSEQ’

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**Type** Package

**Title** Ethnicity Annotation from Whole-Exome and Targeted Sequencing Data

**Version** 3.0.2

**Description** Reliable and rapid ethnicity annotation from whole exome and targeted sequencing data.

**License** GPL-3

**Depends** R (>= 2.15)

**Imports** graphics, utils, parallel, grDevices, MASS (>= 7.3-47), geometry (>= 0.3-6), data.table (>= 1.10.0), SNPRelate (>= 1.8.0), gdsfmt (>= 1.10.1), plot3D (>= 1.1), Rcpp (>= 0.11.0)

**LinkingTo** Rcpp

**RoxygenNote** 7.2.1

**Suggests** knitr, rmarkdown

**VignetteBuilder** knitr

**NeedsCompilation** yes

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**Repository** CRAN

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`ethseq.Analysis`*Ancestry analysis from whole-exome and targeted sequencing data*

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## Description

This function performs ancestry analysis of a set of samples and reports the results.

## Usage

```
ethseq.Analysis(  
  target.vcf = NA,  
  target.gds = NA,  
  bam.list = NA,  
  out.dir = tempdir(),  
  model.gds = NA,  
  model.available = NA,  
  model.assembly = "hg38",  
  model.pop = "All",  
  model.folder = tempdir(),  
  run.genotype = FALSE,  
  aseq.path = tempdir(),  
  mbq = 20,  
  mrq = 20,  
  mdc = 10,  
  cores = 1,  
  verbose = TRUE,  
  composite.model.call.rate = 1,  
  refinement.analysis = NA,  
  space = "2D",  
  bam.chr.encoding = FALSE  
)
```

## Arguments

|                              |   |
|------------------------------|---|
| <code>target.vcf</code>      | Path to the sample's genotypes in VCF format                                      |
| <code>target.gds</code>      | Path to the sample's genotypes in GDS format                                      |
| <code>bam.list</code>        | Path to a file containing a list of BAM files paths                               |
| <code>out.dir</code>         | Path to the folder where the output of the analysis is saved                      |
| <code>model.gds</code>       | Path to a GDS file specifying the reference model                                 |
| <code>model.available</code> | String specifying the pre-computed reference model to use                         |
| <code>model.assembly</code>  | String value indicating the assembly version to download for the pre-build models |
| <code>model.pop</code>       | String value indicating the population to download for the pre-build models       |

|                           |   |
|---------------------------|---|
| model.folder              | Path to the folder where reference models are already present or downloaded when needed     |
| run.genotype              | Logical values indicating whether the ASEQ genotype should be run                           |
| aseq.path                 | Path to the folder where ASEQ binary is available or is downloaded when needed              |
| mbq                       | Minimum base quality used in the pileup by ASEQ   |
| mrq                       | Minimum read quality used in the pileup by ASEQ   |
| mdc                       | Minimum read count acceptable for genotype inference by ASEQ                                |
| cores                     | Number of parallel cores used for the analysis  |
| verbose                   | Print detailed information  |
| composite.model.call.rate | SNP call rate used to run Principal Component Analysis (PCA)                                |
| refinement.analysis       | Matrix specifying a tree of ancestry sets   |
| space                     | Dimensions of PCA space used to infer ancestry (2D or 3D)                                   |
| bam.chr.encoding          | Logical value indicating whether input BAM files have chromosomes encoded with "chr" prefix |

**Value**

Logical value indicating the success of the analysis

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ethseq.RM

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*Create Reference Model for Ancestry Analysis*


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**Description**

This function creates a GDS reference model that can be used to perform EthSEQ ancestry analysis

**Usage**

```
ethseq.RM(
  vcf.fn,
  annotations,
  out.dir = "./",
  model.name = "Reference.Model",
  bed.fn = NA,
  verbose = TRUE,
  call.rate = 1,
  cores = 1
)
```

**Arguments**

|                          |   |
|--------------------------|---|
| <code>vcf.fn</code>      | vector of paths to genotype files in VCF format                     |
| <code>annotations</code> | data.frame with mapping of all samples names, ancestries and gender |
| <code>out.dir</code>     | Path to output folder   |
| <code>model.name</code>  | Name of the output model  |
| <code>bed.fn</code>      | path to BED file with regions of interest                           |
| <code>verbose</code>     | Print detailed information  |
| <code>call.rate</code>   | SNPs call rate cutoff for inclusion in the final reference model    |
| <code>cores</code>       | How many parallel cores to use in the reference model generation    |

**Value**

Logical value indicating the success of the analysis

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|                           |                                  |
|---------------------------|----------------------------------|
| <code>getModelList</code> | <i>List the models available</i> |
|---------------------------|----------------------------------|

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**Description**

This function prints the list of all available models.

**Usage**

```
getModelList()
```

**Value**

data.frame of all available models to use with specified assembly and population

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|                             |                                    |
|-----------------------------|------------------------------------|
| <code>getSamplesInfo</code> | <i>List the samples annotation</i> |
|-----------------------------|------------------------------------|

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**Description**

This function prints the list of all 1,000 Genomes Project samples used to build the reference models.

**Usage**

```
getSamplesInfo()
```

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