

# Searching a DNA sequence using the `matchPattern` method (work in progress)

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November 7, 2006

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## 1 Load a genome

Load the *Caenorhabditis elegans* genome:

```
> library(BSgenome.Celegans.UCSC.ce2)
> ls(2)
```

```
[1] "Celegans"
```

```
> Celegans
```

Caenorhabditis elegans genome:

```
[1] "chrI"      "chrII"      "chrIII"      "chrIV"      "chrV"
[6] "chrM"      "chrX"      "upstream1000" "upstream2000" "upstream5000"
```

(use the '\$' or '[' operator to access a given chromosome)

```
> comment(Celegans$chrI)
```

```
[1] "Caenorhabditis elegans - chromosome I (generated from FASTA file chrI.fa)"
```

Display chromosome I:

```
> Celegans$chrI
```

Views on a 15080483-letter DNAString subject

Subject: GCCTAAGCCTAAGCCTAAGCCTAAGCCTAAGCCT...GGCTTAGGCTTAGGCTTAGGTTTAGGCTTAGGC

Views:

	first	last	width
[1]	1	15080483	15080483 [GCCTAAGCCTAAGCCTAAGCCTA...TAGGCTTAGGTTTAGGCTTAGGC]

Note that `Celegans$chrI` is a *BStringViews* object with a single view (`length(Celegans$chrI)` is 1). The number of letters in this sequence can be retrieved with:

```
> cI <- Celegans$chrI[[1]]
> length(cI)

[1] 15080483
```

Try with an *upstream* object:

```
> length(Celegans$upstream2000)

[1] 992

> length(Celegans$upstream2000[[1]])

[1] 2000
```

Some basic stats:

```
> af <- alphabetFrequency(cI)
> af
```

A	C	G	T	-	other
4838561	2697177	2693544	4851201	0	0

```
> sum(af) == length(cI)

[1] TRUE
```

## 2 Find patterns in a DNA sequence

To find all exact matches of pattern "ACCCAGGGC":

```
> p <- "ACCCAGGGC"
> countPattern(p, cI)

[1] 0

> countPattern(p, cI, mismatch = 1)

[1] 235
```

The matches can be stored in a *BStringViews* object by using the `matchPattern` method:

```
> m <- matchPattern(p, cI, mismatch = 1)
> m[4:6]
```

```
Views on a 15080483-letter DNAString subject
Subject: GCCTAAGCCTAAGCCTAAGCCTAAGCCTAAGCCT...GGCTTAGGCTTAGGCTTAGGTTTAGGCTTAGGC
Views:
```

```
      first  last width
[1] 187350 187358      9 [ACCCAAGGC]
[2] 213236 213244      9 [ACCCAGGGG]
[3] 424133 424141      9 [ACCCAGGAC]
```

```
> mismatch(p, m[4:6])
```

```
[[1]]
[1] 6
```

```
[[2]]
[1] 9
```

```
[[3]]
[1] 8
```

The `mismatch` method (new in *Biostrings* 2) returns the positions of the mismatching letters.

Note: The `mismatch` method is in fact a particular case of a (vectorized) *alignment* function where only “replacements” are allowed. Current implementation is slow but (but this will change).

It may happen that a match is *out of limits* like here:

```
> p2 <- DNAString("AAGCCTAAGCCTAAGCCTAA")
> m2 <- matchPattern(p2, cI, mismatch = 2)
> m2[1:4]
```

```
Views on a 15080483-letter DNAString subject
Subject: GCCTAAGCCTAAGCCTAAGCCTAAGCCTAAGCCT...GGCTTAGGCTTAGGCTTAGGTTTAGGCTTAGGC
Views:
```

```
      first last width
[1]    -1   18    20 [ GCCTAAGCCTAAGCCTAA]
[2]     5   24    20 [AAGCCTAAGCCTAAGCCTAA]
[3]    11   30    20 [AAGCCTAAGCCTAAGCCTAA]
[4]    17   36    20 [AAGCCTAAGCCTAAGCCTAA]
```

```
> p2 == m2[1:4]
```

```
[1] FALSE  TRUE  TRUE  TRUE
```

```
> mismatch(p2, m2[1:4])
```

```
[[1]]
[1] 1 2
```

```
[[2]]
integer(0)
```

```
[[3]]
integer(0)
```

```
[[4]]
integer(0)
```

The list of exact matches and the list of inexact matches can both be obtained with:

```
> m2[p2 == m2]
> m2[p2 != m2]
```

Note that the length of `m2[p2 == m2]` should be equal to `countPattern(p2, cI, mismatch=0)`.

### 3 A note on performance

If needed, the `matchPattern` and `countPattern` methods convert their first argument (the pattern) to an object of the same class than their second argument (the subject) before they pass it to the function that actually implements the fast search algorithm.

So if you need to reuse the same pattern a high number of times, it's a good idea to convert it *before* to pass it to the `matchPattern` or `countPattern` method. This way the conversion is done only once:

```
> library(hgu95av2probe)
> tmpseq <- BStringViews(hgu95av2probe$sequence, "DNASTring")
> someStats <- function(v) {
+   GC <- DNASTring("GC")
+   CG <- DNASTring("CG")
+   sapply(1:length(v), function(i) {
+     y <- v[i]
+     c(alphabetFrequency(y)[1:4], GC = countPattern(GC, y),
+       CG = countPattern(CG, y))
+   })
+ }
> someStats(tmpseq[1:10])
```

	[,1]	[,2]	[,3]	[,4]	[,5]	[,6]	[,7]	[,8]	[,9]	[,10]
A	2	6	5	5	5	4	6	8	5	6
C	9	9	9	8	3	9	7	5	8	7
G	5	7	7	6	10	8	4	6	5	8
T	9	3	4	6	7	4	8	6	7	4
GC	3	4	4	2	1	3	0	0	2	2
CG	0	0	0	0	0	0	0	0	0	0