Sequence manipulation and scanning

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Abstract

Sequences stored as XStringSet objects (from the Biostrings package) can be used by several functions in the universalmotif package. These functions are demonstrated here and fall into two categories: sequence manipulation and motif scanning. Sequences can be generated, shuffled, and background frequencies of any order calculated. Scanning can be done simply to find locations of motif hits above a certain threshold, or to find instances of enriched motifs.

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1 Introduction

This vignette goes through generating your own sequences from a specified background model, shuffling sequences whilst maintaining a certain k-let size, and the scanning of sequences and scoring of motifs. For an introduction to sequence motifs, see the introductory vignette. For a basic overview of available motif-related functions, see the motif manipulation vignette. For a discussion on motif comparisons and P-values, see the motif comparisons and P-values vignette.

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2 Creating random sequences

The Biostrings package offers an excellent suite of functions for dealing with biological sequences. The universalmotif package hopes to help extend these by providing the create_sequences() and shuffle_sequences() functions. The first of these, create_sequences(), in it's simplest form generates a set of letters in random order, then passes these strings to the Biostrings package. The number and length of sequences can be specified. The probabilities of individual letters can also be set.

The freqs option of create_sequences() also takes higher order backgrounds. In these cases the sequences are constructed in a Markov-style manner, where the probability of each letter is based on which letters precede it.

```
library(universalmotif)
library(Biostrings)
## Create some DNA sequences for use with an external program (default
## is DNA):
sequences.dna <- create_sequences(seqnum = 500,</pre>
                                  freqs = c(A=0.3, C=0.2, G=0.2, T=0.3))
## writeXStringSet(sequences.dna, "dna.fasta")
sequences.dna
#> DNAStringSet object of length 500:
#>
         width seq
#>
     [1]
           100 CTTATGATACGTAAGTAGGGCCGCCGCCGC...TTTAGCATGGGATTTATCCATTCCAATCCATC
#>
     [21]
           100 TCCAGAATATATCACTACTAAAATTATGAGTAT...GTTACTAGCCACGTTGTTGAGGTTAAGTTCTG
           100 AAACCGCTAATGGAATAAGATGTCAGCAGACGC...AGTAAAAGATGGGTGTTCAAGCCCAAAAATGT
     [3]
#>
#>
     [4]
           100 GTTCAACTATCGGATTATCCCACTCATCACTAT...ACTATGACTGATCTGAGTCCCATTAAAGTTGA
     Γ51
#>
           100 TTTGTGGTAAACTCCTACCCTGACTTAGGGAGA...TCTACTAACAGAATCAGCTCATCCAATAAACG
#>
#> [496]
           100 CACCGAACTTGTAAATCGGTGATGGAAGCAGCT...TTCTTTGTTAGAAATTTCTATCAGACTATGTA
           100 TTAGCTCCAAAGAATTCAGTAGGAAGCATGATC...GTCTGAGGGTAGTATATTACGAGTACGTCTTC
#> [497]
#> [498]
           100 AGTGAACCAGTCCCCGCTCATAAGCATCCTGCT...CGCTAGGGCCGTGAGCGCCCCTACAGCATAGT
           100\ GCGTTTGCAACCGCAGCGAACGTATGCGTATTA\dots AGCATTGCAAATGAGGGTTGATGTAGAGAAGA
#> [499]
100 TACTGATCACTATCTGCAAATACGAATAGGCCG...TCCAGCGATTGACACTCAAGACACAGTTGGCC
## Amino acid:
create_sequences(alphabet = "AA")
#> AAStringSet object of length 100:
#>
         width seq
#>
           100 QQPGFDNKVHHMYHPKIGFWTEVTTYFKWHEAL...SVPFTLAGPALDRDHCKCKWNCHKWQYETRIN
#>
     [2]
           100 HIFNLGDVSPPCYQGVSMKVPIRPPWLVVQIAW...MRGKQYCNGCYFLFPDVDVVDDPVVIYHPKRD
     [3]
           100 YATTRKQGPWYNYYVHDSPTKNNLKVQHTANAI...GNVPLLCVWCLHEHYEHEHTQEWGTCYMLDCQ
#>
#>
     [4]
           100 QRKDYNGRMFGDYHMSMCTRFSKGFVWSSISCT...AKMTHWDENDWLWKGFSFSSFFFRRSYQGTKG
#>
     [5]
           100 HKAIERTEINPQYPEFVHAQAYHCCTEFRRYCG...SGDAEKDLFDKNPMPGFHFQSHNDQLWVCMST
#>
#>
    [96]
           100 VIVMYHIGIVDAIMPMFKTVGKNNWRERSYLGF...TFFPRPLSETKEPAVLNVTCQWWWPRYRLPAE
#> [97]
           100 NCLTFLYRGEKMHVGYPQATCPLKRQKERHQHQ...PCTTNDMCTTVHHDEMAYGAFYGIMKYVGHIS
           100\ FSCCLPMDDMSCHFVKYWFSVTHFMQPPQPVHD...IWLAINNILVIKCFLPLARYSANSLDYCDARH
   [98]
           100 VKQIRSCNATCNHMMVIEKRRAECHPVCPYCHN...DSCFFCNQDVVMSHTQWCDWICVFKSWFWRDV
#> [99]
#> [100]
           100 MDGPWWQAVDKDHVEHRLPPMFAVENDMNHGHA...VPRKANPYSWHQMKDRIDNTWEEPHLWKRLLK
## Any set of characters can be used
```

```
create_sequences(alphabet = paste0(letters, collapse = ""))
#> BStringSet object of length 100:
#>
         width seq
#>
           100 xawpwqikhoetkjsknawxuclwlfmrkzuna...dndyqplbxzamqjchdqfxumaibpkpmchh
#>
           100 vbsetnrvocimutlvbatvqfwtwlyjvzoaa...hbgxhfxcvyaymsephhlvrzbrcfueyfop
#>
     [3]
           100 scotgtagwamfecdgpbgsmiipirlbfyina...lpjvyvjdsxaktcgwlyrtnmcaeuftkivw
#>
     [4]
           100 \ qdkjnajrdeqzpmwrcbmqhltmtxxtqydaa...pcmuplveqxaxzlieopxqjzcjfkpiwlde
#>
     [5]
           100 \,\, ndgykhscksuszwocgbjodoejfdklayxna...tqptfbhqowajfvkmsqdofmdshzaxiokl
#>
#>
    [96]
           100 xuijunueciedbokjpkujcsxqdywzwrval...emqxyivlqqbnlxodsqlputqbilnuklzz
#>
    [97]
           100 uveysuepjwiwmxdudkshxvjnoejrhqqnm...iztwoyqmdfbarqqlvhrmqqrjkaxjwoqq
    [98]
           100 swanpanarkmpwhufqloetyujakujrqkam...mnwufosobebmyqstzyxkmtsslqiyiqoo
    [99]
#>
           100 pwxdmhwlyyqjgroqellcobgglqhacqenm...qbztweepzdbyezuadpdiigtbnfsnutvv
  [100]
           100 nxtsjnewfnucragasmiakerdxwusmpzam...uocrnugqwdblkjwigfjgettkovccqwcd
```

3 Calculating sequence background

Sequence backgrounds can be retrieved for DNA and RNA sequences with oligonucleotideFrequency() from "Biostrings. Unfortunately, no such Biostrings function exists for other sequence alphabets. The universalmotif package proves get_bkg() to remedy this. Similarly, the get_bkg() function can calculate higher order backgrounds for any alphabet as well. It is recommended to use the original Biostrings for very long DNA and RNA sequences whenever possible though, as it is much faster than get_bkg().

```
library(universalmotif)
## Background of DNA sequences:
dna <- create sequences()</pre>
get bkg(dna, k = 1:2, list.out = FALSE)
                       C
                                  G
                                              T
                                                                   AC
           Α
#> 0.24580000 0.25300000 0.24880000 0.25240000 0.06191919 0.06272727 0.06121212
                   CA
                                 CC
                                             CG
                                                        CT
         AT
                                                                   GA
#> 0.05979798 0.06373737 0.06171717 0.06303030 0.06464646 0.05979798 0.06636364
           GG
                      GT
                                 TA
                                             TC
                                                        TG
                                                                   TT
#> 0.06242424 0.06010101 0.06020202 0.06212121 0.06222222 0.06797980
## Background of non DNA/RNA sequences:
qwerty <- create_sequences("QWERTY")</pre>
get_bkg(qwerty, k = 1:2, list.out = FALSE)
                                              T
                                                                    Y
           E
                       Q
                                  R
#> 0.16530000 0.16790000 0.16210000 0.16740000 0.16700000 0.17030000 0.02363636
           EQ
                      ER
                                 ET
                                                        EY
                                                                   QE
                                            EW
#> 0.02616162 0.02686869 0.02929293 0.02666667 0.03262626 0.02727273 0.03040404
                                 QW
                                             QY
#>
           QR
                      QT
                                                        RE
                                                                   RQ
#> 0.02898990 0.02727273 0.02676768 0.02717172 0.02909091 0.02686869 0.02656566
           RT
                      RW
                                 RY
                                             TE
                                                        TQ
#> 0.02787879 0.02565657 0.02595960 0.02676768 0.02888889 0.02868687 0.02525253
           TW
                      TY
                                 WE
                                             WQ
                                                        WR
                                                                   WT
#> 0.02979798 0.02808081 0.02838384 0.02878788 0.02646465 0.02767677 0.02848485
                                                        YT
#> 0.02717172 0.03030303 0.02707071 0.02464646 0.03030303 0.02919192 0.02888889
```

4 Shuffling sequences

When performing *de novo* motif searches or motif enrichment analyses, it is common to do so against a set of background sequences. In order to properly identify consistent patterns or motifs in the target sequences, it is important that there be maintained a certain level of sequence composition between the target and background sequences. This reduces results which are derived purely from differential letter frequency biases.

In order to avoid these results, typically it desirable to use a set of background sequences which preserve a certain k-let size (such as dinucleotide or trinucleotide frequencies in the case of DNA sequences). Though for some cases a set of similar sequences may already be available for use as background sequences, usually background sequences are obtained by shuffling the target sequences, while preserving a desired k-let size. For this purpose, a commonly used tool is uShuffle (Jiang et al. 2008). The universalmotif package aims to provide its own k-let shuffling capabilities for use within R via shuffle_sequences().

The universalmotif package offers three different methods for sequence shuffling: euler, markov and linear. The first method, euler, can shuffle sequences while preserving any desired k-let size. Furthermore 1-letter counts will always be maintained. However in order for this to be possible, the first and last letters will remain unshuffled. This method is based on the initial random Eulerian walk algorithm proposed by Altschul and Erickson (1985) and the subsequent cycle-popping algorithm detailed by Propp and Wilson (1998) for quickly and efficiently finding Eulerian walks.

The second method, markov can only guarantee that the approximate k-let frequency will be maintained, but not that the original letter counts will be preserved. The markov method involves determining the original k-let frequencies, then creating a new set of sequences which will have approximately similar k-let frequency. As a result the counts for the individual letters will likely be different. Essentially, it involves a combination of determining k-let frequencies followed by create_sequences(). This type of shuffling is discussed by Fitch (1983).

The third method linear preserves the original 1-letter counts exactly, but uses a more crude shuffling technique. In this case the sequence is split into sub-sequences every k-let (of any size), which are then re-assembled randomly. This means that while shuffling the same sequence multiple times with method = "linear" will result in different sequences, they will all have started from the same set of k-length sub-sequences (just re-assembled differently).

```
library(universalmotif)
library(Biostrings)
data(ArabidopsisPromoters)

## Potentially starting off with some external sequences:
# ArabidopsisPromoters <- readDNAStringSet("ArabidopsisPromoters.fasta")

euler <- shuffle_sequences(ArabidopsisPromoters, k = 2, method = "euler")
markov <- shuffle_sequences(ArabidopsisPromoters, k = 2, method = "markov")
linear <- shuffle_sequences(ArabidopsisPromoters, k = 2, method = "linear")
k1 <- shuffle_sequences(ArabidopsisPromoters, k = 1)</pre>
```

Let us compare how the methods perform:

```
o.letter <- get_bkg(ArabidopsisPromoters, 1, as.prob = FALSE, list.out = FALSE)
e.letter <- get_bkg(euler, 1, as.prob = FALSE, list.out = FALSE)
m.letter <- get_bkg(markov, 1, as.prob = FALSE, list.out = FALSE)
l.letter <- get_bkg(linear, 1, as.prob = FALSE, list.out = FALSE)

data.frame(original=o.letter, euler=e.letter, markov=m.letter, linear=l.letter)
#> original euler markov linear
#> A 17384 17384 17451 17384
#> C 8081 8081 8121 8081
```

```
7583 7583
                      7616
                              7583
#> T
        16952 16952 16862
                           16952
o.counts <- get_bkg(ArabidopsisPromoters, 2, as.prob = FALSE, list.out = FALSE)
e.counts <- get_bkg(euler, 2, as.prob = FALSE, list.out = FALSE)</pre>
m.counts <- get_bkg(markov, 2, as.prob = FALSE, list.out = FALSE)</pre>
1.counts <- get_bkg(linear, 2, as.prob = FALSE, list.out = FALSE)</pre>
data.frame(original=o.counts, euler=e.counts, markov=m.counts, linear=1.counts)
      original euler markov linear
#> AA
          6893 6893
                       6151
                               6488
#> AC
          2614 2614
                       2781
                               2693
#> AG
          2592 2592
                       2627
                               2647
          5276 5276
#> AT
                       5874
                               5536
          3014 3014
#> CA
                       2791
                               2932
#> CC
          1376 1376
                       1364
                               1309
#> CG
          1051 1051
                       1211
                               1131
#> CT
          2621 2621
                       2747
                               2701
#> GA
          2734 2734
                       2705
                               2657
#> GC
         1104 1104
                       1210
                               1171
          1176 1176
#> GG
                       1186
                               1150
#> GT
          2561 2561
                       2508
                               2598
#> TA
          4725 4725
                       5787
                               5292
          2977 2977
#> TC
                       2757
                               2899
#> TG
          2759 2759
                       2585
                               2650
#> TT
          6477 6477
                       5716
                               6096
```

5 Miscellaneous string utilities

Since biological sequences are usually contained in XStringSet class objects, get_bkg() and shuffle_sequences() are designed to work with such objects. For cases when strings are not XStringSet objects, the following functions are available:

- count_klets(): alternative to get_bkg()
- shuffle_string(): alternative to shuffle_sequences()

```
library(universalmotif)
string <- "DASDSDDSASDSSA"
count_klets(string, 2)
     klets counts
#> 1
        AA
#> 2
        AD
                 0
#> 3
        AS
                 2
#> 4
        DA
                 1
#> 5
        DD
                1
                3
#> 6
        DS
#> 7
        SA
                 2
#> 8
        SD
                 3
#> 9
        SS
                 1
```

```
shuffle_string(string, 2)
#> [1] "DSDSSDASDDSASA"
```

Finally, the get_klets() function can be used to get a list of all possible k-lets for any sequence alphabet:

```
library(universalmotif)

get_klets(c("A", "S", "D"), 2)
#> [1] "AA" "AS" "AD" "SA" "SS" "SD" "DA" "DS" "DD"
```

6 Scanning sequences for motifs

There are many motif-programs available with sequence scanning capabilities, such as HOMER and tools from the MEME suite. The universalmotif package does not aim to supplant these, but rather provide convenience functions for quickly scanning a few sequences without needing to leave the R environment. Furthermore, these functions allow for taking advantage of the higher-order (multifreq) motif format described here.

Two scanning-related functions are provided: scan_sequences() and enrich_motifs(). The latter simply runs scan_sequences() twice on a set of target and background sequences. Given a motif of length n, scan_sequences() considers every possible n-length subset in a sequence and scores it using the PWM format. If the match surpasses the minimum threshold, it is reported. This is case regardless of whether one is scanning with a regular motif, or using the higher-order (multifreq) motif format (the multifreq matrix is converted to a PWM).

Before scanning a set of sequences, one must first decide the minimum logodds threshold for retrieving matches. This decision is not always the same between scanning programs out in the wild, nor is it usually told to the user what the cutoff is or how it is decided. As a result, universalmotif aims to be as transparent as possible in this regard by allowing for complete control of the threshold. For more details on PWMs, see the introductory vignette.

One way is to set a cutoff between 0 and 1, then multiplying the highest possible PWM score to get a threshold. The matchPWM() function from the Biostrings package for example uses a default of 0.8 (shown as "80%"). This is quite arbitrary of course, and every motif will end up with a different threshold. For high information content motifs, there is really no right or wrong threshold; as they tend to have fewer non-specific positions. This means that incorrect letters in a match will be more punishing. To illustrate this, contrast the following PWMs:

```
#> C -5.672425 -5.672425 -5.672425
#> G -5.672425 -5.672425 -5.672425
#> T -5.672425 1.978626 -5.672425
m2["motif"]
#>
                                     C
#> A -1.3219281 0.09667602 -0.12029423 -0.3959287 0.2141248
                                                             0.1491434
#> C 0.5260688
                0.19976951 1.02856915 0.6040713 -0.1202942 -0.6582115
#> G 0.8479969 -2.33628339 -3.64385619 -0.9434165 0.1110313 0.5897160
                0.66371661 -0.05889369 0.2630344 -0.2515388 -0.4102840
#> T -1.4739312
             R
                        N
                                   N
#> A 1.0430687 -1.0732490 0.4436067
                                     0.04222824
#> C -0.5418938 -0.2658941 -0.1202942 0.51171352
#> G 0.0710831 0.5897160 -1.0588937 0.29598483
#> T -2.3074285 0.2486791 0.3103401 -1.65821148
```

In the first example, sequences which do not have a matching base in every position are punished heavily. The maximum logodds score in this case is approximately 20, and for each incorrect position the score is reduced approximately by 5.7. This means that a threshold of zero would allow for at most three mismatches. At this point, it is up to you how many mismatches you would deem appropriate.

This thinking becomes impossible for the second example. In this case, mismatches are much less punishing, to the point that one could ask: what even constitutes a mismatch? The answer to this question is much more difficult in cases such as these. An alternative to manually deciding upon a threshold is to instead start with maximum P-value one would consider appropriate for a match. If, say, we want matches with a P-value of at most 0.001, then we can use motif_pvalue() to calculate the appropriate threshold (see the comparisons and P-values vignette for details on motif P-values).

```
motif_pvalue(m2, pvalue = 0.001)
#> [1] 4.8779
```

Furthermore, the scan_sequences() function offers the ability to scan using the multifreq slot, if available. This allows to take into account inter-positional dependencies, and get matches which more faithfully represent the original sequences from which the motif originated.

```
library(universalmotif)
library(Biostrings)
data(ArabidopsisPromoters)

## A 2-letter example:

motif.k2 <- create_motif("CWWWWCC", nsites = 6)
sequences.k2 <- DNAStringSet(rep(c("CAAAACC", "CTTTTCC"), 3))
motif.k2 <- add_multifreq(motif.k2, sequences.k2)</pre>
```

Regular scanning:

```
head(scan_sequences(motif.k2, ArabidopsisPromoters, RC = TRUE,
                    threshold = 0.9, threshold.type = "logodds"))
#> Note: detected non-finite positional weights, normalising
#> Note: since this motif has a pseudocount of 0, 1 will be used
#> DataFrame with 6 rows and 12 columns
#>
           motif
                   motif.i
                              sequence
                                            start
                                                       stop
                                                                            match
                                                                score
     <character> <integer> <character> <integer> <numeric> <character>
#>
#> 1
           motif
                             AT4G28150
                                              621
                                                        627
                                                                 9.08
                                                                          CTAAACC
                         1
#> 2
                             AT1G19380
                                              139
                                                        145
                                                                 9.08
                                                                          CTTATCC
           motif
                         1
           motif
                             AT1G19380
                                             204
                                                        210
                                                                 9.08
                                                                          CTAAACC
```

```
#> 4
                               AT1G03850
                                                 203
                                                           209
                                                                     9.08
                                                                               CTAATCC
            motif
#> 5
                                                           827
                                                                     9.08
            motif
                           1
                               AT5G01810
                                                 821
                                                                               CATATCC
#> 6
                                                                     9.08
            motif
                           1
                               AT5G01810
                                                 840
                                                           846
                                                                               CAAATCC
#>
     thresh.score min.score max.score score.pct
                                                         strand
#>
         <numeric> <numeric> <numeric> <numeric> <character>
#> 1
             8.172
                      -19.649
                                    9.08
                                                100
#> 2
             8.172
                      -19.649
                                    9.08
                                               100
#> 3
             8.172
                      -19.649
                                    9.08
                                                100
#> 4
             8.172
                      -19.649
                                    9.08
                                               100
#> 5
             8.172
                      -19.649
                                    9.08
                                                100
#> 6
             8.172
                      -19.649
                                    9.08
                                                100
```

Using 2-letter information to scan:

```
head(scan_sequences(motif.k2, ArabidopsisPromoters, use.freq = 2, RC = TRUE,
                     threshold = 0.9, threshold.type = "logodds"))
#> Note: detected non-finite positional weights, normalising
#> Note: since this motif has a pseudocount of 0, 1 will be used
#> DataFrame with 6 rows and 12 columns
#>
           motif
                    motif.i
                               sequence
                                             start
                                                         stop
                                                                  score
                                                                               match
     <character> <integer> <character> <integer> <integer> <numeric> <character>
#>
#> 1
           motif
                              AT4G12690
                                               938
                                                          943
                                                                 17.827
                                                                              CAAAAC
                          1
#> 2
                              AT2G37950
                                               751
                                                          756
                                                                 17.827
                                                                              CAAAAC
           motif
                          1
#> 3
           motif
                          1
                              AT1G49840
                                               959
                                                          964
                                                                 17.827
                                                                              CTTTTC
#> 4
           motif
                          1
                              AT1G77210
                                               184
                                                          189
                                                                 17.827
                                                                              CAAAAC
#> 5
           motif
                          1
                              AT1G77210
                                               954
                                                          959
                                                                 17.827
                                                                              CAAAAC
#> 6
                          1
                              AT3G57640
                                               917
                                                          922
                                                                 17.827
                                                                              CTTTTC
           motif
#>
     thresh.score min.score max.score score.pct
                                                        strand
#>
        <numeric> <numeric> <numeric> <numeric> <character>
#> 1
          16.0443
                     -16.842
                                17.827
                                              100
#> 2
          16.0443
                     -16.842
                                17.827
                                              100
#> 3
          16.0443
                     -16.842
                                17.827
                                              100
#> 4
          16.0443
                     -16.842
                                 17.827
                                              100
#> 5
          16.0443
                     -16.842
                                 17.827
                                              100
          16.0443
                     -16.842
                                 17.827
                                              100
```

As an aside: the previous example involved calling create_motif() and add_multifreq() separately. In this case however this could have been simplified to just calling create_motif() and using the add.multifreq option:

```
library(universalmotif)
library(Biostrings)

sequences <- DNAStringSet(rep(c("CAAAACC", "CTTTTCC"), 3))
motif <- create_motif(sequences, add.multifreq = 2:3)</pre>
```

7 Enrichment analyses

The universalmotif package offers the ability to search for enriched motif sites in a set of sequences via enrich_motifs(). There is little complexity to this, as it simply runs scan_sequences() twice: once on a set of target sequences, and once on a set of background sequences. After which the results between the two sequences are collated and run through enrichment tests. The background sequences can be given explicitly, or else enrich_motifs() will create background sequences on its own by using shuffle_sequences() on

the target sequences.

Let us consider the following basic example:

```
library(universalmotif)
data(ArabidopsisMotif)
data(ArabidopsisPromoters)
enrich_motifs(ArabidopsisMotif, ArabidopsisPromoters, shuffle.k = 3,
              threshold = 0.001, RC = TRUE)
#> DataFrame with 1 row and 11 columns
#>
               motif motif.i target.hits target.seq.hits target.seq.count
#>
         <character> <integer>
                                 <integer>
                                                  <integer>
#> 1 YTTTYTTTTTYTTY
                             1
                                       681
                                                                          50
      bkq.hits bkq.seq.hits bkq.seq.count
                                                  Pval
                                                              Qval
                                                                          Enal.
                  <integer>
#>
     <integer>
                                <integer>
                                                         <numeric>
                                                                     <numeric>
                                             <numeric>
                         48
#> 1
                                        50 2.17634e-37 2.17634e-37 4.35268e-37
```

Here we can see that the motif is significantly enriched in the target sequences. The Pval was calculated by calling stats::fisher.test().

One final point: always keep in mind the threshold parameter, as this will ultimately decide the number of hits found. (A bad threshold can lead to a false negative.)

8 Testing for motif positional preferences in sequences

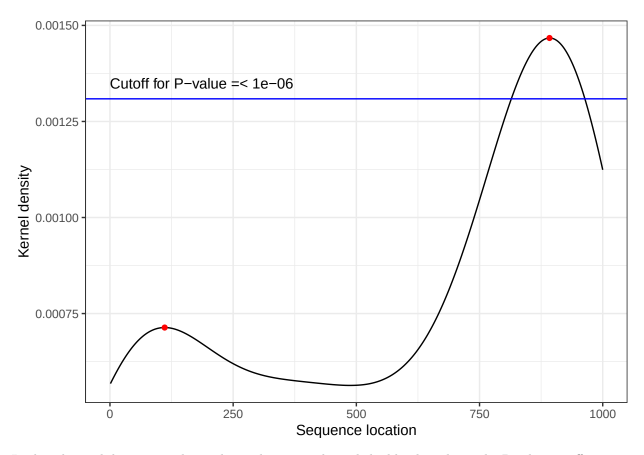
The universalmotif package provides the motif_peaks() function, which can test for positionally preferential motif sites in a set of sequences. This can be useful, for example, when trying to determine whether a certain transcription factor binding site is more often than not located at a certain distance from the transcription start site (TSS). The motif_peaks() function finds density peaks in the input data, then creates a null distribution from randomly generated peaks to calculate peak P-values.

```
library(universalmotif)
data(ArabidopsisMotif)
data(ArabidopsisPromoters)
hits <- scan_sequences(ArabidopsisMotif, ArabidopsisPromoters, RC = FALSE)
res <- motif_peaks(hits$start,
                   seq.length = unique(width(ArabidopsisPromoters)),
                   seq.count = length(ArabidopsisPromoters))
## Significant peaks:
res$Peaks
#> DataFrame with 1 row and 2 columns
                      Pval
          Peak
                 <numeric>
#>
     <numeric>
#> 1
         892 4.26875e-14
```

Using the datasets provided in this package, a significant motif peak was found about 100 bases away from the TSS. If you'd like to simply know the locations of any peaks, this can be done by setting max.p = 1.

The function can also output a plot:

```
res$Plot
```



In this plot, red dots are used to indicate density peaks and the blue line shows the P-value cutoff.

9 Motif discovery with MEME

The universalmotif package provides a simple wrapper to the powerful motif discovery tool MEME (Bailey and Elkan 1994). To run an analysis with MEME, all that is required is a set of XStringSet class sequences (defined in the Biostrings package), and run_meme() will take care of running the program and reading the output for use within R.

The first step is to check that R can find the MEME binary in your \$PATH by running run_meme() without any parameters. If successful, you should see the default MEME help message in your console. If not, then you'll need to provide the complete path to the MEME binary. There are two options:

```
library(universalmotif)

## 1. Once per session: via `options()`

options(meme.bin = "/path/to/meme/bin/meme")

run_meme(...)

## 2. Once per run: via `run_meme()`

run_meme(..., bin = "/path/to/meme/bin/meme")
```

Now we need to get some sequences to use with run_meme(). At this point we can read sequences from disk or extract them from one of the Bioconductor BSgenome packages.

```
library(universalmotif)
data(ArabidopsisPromoters)
## 1. Read sequences from disk (in fasta format):
library(Biostrings)
# The following `read*()` functions are available in Biostrings:
# DNA: readDNAStringSet
# DNA with quality scores: readQualityScaledDNAStringSet
# RNA: readRNAStringSet
# Amino acid: readAAStringSet
# Any: readBStringSet
sequences <- readDNAStringSet("/path/to/sequences.fasta")</pre>
run_meme(sequences, ...)
## 2. Extract from a `BSgenome` object:
library(GenomicFeatures)
library(TxDb.Athaliana.BioMart.plantsmart28)
library(BSgenome.Athaliana.TAIR.TAIR9)
# Let us retrieve the same promoter sequences from ArabidopsisPromoters:
gene.names <- names(ArabidopsisPromoters)</pre>
# First get the transcript coordinates from the relevant `TxDb` object:
transcripts <- transcriptsBy(TxDb.Athaliana.BioMart.plantsmart28,</pre>
                             by = "gene") [gene.names]
# There are multiple transcripts per gene, we only care for the first one
# in each:
transcripts <- lapply(transcripts, function(x) x[1])</pre>
transcripts <- unlist(GRangesList(transcripts))</pre>
# Then the actual sequences:
# Unfortunately this is a case where the chromosome names do not match
# between the two databases
seqlevels(TxDb.Athaliana.BioMart.plantsmart28)
#> [1] "1" "2" "3" "4" "5" "Mt" "Pt"
seqlevels(BSgenome.Athaliana.TAIR.TAIR9)
#> [1] "Chr1" "Chr2" "Chr3" "Chr4" "Chr5" "ChrM" "ChrC"
# So we must first rename the chromosomes in `transcripts`:
seqlevels(transcripts) <- seqlevels(BSgenome.Athaliana.TAIR.TAIR9)</pre>
# Finally we can extract the sequences
promoters <- getPromoterSeq(transcripts,</pre>
                             BSgenome.Athaliana.TAIR.TAIR9,
```

```
upstream = 1000, downstream = 0)
run_meme(promoters, ...)
```

Once the sequences are ready, there are few important options to keep in mind. One is whether to conserve the output from MEME. The default is not to, but this can be changed by setting the relevant option:

```
run_meme(sequences, output = "/path/to/desired/output/folder")
```

The second important option is the search function (objfun). Some search functions such as the default classic do not require a set of background sequences, whilst some do (such as de). If you choose one of the latter, then you can either let MEME create them for you (it will shuffle the target sequences) or you can provide them via the control.sequences parameter.

Finally, choose how you'd like the data imported into R. Once the MEME program exits, run_meme() will import the results into R with read_meme(). At this point you can decide if you want just the motifs themselves (readsites = FALSE) or if you'd like the original sequence sites as well (readsites = TRUE, the default). Doing the latter gives you the option of generating higher order representations for the imported MEME motifs as shown here:

```
motifs <- run_meme(sequences)
motifs.k23 <- mapply(add_multifreq, motifs$motifs, motifs$sites)</pre>
```

There are a wealth of other MEME options available, such as the number of desired motifs (nmotifs), the width of desired motifs (minw, maxw), the search mode (mod), assigning sequence weights (weights), using a custom alphabet (alph), and many others. See the output from run_meme() for a brief description of the options, or visit the online manual for more details.

Session info

```
#> R version 4.0.2 (2020-06-22)
#> Platform: x86_64-pc-linux-gnu (64-bit)
#> Running under: Ubuntu 18.04.4 LTS
#>
#> Matrix products: default
           /home/biocbuild/bbs-3.11-bioc/R/lib/libRblas.so
#> BLAS:
#> LAPACK: /home/biocbuild/bbs-3.11-bioc/R/lib/libRlapack.so
#>
#> locale:
    [1] LC CTYPE=en US.UTF-8
                                   LC NUMERIC=C
#>
    [3] LC_TIME=en_US.UTF-8
                                   LC_COLLATE=C
#>
   [5] LC MONETARY=en US.UTF-8
                                   LC MESSAGES=en US.UTF-8
#>
   [7] LC_PAPER=en_US.UTF-8
                                   LC_NAME=C
#>
    [9] LC ADDRESS=C
                                   LC TELEPHONE=C
#> [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
#>
#> attached base packages:
#> [1] stats4
                 parallel stats
                                     graphics grDevices utils
                                                                    datasets
#> [8] methods
                 base
#>
#> other attached packages:
#>
   [1] TFBSTools_1.26.0
                             Logolas_1.12.0
                                                   dplyr_1.0.2
#>
   [4] ggtree_2.2.4
                             ggplot2_3.3.2
                                                   MotifDb_1.30.0
  [7] GenomicRanges 1.40.0 GenomeInfoDb 1.24.2 Biostrings 2.56.0
```

```
#> [10] XVector 0.28.0
                             IRanges 2.22.2
                                                  S4Vectors_0.26.1
#> [13] BiocGenerics_0.34.0 universalmotif_1.6.4
#> loaded via a namespace (and not attached):
#>
     [1] colorspace_1.4-1
                                     grImport2_0.2-0
#>
     [3] ellipsis 0.3.1
                                     base64enc 0.1-3
                                     rGADEM 2.36.0
#>
     [5] aplot_0.0.6
                                     bit64 4.0.5
#>
     [7] farver_2.0.3
#>
     [9] AnnotationDbi_1.50.3
                                     R.methodsS3_1.8.1
#>
    [11] motifStack_1.32.1
                                     knitr_1.29
  [13] ade4_1.7-15
                                     jsonlite_1.7.1
#>
   [15] splitstackshape_1.4.8
                                     Rsamtools_2.4.0
#>
  [17] seqLogo_1.54.3
                                     gridBase_0.4-7
#> [19] annotate_1.66.0
                                     GO.db_3.11.4
#> [21] png_0.1-7
                                     R.oo_1.24.0
#>
   [23] BiocManager_1.30.10
                                     readr_1.3.1
#> [25] compiler_4.0.2
                                     httr_1.4.2
#> [27] rvcheck 0.1.8
                                     Matrix_1.2-18
#> [29] lazyeval_0.2.2
                                     htmltools_0.5.0
   [31] tools_4.0.2
                                     gtable_0.3.0
#> [33] glue_1.4.2
                                     TFMPvalue_0.0.8
#> [35] GenomeInfoDbData_1.2.3
                                     reshape2_1.4.4
#> [37] tinytex_0.25
                                     Rcpp_1.0.5
#> [39] Biobase_2.48.0
                                     vctrs 0.3.4
#> [41] ape_5.4-1
                                     nlme_3.1-149
#> [43] rtracklayer_1.48.0
                                     ggseqlogo_0.1
#> [45] gbRd_0.4-11
                                     xfun_0.17
#> [47] CNEr_1.24.0
                                     stringr_1.4.0
#> [49] ps_1.3.4
                                     lifecycle_0.2.0
#> [51] poweRlaw_0.70.6
                                     gtools_3.8.2
#> [53] XML_3.99-0.5
                                     zlibbioc_1.34.0
#> [55] MASS_7.3-53
                                     scales_1.1.1
#> [57] BSgenome_1.56.0
                                     hms_0.5.3
#> [59] SummarizedExperiment_1.18.2 RColorBrewer_1.1-2
#>
   [61] vaml 2.2.1
                                     memoise_1.1.0
#> [63] MotIV_1.44.0
                                     stringi_1.5.3
#> [65] RSQLite 2.2.0
                                     SQUAREM_2020.4
#> [67] highr_0.8
                                     tidytree_0.3.3
                                     BiocParallel_1.22.0
#> [69] caTools_1.18.0
#> [71] bibtex_0.4.2.2
                                     Rdpack_1.0.0
#> [73] rlang_0.4.7
                                     pkgconfig_2.0.3
#> [75] matrixStats_0.56.0
                                     bitops_1.0-6
#> [77] pracma_2.2.9
                                     evaluate_0.14
#> [79] lattice_0.20-41
                                     purrr_0.3.4
#> [81] htmlwidgets_1.5.1
                                     GenomicAlignments_1.24.0
#> [83] treeio_1.12.0
                                     patchwork_1.0.1
#> [85] labeling_0.3
                                     bit_4.0.4
#> [87] processx_3.4.4
                                     tidyselect_1.1.0
#> [89] plyr_1.8.6
                                     magrittr_1.5
#>
   [91] bookdown_0.20
                                     R6_2.4.1
#> [93] generics_0.0.2
                                     DelayedArray_0.14.1
#> [95] DBI_1.1.0
                                     pillar_1.4.6
#> [97] withr_2.2.0
                                     KEGGREST_1.28.0
#> [99] RCurl_1.98-1.2
                                     tibble_3.0.3
```

References

Altschul, Stephen F., and Bruce W. Erickson. 1985. "Significance of Nucleotide Sequence Alignments: A Method for Random Sequence Permutation That Preserves Dinucleotide and Codon Usage." *Molecular Biology and Evolution* 2 (6):526–38.

Bailey, T.L., and C. Elkan. 1994. "Fitting a Mixture Model by Expectation Maximization to Discover Motifs in Biopolymers." *Proceedings of the Second International Conference on Intelligent Systems for Molecular Biology* 2:28–36.

Fitch, Walter M. 1983. "Random Sequences." Journal of Molecular Biology 163 (2):171-76.

Jiang, M., J. Anderson, J. Gillespie, and M. Mayne. 2008. "uShuffle: A Useful Tool for Shuffling Biological Sequences While Preserving K-Let Counts." *BMC Bioinformatics* 9 (192).

Propp, J.G., and D.W. Wilson. 1998. "How to Get a Perfectly Random Sample from a Generic Markov Chain and Generate a Random Spanning Tree of a Directed Graph." *Journal of Algorithms* 27:170–217.