NAME

SequenceFileUtil

SYNOPSIS

```
use SequenceFileUtil;
use SequenceFileUtil qw(:all);
```

DESCRIPTION

SequenceFileUtil module provides the following functions:

Are Sequence Lengths I dentical, Calcuate Percent Sequence I dentity, Calculate Percent Sequence I dentity Matrix, Get Longest Sequence, Get Sequence Length, Get Shortest Sequence, Is Clustal W Sequence File, Is Gap Residue, Is MSF Sequence File, Is PIRFasta Sequence File, Is Pearson Fasta Sequence File, Is Supported Sequence File, Read Clustal W Sequence File, Read MSF Sequence File, Read PIRFasta Sequence File, Read Pire Read Sequence File, Remove Sequence Alignment Gap Columns, Remove Sequence Gaps, Write Pearson Fasta Sequence File Sequence File Util module provides various methods to process sequence files and retreive appropriate information.

FUNCTIONS

Are Sequence Lengths I dentical

```
$Status = AreSequenceLengthsIdentical($SequencesDataRef);
```

Checks the lengths of all the sequences available in SequencesDataRef and returns 1 or 0 based whether lengths of all the sequence is same.

CalcuatePercentSequenceI dentity

```
$PercentIdentity =
AreSequenceLengthsIdenticalAreSequenceLengthsIdentical(
    $Sequence1, $Sequence2, [$IgnoreGaps, $Precision]);
```

Returns percent identity between *Sequence1* and *Sequence2*. Optional arguments *IgnoreGaps* and *Precision* control handling of gaps in sequences and precision of the returned value. By default, gaps are ignored and precision is set up to 1 decimal.

CalculatePercentSequenceI dentityMatrix

Calculate pairwise percent identity between all the sequences available in *SequencesDataRef* and returns a reference to identity matrix hash. Optional arguments *IgnoreGaps* and *Precision* control handling of gaps in sequences and precision of the returned value. By default, gaps are ignored and precision is set up to 1 decimal.

GetSequenceLength

```
$SeqquenceLength = GetSequenceLength($Sequence, [$IgnoreGaps]);
```

Returns length of the specified sequence. Optional argument *IgnoreGaps* controls handling of gaps. By default, gaps are ignored.

GetShortestSequence

```
($ID, $Sequence, $SeqLen, $Description) = GetShortestSequence(
    $SequencesDataRef, [$IgnoreGaps]);
```

Checks the lengths of all the sequences available in \$SequencesDataRef and returns \$ID, \$Sequence, \$SeqLen, and \$Description values for the shortest sequence. Optional arguments \$IgnoreGaps controls handling of gaps in sequences. By default, gaps are ignored.

GetLongestSequence

```
($ID, $Sequence, $SeqLen, $Description) = GetLongestSequence(
    $SequencesDataRef, [$IgnoreGaps]);
```

Checks the lengths of all the sequences available in *SequencesDataRef* and returns ID, Sequence, SeqLen, and Description values for the longest sequence. Optional argument \$*IgnoreGaps* controls handling of gaps in sequences. By default, gaps are ignored.

IsGapResidue

```
$Status = AreSequenceLengthsIdentical($Residue);
```

Returns 1 or 0 based on whether *Residue* corresponds to a gap. Any character other than A to Z is considered a gap residue.

IsSupportedSequenceFile

```
$Status = IsSupportedSequenceFile($SequenceFile);
```

Returns 1 or 0 based on whether SequenceFile corresponds to a supported sequence format.

IsClustalWSequenceFile

```
$Status = IsClustalWSequenceFile($SequenceFile);
```

Returns 1 or 0 based on whether SequenceFile corresponds to Clustal sequence alignment format.

IsPearsonFastaSequenceFile

```
$Status = IsPearsonFastaSequenceFile($SequenceFile);
```

Returns 1 or 0 based on whether SequenceFile corresponds to Pearson FASTA sequence format.

IsPIRFastaSequenceFile

```
$Status = IsPIRFastaSequenceFile($SequenceFile);
```

Returns 1 or 0 based on whether SequenceFile corresponds to PIR FASTA sequence format.

IsMSFSequenceFile

```
$Status = IsClustalWSequenceFile($SequenceFile);
```

Returns 1 or 0 based on whether SequenceFile corresponds to MSF sequence alignment format.

ReadSequenceFile

```
$SequenceDataMapRef = ReadSequenceFile($SequenceFile);
```

Reads SequenceFile and returns reference to a hash containing following key/value pairs:

```
$SequenceDataMapRef->{IDs} - Array of sequence IDs
$SequenceDataMapRef->{Count} - Number of sequences
$SequenceDataMapRef->{Description}{$ID} - Sequence description
$SequenceDataMapRef->{Sequence}{$ID} - Sequence for a specific ID
$SequenceDataMapRef->{Sequence}{InputFileType} - File format
```

ReadClustalWSequenceFile

```
$SequenceDataMapRef = ReadClustalWSequenceFile($SequenceFile);
```

Reads ClustalW *SequenceFile* and returns reference to a hash containing following key/value pairs as describes in ReadSequenceFile method.

ReadMSFSequenceFile

```
$SequenceDataMapRef = ReadMSFSequenceFile($SequenceFile);
```

Reads MSF SequenceFile and returns reference to a hash containing following key/value pairs as describes in ReadSequenceFile method.

ReadPI RFastaSequenceFile

```
$SequenceDataMapRef = ReadPIRFastaSequenceFile($SequenceFile);
```

Reads PIR FASTA *SequenceFile* and returns reference to a hash containing following key/value pairs as describes in ReadSequenceFile method.

ReadPearsonFastaSequenceFile

```
$SequenceDataMapRef = ReadPearsonFastaSequenceFile($SequenceFile);
```

Reads Pearson FASTA SequenceFile and returns reference to a hash containing following key/value pairs as describes in ReadSequenceFile method.

RemoveSequenceGaps

```
$SeqWithoutGaps = RemoveSequenceGaps($Sequence);
```

Removes gaps from Sequence and return a sequence without any gaps.

RemoveSequenceAlignmentGapColumns

Using input alignment data map ref containing following keys, generate a new hash with same set of keys after residue columns containg only gaps have been removed:

```
{IDs} : Array of IDs in order as they appear in file
{Count}: ID count
{Description}{$ID} : Description data
{Sequence}{$ID} : Sequence data
```

WritePearsonFastaSequenceFile

Using sequence data specified via *SequenceDataRef*, write out a Pearson FASTA sequence file. Optional argument *MaxLength* controls maximum length sequence in each line; default is 80.

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SEE ALSO

PDBFileUtil.pm

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