Package 'MPXV'

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Title Variable Number Tandem Repeats Analysis
Version 0.1.0
Description Analysis of calling Variable Number Tandem Repeats for the genome sequence of mon keypox virus and virus tracking.
LazyData true
Imports seqinr, stringr, Biostrings
RoxygenNote 7.2.1
R topics documented:

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 VNTRcaller
                Calling Variable Number Tandem Repeats (VNTR) for the genome se-
                quence of monkeypox virus (MPXV)
```

Description

The function VNTRcaller estimates the copy numbers of VNTRs.

Usage

```
VNTRcaller(
  data,
  vntr = vntr,
  match_s = match_s,
  mismatch_s = mismatch_s,
  regionStart = regionStart,
  regionEnd = regionEnd,
  baseonly = T,
  fseqExtend = fseqExtend,
  VNTRoutput = F,
  tracker = F
)
```

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Arguments

data MPXV sequences from a file in FASTA format

vntr sequence of the repetitive unit

match_s matching weight mismatch_s mismatching penalty

regionStart starting position of a VNTR region regionEnd ending position of a VNTR region

baseonly logical. If TRUE, only the letters of the nucleotide bases (i.e. A,C,G,T) are

considered. If FALSE, the degenerate codes are also considered.

fseqExtend the length of extended flanking sequence

VNTRoutput logical. If TRUE, the output is written to .csv files. tracker logical. If TRUE, call function VNTRtracker.

Value

ID name of MPXV sequencesr copy of tandem repeatsmatch number of matchesmismatch number of mismatches

indel number of insertions and deletions (indels)

score alignment score of a VNTR region for a query strain start_pos starting position of the VNTR region for a query strain

References

Yang, H.-C. et al (2022) Monkeypox genome contains variable number tandem repeats enabling accurate virus tracking.

Pages H, Aboyoun P, Gentleman R, DebRoy S (2022) *Biostrings: Efficient manipulation of biological strings*. R package version 2.64.0.

Examples

```
## Read MPXV example sequences from a file in FASTA format.
MPXVseq <- read.fasta(system.file("extdata/MPXV_seq_example.fasta.gz", package = "MPXV"), as.string = T)

## VNTR
vntr <- c("T", "TATGATGGA", "AT", "ATATACATT")
regionStart <- c(132436,150542,173240,178413)
regionEnd <- c(133216,151501,173320,179244)

## parameter settings
match_s = 2
mismatch_s = -5
baseonly = TRUE
fseqExtend = 150
VNTRoutput = FALSE
tracker = FALSE

## computes the copy of the variable number tandem repeats</pre>
```

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For more details about the output of VNTRcaller, please vitsit https://github.com/teresayang/MPXV_VNTR.

VNTRtracker

Tracking the origin of monkeypox virus (MPXV) by using Variable Number Tandem Repeats (VNTR) markers

Description

The function VNTRtracker calculates the MPXV strain-strain genetic distance to track the origin of a given MPXV by using VNTRs.

Usage

```
VNTRtracker(r, file_name = "Tracking_output")
```

Arguments

r A vector of copies of VNTR of MPXV.

file_name Filename of the output.

Details

The four VNTR markers nt133095[A25R - A26L] (T), nt150560[A47R - A49R] (TATGATGGA), nt173267[B14R - B15L] (AT), and nt179074[B18R - B19R] (ATATACATT) are proposed to track the original of MPXV strains. Based on these four VNTRs, we calculate three distances (ploymorphism information content, entropy, and length of repetitive unit (see Yang et al. (2022)) between a query MPXV strain and the strains in our database.

Value

The output contains multiple columns, including accession ID, country name, collection date, clade, lineage, and three distances mentioned above.

References

Yang, H.-C. et al (2022) Monkeypox genome contains variable number tandem repeats enabling accurate virus tracking.

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Examples

```
r=c(27,7,24,16)
```

The output is a csv file with the file name given by the parameter "file_name".

In this file, each row represents a strain in our database, and column variables contain

accession ID, country name, collection date, clade, lineage, and three distances (see Yang et al. (2022))

between a query MPXV strain and a strains in our database.

The rank of strains are ordered by ploymorphism information content distance (see Yang et al. (2022)) from low VNTRtracker(r, file_name="Tracking_output")

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