Package 'CleanBSequences'

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Type Package
Title Curing of Biological Sequences
Version 2.3.0
Author Florencia I. Pozzi, Silvina A. Felitti
Maintainer Florencia I. Pozzi <florenciapoz@hotmail.com></florenciapoz@hotmail.com>
Description Curates biological sequences massively, quickly, without errors and without internet connection. Biological sequences curing is performed by aligning the forward and / or revers primers or ends of cloning vectors with the sequences to be cleaned. After the alignment, new subsequences are generated without biological fragment not desired by the user. Pozzi et al (2020) <doi:10.1007 s00438-020-01671-z="">.</doi:10.1007>
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Curing of biological sequences

Description

Curates biological sequences of primer reverse. This cleaning is required for techniques such as cDNA-AFLP.

Usage

```
DNAStringSetOPR(SEQs, PrimerR)
```

Arguments

SEQs file with fasta format containing biological sequences that are to be cleaned.

PrimerR dnastring containing the reverse primer/vector sequences to be removed.

Value

clean biological sequences and visualization of the alignments

Author(s)

Florencia I Pozzi, Silvina A. Felitti

Examples

```
SEQs = readDNAStringSet(system.file("sequences","SeqInputOPR.fasta", package = "CleanBSequences"))
PrimerR= DNAString ("GACTGCGTACCATGC")
DNAStringSetOPR (SEQs,PrimerR)
```

DNAStringSetTPR

Curing of biological sequences

Description

Curates biological sequences of two restriction enzyme primers or cloning vectors. This cleaning is required for techniques such as cDNA-AFLP. This cleaning is required for techniques such as cDNA-AFLP.

Usage

```
DNAStringSetTPR(SEQs, PrimerF, PrimerR)
```

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Arguments

SEQs file with fasta format containing biological sequences that are to be cleaned.

PrimerF dnastring containing the foward primer/vector sequences to be removed.

PrimerR dnastring containing the reverse primer/vector sequences to be removed.

Value

clean biological sequences and visualization of the alignments

Author(s)

Florencia I Pozzi, Silvina A. Felitti

Examples

```
SEQs = readDNAStringSet(system.file("sequences","SeqInputTPR.fasta", package = "CleanBSequences"))
PrimerR= DNAString ("GACTGCGTACCATGC")
PrimerF = DNAString("GATGAGTCCTGACCGAA")
DNAStringSetTPR (SEQs,PrimerF,PrimerR)
```

OnePrimerRemove

Clean biological secuences

Description

Curates biological sequences of primer reverse. This cleaning is required for techniques such as cDNA-AFLP.

Usage

```
OnePrimerRemove(SEQs, PrimerR)
```

Arguments

SEQs dnastring containing biological sequences that are to be cleaned.

PrimerR dnastring containing the reverse primer/vector sequences to be removed.

Value

clean biological sequences and visualization of the alignments

Author(s)

Florencia I Pozzi and Silvina A. Felitti

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Examples

```
SEQs = DNAString(paste("GCCTCGCCTCCTCTTTGATCAGCTTCGCATATCAGGCAACAGCTCAATTT",
"GGTACTTGTTCAAATAAGCATTTAGACCATCTGTTCCAAGAACCTTTGCAATCTT",
"CACAAGGTGGTCATGGTACGCAGTC", sep=""))
PrimerR= DNAString("GACTGCGTACCATGC")
OnePrimerRemove (SEQs,PrimerR)
```

TwoPrimerRemove

Clean biological secuences

Description

Curates biological sequences of two restriction enzyme primers or cloning vectors. This cleaning is required for techniques such as cDNA-AFLP.

Usage

```
TwoPrimerRemove(SEQs, PrimerF, PrimerR)
```

Arguments

SEQs DNAString containing biological sequences that are to be cleaned.

PrimerF dnastring containing the foward primer/vector sequences to be removed.

PrimerR dnastring containing the reverse primer/vector sequences to be removed.

Value

clean biological sequences and visualization of the alignments

Author(s)

Florencia I Pozzi, Silvina A. Felitti

Examples

```
SEQs = DNAString(paste("ACTTTCTGCTGCTTGTGGTCGCAATCAGAGTCCTGATGATGAGTCCTGA",
"CCGAACCCTTTTTCTCCGTCATCCGTTGGTCCATGGTACGCAATCAGAG", sep = ""))
PrimerF = DNAString("GATGAGTCCTGACCGAA")
PrimerR = DNAString("GACTGCGTACCATGC")
TwoPrimerRemove (SEQs,PrimerF,PrimerR)
```

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