Package 'plasmut'

November 6, 2025

Type Package

Title Stratifying mutations observed in cell-free DNA and white blood cells as germline, hematopoietic, or somatic

Version 1.9.0

Description A Bayesian method for quantifying the liklihood that a given plasma mutation arises from clonal hematopoesis or the underlying tumor. It requires sequencing data of the mutation in plasma and white blood cells with the number of distinct and mutant reads in both tissues. We implement a Monte Carlo importance sampling method to assess the likelihood that a mutation arises from the tumor relative to non-tumor origin.

License Artistic-2.0
Encoding UTF-8
biocViews Bayesian, SomaticMutation, GermlineMutation, Sequencing
RoxygenNote 7.2.3
Imports tibble, stats, dplyr
Depends R (>= 4.3.0)
Suggests knitr, rmarkdown, tidyverse, ggrepel, magrittr, qpdf, BiocStyle, biocViews, testthat (>= 3.0.0)

VignetteBuilder knitr

Config/testthat/edition 3

git_url https://git.bioconductor.org/packages/plasmut

git_branch devel

git_last_commit e0d320c

git_last_commit_date 2025-10-29

Repository Bioconductor 3.23

Date/Publication 2025-11-05

Author Adith Arun [aut, cre], Robert Scharpf [aut]

Maintainer Adith Arun <adith.3.arun@gmail.com>

2 importance_sampler

Contents

crcse	eq	This	s da	ıta	is	an	ex	an	ıр	le	de	ıta	sei	t t	o s	sh	ow	h	oи	, to	าเ	ıse	e ti	he	po	ıcl	kaş	зe		
Index																														7
	wbc_somatic																													5
	plasmut																													
	plasma_somatic																													
	model_w																													3
	importance_sampler																													2
	crcseq																													2

Description

A cohort of metastatic colorectal cancer patients whose plasma and buffy coat were sequenced as part of the CAIRO5 trial. The cohort and analyses are described here: https://pubmed.ncbi.nlm.nih.gov/36534496/

Value

An example DNA sequencing dataset of matched plasma and wbc colorectal cancer samples croseq

 ${\it importance_sampler} \quad \quad {\it Importance\ sampler\ to\ estimate\ marginal\ likelihoods\ and\ Bayes\ factors}$

Description

Importance sampler to estimate marginal likelihoods and Bayes factors

Usage

```
importance_sampler(dat, params, save_montecarlo = TRUE)
```

Arguments

data frame with observed mutant and total counts and the analyte (plasma or buffy coat) it was taken from and the identifiers on what the mutation is (e.g.,

KRASG12C) and pt id

params list with ctc, ctdna and chip a and b beta parameters reflect beliefs on what frac-

tion of fragments belong to each class; montecarlo.samples being the number of MC samples; prior weight is the prior.weight reflects how much importance sampling to implement, closer to zero means more importance density consid-

ered

save_montecarlo

save more indepth monte carlo results

model_w 3

Value

implement importance sampling for a data set to assess probability of tumor derived mutations from sequencing results

Examples

model_w

Estimate the marginal likelihood that mutations in buffy coat and cfDNA reflect CH or correspond to germline mutations. If germline, the allele frequency should be 50 percent. The prior should be diffuse enough to handle CHIP mutations which are potentially way less than 50 percent

Description

Estimate the marginal likelihood that mutations in buffy coat and cfDNA reflect CH or correspond to germline mutations. If germline, the allele frequency should be 50 percent. The prior should be diffuse enough to handle CHIP mutations which are potentially way less than 50 percent

Usage

```
model_w(dat, params)
```

Arguments

dat

tibble containing vectors yand n. y and n should be named

params

a list with named elements that must include the following a which is the prior expectation for number of CH or germline variants observed in the sequencing data b which is the prior expectation for number of fragments reflecting CH or germline

Value

list of samples, probability densities, and likelihood for non-tumor assumption

plasma_somatic

Examples

plasma_somatic

Estimate the marginal likelihood that variants identified in cell-free DNA are derived from tumor cells (ctDNA-derived)

Description

Estimate the marginal likelihood that variants identified in cell-free DNA are derived from tumor cells (ctDNA-derived)

Usage

```
plasma_somatic(dat, params)
```

Arguments

dat tibble containing vectors 'y'and 'n'; 'y' and 'n' should be named

params a list with named elements that must include the following: 'a': prior expecta-

tion for number of plasma somatic variants observed in the plasma sequencing data 'b': prior expectation for number of plasma fragments not containing vari-

ants

Value

generate importance samples for plasma somatic model

Examples

plasmut 5

plasmut Bayesian models for estimating the origin of a sequenced DNA fragment

Description

The plasmut package provides a Bayesian importance sampling based approach to estimate the liklihood of a mutation arising from clonal hematopoiesis or tumor

wbc_somatic

Estimate the marginal likelihood of observing somatic mutations from CTCs present in buffy coat $p(y_w \mid theta_w, n_w, model_S) x$ $p(theta_w \mid model_S) theta_w \mid model_S \sim beta(1, 999) \# sequencing error or CTC$

Description

Estimate the marginal likelihood of observing somatic mutations from CTCs present in buffy coat $p(y_w \mid theta_w, n_w, model_S) \times p(theta_w \mid model_S) \times p(thet$

Usage

```
wbc_somatic(dat, params)
```

Arguments

dat tibble containing vectors 'y' and 'n'; 'y' and 'n' should be named

params a list with named elements that must include the following: 'a': prior expec-

tation for number of somatic variants observed in the WBC sequencing data (either by error or from a CTC) 'b': prior expectation for number of WBCs not

containing the variant

Value

generate importance samples for wbc somatic model

6 wbc_somatic

Examples

Index

```
crcseq, 2
importance_sampler, 2
model_w, 3
plasma_somatic, 4
plasmut, 5
wbc_somatic, 5
```