# Package 'pqsfinder' 

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## Type Package

Title Identification of potential quadruplex forming sequences
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Description Pqsfinder detects
DNA and RNA sequence patterns that are likely to fold into an intramolecular G-quadruplex (G4). Unlike many other approaches, pqsfinder is able to detect G4s
folded from imperfect G-runs containing bulges or mismatches or G4s having long loops. Pqsfinder also assigns an integer score to each hit that was fitted on G4 sequencing data and corresponds to expected stability of the folded G4.
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## $R$ topics documented:

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as.character, PQSViews-method
Coerce to character vector

## Description

Coerce to character vector

## Usage

\#\# S4 method for signature 'PQSViews'
as.character (x)

## Arguments

$x \quad$ PQSViews object.

## Value

Character vector representing PQS.

```
density,PQSViews-method
```

    Get density vector
    
## Description

Desity vector represents numbers of PQS (potential quadruplex forming sequences) overlapping at each position in input sequence.

## Usage

\#\# S4 method for signature 'PQSViews' density (x)

## Arguments

$x \quad$ PQSViews object.

## Value

Density vector.

## Examples

pqs <- pqsfinder(DNAString("CCCCCCGGGTGGGTGGGTGGGAAAA")) density(pqs)
maxScores Get vector of maximal scores

## Description

Get vector of maximal scores for a given object.

## Usage

maxScores(x, ...)

## Arguments

x
An object.
... Additional arguments, for use in specific methods.

## Value

Vector of maximal scores.

## Examples

```
    showMethods("maxScores")
```

maxScores, PQSViews-method
Get vector of maximal scores

## Description

For each sequence position it gives the maximal score of all PQS conformations which overlap that position.

## Usage

\#\# S4 method for signature 'PQSViews'
maxScores ( x )

## Arguments

x
PQSViews object.

## Value

Vector of maximal scores.

## Examples

```
pqs <- pqsfinder(DNAString("CCCCCCGGGTGGGTGGGTGGGAAAA"))
maxScores(pqs)
```

pqsfinder Identify potential quadruplex forming sequences.

## Description

Function for identification of all potential intramolecular quadruplex patterns (PQS) in DNA or RNA sequence.

```
Usage
    pqsfinder(
        subject,
        strand = "*",
        overlapping = FALSE,
        max_len = 50L,
        min_score = 47L,
        run_min_len = 2L,
        run_max_len = 11L,
        loop_min_len = 0L,
        loop_max_len = 30L,
        max_bulges = 3L,
        max_mismatches = 3L,
        max_defects = 3L,
        tetrad_bonus = 40L,
        mismatch_penalty = 28L,
        bulge_penalty = 20L,
        bulge_len_factor = 0.2,
        bulge_len_exponent = 1,
        loop_mean_factor = 6.6,
        loop_mean_exponent = 0.8,
        run_re = "G{1,10}.{0,9}G{1,10}",
        custom_scoring_fn = NULL,
        use_default_scoring = TRUE,
        deep = FALSE,
        verbose = FALSE
)
```


## Arguments

subject DNAString or RNAString object.
strand Strand specification. Allowed values are "+", "-" or "*", where the last one represents both strands. Implicitly, the input DNAString object is assumed to encode the " + " strand.
overlapping If true, than all overlapping PQS will be reported.
max_len Maximal lenth of PQS.
min_score Minimal PQS score. The default value 52 shows the best balanced accuracy on G4 sequencing data provided by Chambers et al. 2015.
run_min_len Minimal length of quadruplex run.
run_max_len Maximal length of quadruplex run.
loop_min_len Minimal length of quadruplex loop. Unless the default scoring system is disabled, at most one loop can have zero length.
loop_max_len Maxmimal length of quadruplex loop.
max_bulges Maximal number of runs with bulge.
max_mismatches Maximal number of runs with mismatch.

```
max_defects Maximum number of defects in total (max_bulges + max_mismatches).
tetrad_bonus Score bonus for one complete G tetrade.
mismatch_penalty
    Penalization for a mismatch in tetrad.
bulge_penalty Penalization for a bulge in quadruplex run.
bulge_len_factor
    Penalization factor for a bulge length.
bulge_len_exponent
    Exponent of bulge length.
loop_mean_factor
    Penalization factor of loop length mean.
loop_mean_exponent
    Exponent of loop length mean.
run_re Regular expression specifying one run of quadruplex.
custom_scoring_fn
                    Custom quadruplex scoring function. It takes the following 10 arguments: subject
            - Input DNAString or RNAString object, score - implicit PQS score, start -
            PQS start position, width - PQS width, loop_1 - start pos. of loop #1, run_2
            - start pos. of run #2, loop_2 - start pos. of loop #2, run_3 - start pos. of run
            #3, loop_3 - start pos. of loop #3, run_4 - start pos. of run #4. Return value
            of the function has to be new score represented as a single integer value. Please
                note that if use_default_scoring is enabled, the custom scoring function is
                evaluated AFTER the default scoring system but ONLY IF the default scoring
                system resulted in non-zero score (for performance reasons). On the other hand,
                when use_default_scoring is disabled, custom scoring function is evaluated
                on every PQS.
use_default_scoring
                            Enables default internal scoring system. This option is particularly useful in
                        case you intend to radically change the default behavior and specify your own
                    scoring function. By disabling the default scoring you will get a full control
                    above the underlying detection algorithm.
deep Perform deep search. With this option enabled, maxScores and density vectors
        are computed. Deep search is much more computationaly demanding.
verbose Enables detailed output. Turn it on if you want to see all possible PQS found
        at each positions and not just the best one. It is highly recommended to use
        this option for debugging custom quadruplex scoring function. Each PQS is
        reported on separate row in the following format: start cnt pqs_sequence
        score, where start is the PQS starting position, pqs_sequence shows the PQS
        sequence structure with each run surrounded by square brackets and score is the
        score assigned to the particular PQS by all applied scoring functions.
```


## Details

Use elementMetadata function to get extra PQS features like number of tetrads (nt), bulges (nb), mismatches ( nm ) or loop lengths ( $111,112,113$ ).

## Value

PQSViews object

## Examples

```
pv <- pqsfinder(DNAString("CCCCCCGGGTGGGTGGGTGGGTAAAA"))
pv
elementMetadata(pv)
```

PQSViews PQSViews class constructor

## Description

User friendly constructor for PQSViews class representing potential quadruplex forming sequences (PQS). PQSViews is a subclass of XStringViews class and adds two more slots to store PQS density and PQS score distribution.

## Usage

PQSViews( subject, start, width, strand, score, density, max_scores, $n t$,
nb,
nm,
rl1, rl2, rl3, 111, 112, 113
)

## Arguments

| subject | DNAString or RNAString object. |
| :--- | :--- |
| start | Start positions. |
| width | Lengths. |
| strand | Strand specifications. |


| score | Scores. |
| :--- | :--- |
| density | Numbers of PQS overlapping at each position in subject. |
| max_scores | Score of the best PQS found at each position. |
| nt | Tetrad numbers. |
| nb | Bulge counts. |
| nm | Mismatch counts. |
| $\mathrm{rl1}$ | Run 1 lengths. |
| $\mathrm{rl2}$ | Run 2 lengths. |
| $\mathrm{rl3}$ | Run 3 lengths. |
| 111 | Loop 1 lengths. |
| 112 | Loop 2 lengths. |
| 113 | Loop 3 lengths. |

## Details

Use elementMetadata function to get extra PQS features like number of tetrads, bulges, mismatches or loop lengths.

## Value

PQSViews object.

## Examples

```
pv <- PQSViews(DNAString("GGTGGTGGTGG"), 1, 11, "+", 33, as.integer(rep(1, 11)),
    as.integer(rep(33, 11)), 2, 0, 0, 2, 2, 2, 1, 1, 1)
start(pv)
width(pv)
strand(pv)
score(pv)
density(pv)
maxScores(pv)
elementMetadata(pv)
```


## Description

Represents potential quadruplex forming sequences found by pqsfinder function. This is a subclass of XStringViews-class class and adds one more slot.

## Slots

density Numbers of PQS (potential quadruplex forming sequences) overlapping at each position in input sequence.
max_scores Score of the best PQS found at each position.
score, PQSViews-method Get PQS score vector

## Description

Get PQS score vector

## Usage

\#\# S4 method for signature 'PQSViews' score (x)

## Arguments

x
PQSViews object.

## Value

Score vector.

## Examples

```
pqs <- pqsfinder(DNAString("CCCCCCGGGTGGGTGGGTGGGAAAA"))
score(pqs)
```

show, PQSViews-method Show method

## Description

Show method

## Usage

\#\# S4 method for signature 'PQSViews' show(object)

## Arguments

object PQSViews object.

## Value

PQSViews object printed.
strand, PQSViews-method
Get PQS strand vector

## Description

Get PQS strand vector

## Usage

\#\# S4 method for signature 'PQSViews' strand ( X )

## Arguments

x
PQSViews object.

## Value

Strand vector.

## Examples

```
pqs <- pqsfinder(DNAString("CCCCCCGGGTGGGTGGGTGGGAAAA"))
strand(pqs)
```


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