

Upsize your clustering with Clusterize

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1 Introduction to supersized clustering

You may have found yourself in a familiar predicament for many bioinformaticians: you have a lot of sequences and you need to *downsize* before you can get going. You may also *theorize* that this must be an easy problem to solve—given sequences, output clusters. But what can you *utilize* to solve this problem? This vignette will *familiarize* you with the `Clusterize` function in the DECIPHER package. Clusterize will *revolutionize* all your clustering needs! Why `Clusterize`?:

- Scalability - `Clusterize` will *linearize* the search space so that many sequences can be clustered in a reasonable amount of time.
- Simplicity - Although you can *individualize* `Clusterize`, the defaults are straightforward and should meet most of your needs.
- Accuracy - `Clusterize` will *maximize* your ability to extract biologically meaningful results from your sequences.

This vignette will *summarize* the use of `Clusterize` to cluster DNA, RNA, or protein sequences.

2 Getting started with Clusterize

To get started we need to load the DECIPHER package, which automatically mobilize a few other required packages.

```
> library(DECIPHER)
```

There's no need to memorize the inputs to Clusterize, because its help page can be accessed through:

```
> ? Clusterize
```

Note that, while it's easy to fantasize about using Clusterize, if you only have a moderate number of **homologous** sequences ($\ll 100k$) then it's more accurate to use Treeline with a distance matrix created from a multiple sequence alignment. This function provides hierarchical clustering (i.e., single-linkage, UPGMA, or complete-linkage) that is impossible to criticize as inexact.

3 Optimize your inputs to Clusterize

Clusterize requires that you first digitize your sequences by loading them into memory. For the purpose of this vignette, we will capitalize on the fact that DECIPHER already includes some built-in sets of sequences.

```
> # specify the path to your file of sequences:
> fas <- "<<path to training FASTA file>>"
> # OR use the example DNA sequences:
> fas <- system.file("extdata",
  "50S_ribosomal_protein_L2.fas",
  package="DECIPHER")
> # read the sequences into memory
> dna <- readDNASTringSet(fas)
> dna
DNASTringSet object of length 317:
      width seq
[1] 819 ATGGCTTTAAAAATTTTAATC...ATTTATTGTAAAAAAGAAAA Rickettsia prowaz...
[2] 822 ATGGGAATACGTAAACTCAAGC...CATCATTGAGAGAAGGAAAAAG Porphyromonas gin...
[3] 822 ATGGGAATACGTAAACTCAAGC...CATCATTGAGAGAAGGAAAAAG Porphyromonas gin...
[4] 822 ATGGGAATACGTAAACTCAAGC...CATCATTGAGAGAAGGAAAAAG Porphyromonas gin...
[5] 819 ATGGCTATCGTTAAATGTAAGC...CATCGTACGTCGTCGTGGTAAA Pasteurella multo...
...
[313] 819 ATGGCAATTGTTAAATGTAAAC...TATCGTACGTCGCCGTAATAA Pectobacterium at...
[314] 822 ATGCCTATTCAAAAATGCAAAC...TATTCGCGATCGTCGCGTCAAG Acinetobacter sp....
[315] 864 ATGGGCATTTCGCGTTTACCGAC...GGGTCGCGGTGGTCGTCAGTCT Thermosynechococc...
[316] 831 ATGGCACTGAAGACATTCAATC...AAGCCGCCACAAGCGGAAGAAG Bradyrhizobium ja...
[317] 840 ATGGGCATTTCGCAAATATCGAC...CAAGACGGCTTCGGGCGAGGT Gloeobacter viola...
```

The Clusterize algorithm will generalize to nucleotide or protein sequences, so we must choose which we are going to use. Here, we hypothesize that weaker similarities can be detected between proteins and, therefore, decide to use the translated coding (amino acid) sequences. If you wish to cluster at high similarity, you could also strategize that nucleotide sequences would be better because there would be more nucleotide than amino acid differences.

```
> aa <- translate(dna)
> aa
```

```
AAStringSet object of length 317:
```

```

      width seq
[1] 273 MALKNFPITPSLRELVQVDKT...STKGKKTRKNKRTSKFIVKKRK Rickettsia prowaz...
[2] 274 MGIRKLKPTTPGQRHKVIGAFD...KGLKTRAPKKHSSKYIIERRKK Porphyromonas gin...
[3] 274 MGIRKLKPTTPGQRHKVIGAFD...KGLKTRAPKKHSSKYIIERRKK Porphyromonas gin...
[4] 274 MGIRKLKPTTPGQRHKVIGAFD...KGLKTRAPKKHSSKYIIERRKK Porphyromonas gin...
[5] 273 MAIVKCKPTSAGRRHVVKIVNP...TKGKKTRHNKRTDKFIVRRRGK Pasteurella multo...
...
[313] 273 MAIVKCKPTSPGRRHVVKVNP...TKGKKTRSNKRTDKFIVRRRTK Pectobacterium at...
[314] 274 MPIQKCKPTSPGRRFVEKVVHS...KGYKTRTNKRTTKMIIRDRVK Acinetobacter sp....
[315] 288 MGIRVYRPYTPGVRQKTVSDFA...SDALIVRRRKSSKRGGRGGRQS Thermosynechococc...
[316] 277 MALKTFNPTTPGQRQLVMVDRS...KKTRSNKSTNKFILLSRHKRKK Bradyrhizobium ja...
[317] 280 MGIRKYRPMTPGTRQSGADFA...RKRRKPSSKFIIRRRKTASGRG Gloeobacter viola...
> seqs <- aa # could also cluster the nucleotides
> length(seqs)
[1] 317

```

Now you can choose how to parameterize the function, with the main arguments being *myXStringSet* and *cutoff*. In this case, we will initialize *cutoff* at `seq(0.5, 0, -0.1)` to cluster sequences from 50% to 100% similarity by 10%'s. It is important to recognize that *cutoffs* can be provided in *ascending* or *descending* order and, when *descending*, groups at each *cutoff* will be nested within the previous *cutoff*'s groups.

We must also choose whether to customize the calculation of distance. The defaults will penalize gaps as single events, such that each consecutive set of gaps (i.e., insertion or deletion) is considered equivalent to one mismatch. If you want to standardize the definition of distance to be the same as most other clustering programs then set: *penalizeGapLetterMatches* to TRUE (i.e., every gap position is a mismatch), *method* to "shortest", *minCoverage* to 0, and *includeTerminalGaps* to TRUE. It is possible to rationalize many different measures of distance – see the *DistanceMatrix* function for more information about alternative distance parameterizations.

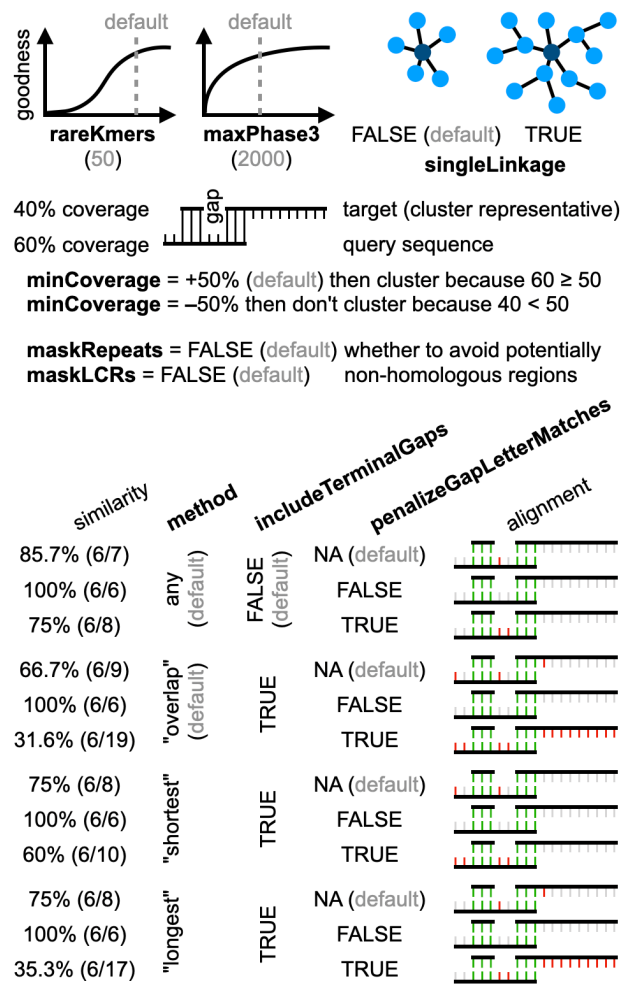


Figure 1: The most important parameters (in **bold**) to customize your use of Clusterize.

We can further *personalize* the inputs as desired. The main function argument to *emphasize* is *processors*, which controls whether the function is parallelized on multiple computer threads (if DECIPHER was built with OpenMP enabled). Setting *processors* to a value greater than 1 will speed up clustering considerably, especially for large size clustering problems. Once we are ready, it's time to run `Clusterize` and wait for the output to *materialize*!

```
> clusters <- Clusterize(seqs, cutoff=seq(0.5, 0, -0.1), processors=1)
Partitioning sequences by 3-mer similarity:
=====

Time difference of 0.04 secs

Sorting by relatedness within 35 groups:

iteration 34 of up to 34 (100.0% stability)

Time difference of 0.57 secs

Clustering sequences by 5-mer similarity:
=====

Time difference of 0.15 secs

Clusters via relatedness sorting: 100% (0% exclusively)
Clusters via rare 3-mers: 100% (0% exclusively)
Estimated clustering effectiveness: 100%
> class(clusters)
[1] "data.frame"
> colnames(clusters)
[1] "cluster_0_5" "cluster_0_4" "cluster_0_3" "cluster_0_2" "cluster_0_1"
[6] "cluster_0"
> str(clusters)
'data.frame':      317 obs. of  6 variables:
 $ cluster_0_5: int  3 1 1 1 3 3 3 2 2 2 ...
 $ cluster_0_4: int  1 21 21 21 3 3 3 10 10 10 ...
 $ cluster_0_3: int  42 1 1 1 35 35 36 23 23 23 ...
 $ cluster_0_2: int  1 67 67 67 12 12 9 34 34 34 ...
 $ cluster_0_1: int  86 1 11 69 69 73 41 41 41 ...
 $ cluster_0   : int  2 102 102 102 25 25 20 59 59 59 ...
> apply(clusters, 2, max) # number of clusters per cutoff
cluster_0_5 cluster_0_4 cluster_0_3 cluster_0_2 cluster_0_1 cluster_0
          3          21          42          67          86         102
> apply(clusters, 2, function(x) which.max(table(x))) # max sizes
cluster_0_5 cluster_0_4 cluster_0_3 cluster_0_2 cluster_0_1 cluster_0
          3           5          30          22          54          45
```

Notice that `Clusterize` will *characterize* the clustering based on how many clustered pairs came from relatedness sorting versus rare k-mers, and `Clusterize` will predict the effectiveness of clustering. Depending on the input sequences, the percentage of clusters originating from relatedness sorting will *equalize* with the number originating from rare k-mers, but more commonly clusters will originate from one source or the other. The clustering effectiveness formalizes the concept of “inexact” clustering by approximating the fraction of possible sequence pairs

that were correctly clustered together. You can incentivize a higher clustering effectiveness by increasing *maxPhase3* at the expense of (proportionally) longer run times.

We can now realize our objective of decreasing the number of sequences. Here, we will prioritize keeping only the longest diverse sequences.

```
> o <- order(clusters[[2]], width(seqs), decreasing=TRUE) # 40% cutoff
> o <- o[!duplicated(clusters[[2]])]
> aa[o]
AAStringSet object of length 21:
      width seq                                     names
[1]    274 MGIRKLKPTTPGQRHKVIGAFDK...KGLKTRAPKKHSSKYIIERRKK Porphyromonas gin...
[2]    274 MGIRKLKPTTPGQRHKVIGAFDK...KGLKTRAPKKHSSKYIIERRKK Porphyromonas gin...
[3]    274 MAVRKLKPTTPGQRHKIIGTFEE...KGLKTRAPKKQSSKYIIERRKK Bacteroides theta...
[4]    277 MGIKTYKPKTSSLRYKTTLSFDD...KGYKTRKKKRYSDKFIKRRNK Borrelia burgdorf...
[5]    280 MAIRKYKPTTPGRRQSSVSMFEE...NPNRYSNNMIVQRRRTNKSCKR Corynebacterium d...
...    ...
[17]   273 MAIVKCKPTSAGRRHVVKIVNPE...TKGKKTRHNKRTDKYIVRRRGK Haemophilus influ...
[18]   273 MAIVKCKPTSAGRRHVVKIVNPE...TKGKKTRHNKRTDKYIVRRRGK Haemophilus influ...
[19]   273 MAIVKCKPTSAGRRFVVKVQNQE...QTGKKTRSNKRTDNMIVRRRK Pseudomonas aerug...
[20]   277 MALKHFNPIPTGQRQLVIVDRSE...KKTRSNKATDKFIMRSRHQRKK Brucella suis VBI22
[21]   274 MAIVKCKPTSAGRRHVVKVNNAD...TKGYKTRSNKRTDKYIVRRRNK Vibrio cholerae PS15
> dna[o]
DNAStrngSet object of length 21:
      width seq                                     names
[1]    822 ATGGGAATACGTAAACTCAAGCC...CATCATTGAGAGAAGGAAAAAG Porphyromonas gin...
[2]    822 ATGGGAATACGTAAACTCAAGCC...CATCATTGAGAGAAGGAAAAAG Porphyromonas gin...
[3]    822 ATGGCAGTACGTAAATTAAGCC...CATTATTGAGAGAAGAAAAAG Bacteroides theta...
[4]    831 ATGGGTATTAAGACTTATAAGCC...TATTATTAAGAAGAATAAA Borrelia burgdorf...
[5]    840 ATGGCTATTCGTAAGTACAAGCC...CACGAACAAGAGCAAGAAGCGC Corynebacterium d...
...    ...
[17]   819 ATGGCTATCGTTAAATGTAAGCC...TATCGTACGTCGTCGTGGCAAA Haemophilus influ...
[18]   819 ATGGCTATCGTTAAATGTAAGCC...TATCGTACGTCGTCGTGGCAAA Haemophilus influ...
[19]   819 ATGGCAATCGTTAAGTGCAAAACC...CATGATCGTCGCCGCCGCAAG Pseudomonas aerug...
[20]   831 ATGGCACTCAAGCATTTTAATCC...TTCGCGCCATCAGCGCAAGAAG Brucella suis VBI22
[21]   822 ATGGCTATTGTTAAATGTAAGCC...CATCGTACGTCGTCGTAATAAG Vibrio cholerae PS15
```

4 Visualize the output of Clusterize

We can scrutinize the clusters by selecting them and looking at their multiple sequence alignment:

```
> t <- table(clusters[[1]]) # select the clusters at a cutoff
> t <- sort(t, decreasing=TRUE)
> head(t)
  3    1    2
218  58  41
> w <- which(clusters[[1]] == names(t[1]))
> AlignSeqs(seqs[w], verbose=FALSE)
AAStringSet object of length 218:
      width seq                                     names
```

```

[1] 288 -MALKNFPITPSLRELVQVDK...TR-KNKRTSKFIVKKRK----- Rickettsia prowaz...
[2] 288 -MAIVKCKPTSAGRRHVVKIVN...TR-HNKRTDKFIVRRRGK---- Pasteurella multo...
[3] 288 -MAIVKCKPTSAGRRHVVKIVN...TR-HNKRTDKFIVRRRGK---- Pasteurella multo...
[4] 288 -MPLMKFKPTSPGRRSAVRVVT...TR-KNKRTQQFIVRDRRG---- Xanthomonas campe...
[5] 288 -MPLMKFKPTSPGRRSAVRVVT...TR-KNKRTQQFIVRDRRG---- Xanthomonas citri...
...
[214] 288 -MAFKHFNPTTPGQRQLVIVDR...TR-SNKATDKFIMHTRHQRKK- Bartonella quinta...
[215] 288 -MAFKHFNPTTPGQRQLVIVDR...TR-SNKATDKFIMHTRHQRKK- Bartonella quinta...
[216] 288 -MAIVKCKPTSPGRRHVVKVNV...TR-SNKRTDKFIVRRRTK---- Pectobacterium at...
[217] 288 -MPIQKCKPTSPGRRFVEKVH...TR-TNKRTTKMIIRDRRVK--- Acinetobacter sp....
[218] 288 -MALKTFNPTTPGQRQLVMVDR...TR-SNKSTNKFILLSRHKRKK- Bradyrhizobium ja...

```

It's possible to utilize the `heatmap` function to view the clustering results.

As can be seen in Figure 2, `Clusterize` will organize its clusters such that each new cluster is within the previous cluster when *cutoff* is provided in descending order. We can also see that sequences from the same species tend to cluster together, which is an alternative way to systematize sequences without clustering.

```

> aligned_seqs <- AlignSeqs(seqs, verbose=FALSE)
> d <- DistanceMatrix(aligned_seqs, verbose=FALSE)
> tree <- Treeline(myDistMatrix=d, method="UPGMA", verbose=FALSE)
> heatmap(as.matrix(clusters), scale="column", Colv=NA, Rowv=tree)

```

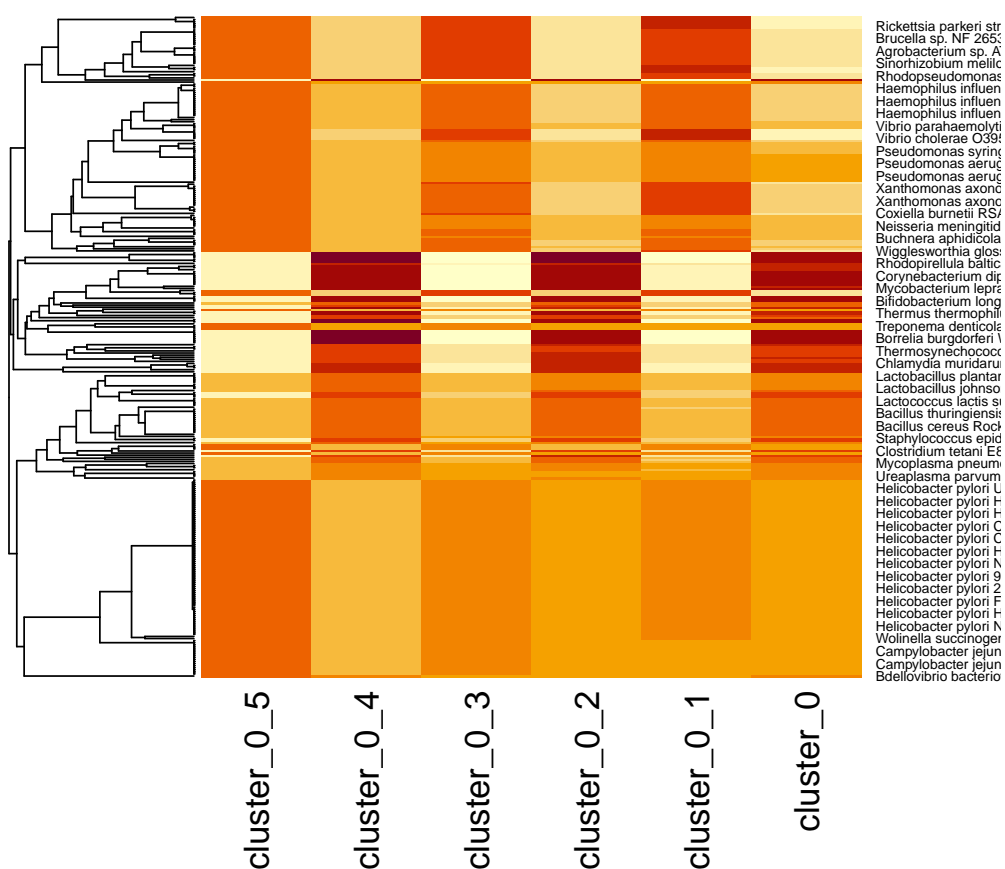


Figure 2: Visualization of the clustering.

5 Specialize clustering for your goals

The most common use of clustering is to categorize sequences into groups sharing similarity above a threshold and pick one representative sequence per group. These settings *empitomize* this typical user scenario:

```
> c1 <- Clusterize(dna, cutoff=0.2, invertCenters=TRUE, processors=1)
Partitioning sequences by 5-mer similarity:
=====

Time difference of 0.15 secs

Sorting by relatedness within 34 groups:

iteration 25 of up to 56 (100.0% stability)

Time difference of 2.4 secs

Clustering sequences by 10-mer similarity:
=====

Time difference of 0.4 secs

Clusters via relatedness sorting: 100% (0% exclusively)
Clusters via rare 5-mers: 100% (0% exclusively)
Estimated clustering effectiveness: 100%
> w <- which(c1 < 0 & !duplicated(c1))
> dna[w] # select cluster representatives (negative cluster numbers)
DNAStrngSet object of length 78:
      width seq                                     names
[1]   819 ATGGCTTTAAAAAATTTTAATCC...ATTTATTGTAAAAAAGAAAA Rickettsia prowaz...
[2]   822 ATGGGAATACGTAAACTCAAGCC...CATCATTGAGAGAAGGAAAAAG Porphyromonas gin...
[3]   837 GTGGGTATTAAGAAGTATAAACC...TGGTCGCCGTCCAGGCAAACAC Lactobacillus pla...
[4]   825 ATGCCATTGATGAAGTTCAAACC...CATCGTCCGCGATCGTAGGGGC Xanthomonas axono...
[5]   828 ATGGGTATTCGTAATTATCGGCC...GATTGTCCGCGTCGCACCAAA Synechocystis sp....
...   ...
[74]  831 ATGGCATTTAAGCACTTTAATCC...TACGCGTCATCAGCGCAAGAAA Bartonella quinta...
[75]  843 ATGTTTAAGAAATATCGACCTGT...CGTGAAACGTCGAAGGAAGAAG Candidatus Protoc...
[76]  822 ATGCCTATTCAAAAATGCAAACC...TATTCGCGATCGTCGCGTCAAG Acinetobacter sp....
[77]  864 ATGGGCATTTCGCGTTTACCGACC...GGGTCGCGGTGGTTCGTCAGTCT Thermosynechococc...
[78]  840 ATGGGCATTTCGCAAATATCGACC...CAAGACGGCTTCCGGGCGAGGT Gloeobacter viola...
```

By default, `Clusterize` will cluster sequences with linkage to the representative sequence in each group, but it is also possible to tell `Clusterize` to *minimize* the number of clusters by establishing linkage to any sequence in the cluster (i.e., single-linkage). This is often how we conceptualize natural groupings and, therefore, may better match alternative classification systems such as taxonomy:

```
> c2 <- Clusterize(dna, cutoff=0.2, singleLinkage=TRUE, processors=1)
Partitioning sequences by 5-mer similarity:
=====

Time difference of 0.11 secs
```

Sorting by relatedness within 34 groups:

iteration 22 of up to 56 (100.0% stability)

Time difference of 2.06 secs

Clustering sequences by 10-mer similarity:

=====

Time difference of 0.78 secs

Clusters via relatedness sorting: 100% (0% exclusively)

Clusters via rare 5-mers: 100% (0% exclusively)

Estimated clustering effectiveness: 100%

> max(abs(c1)) # center-linkage

[1] 78

> max(c2) # single-linkage (fewer clusters, but broader clusters)

[1] 76

It is possible to *synthesize* a plot showing a cross tabulation of taxonomy and cluster number. We may *idealize* the clustering as matching taxonomic labels (3), but this is not exactly the case.

```

> genus <- sapply(strsplit(names(dna), " "), `[`, 1)
> t <- table(genus, c2[[1]])
> heatmap(sqrt(t), scale="none", Rowv=NA, col=hcl.colors(100))

```



Figure 3: Another visualization of the clustering.

6 Resize to fit within less memory

What should you do if you have more sequences than you can cluster on your *midsize* computer? If there are far fewer clusters than sequences (e.g., *cutoff* is high) then it is likely possible to *resize* the clustering problem. This is accomplished by processing the sequences in batches that *miniaturize* the memory footprint and are at least as large as the final number of clusters. The number of sequences processed per batch is critical to *atomize* the problem appropriately while limiting redundant computations. Although not ideal from a speed perspective, the results will not *jeopardize* accuracy relative to as if there was sufficient memory available to process all sequences in one batch.

```
> batchSize <- 2e2 # normally a large number (e.g., 1e6 or 1e7)
> o <- order(width(seqs), decreasing=TRUE) # process largest to smallest
> c3 <- integer(length(seqs)) # cluster numbers
> repeat {
  m <- which(c3 < 0) # existing cluster representatives
  m <- m[!duplicated(c3[m])] # remove redundant sequences
  if (length(m) >= batchSize)
    stop("batchSize is too small")
  w <- head(c(m, o[c3[o] == 0L]), batchSize)
  if (!any(c3[w] == 0L)) {
    if (any(c3[-w] == 0L))
      stop("batchSize is too small")
    break # done
  }
  m <- m[match(abs(c3[-w]), abs(c3[m]))]
  c3[w] <- Clusterize(seqs[w], cutoff=0.05, invertCenters=TRUE)[[1]]
  c3[-w] <- ifelse(is.na(c3[m]), 0L, abs(c3[m]))
}
```

Partitioning sequences by 3-mer similarity:

=====

Time difference of 0.03 secs

Sorting by relatedness within 4 groups:

iteration 1 of up to 29 (100.0% stability)

Time difference of 0.01 secs

Clustering sequences by 5-mer similarity:

=====

Time difference of 0.12 secs

Clusters via relatedness sorting: 100% (0% exclusively)

Clusters via rare 3-mers: 100% (0% exclusively)

Estimated clustering effectiveness: 100%

Partitioning sequences by 3-mer similarity:

=====

Time difference of 0.03 secs

```
Sorting by relatedness within 97 groups:
Clustering sequences by 5-mer similarity:
```

```
=====
```

```
Time difference of 0.22 secs
```

```
Clusters via relatedness sorting: 100% (0% exclusively)
Clusters via rare 3-mers: 100% (0% exclusively)
Estimated clustering effectiveness: 100%
```

```
> table(abs(c3)) # cluster sizes
```

1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26
1	1	1	1	1	1	1	2	1	1	2	1	3	1	1	1	1	1	7	1	1	1	3	1	1	1
27	28	29	30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45	46	47	48	49	50	51	52
3	5	1	3	2	6	3	3	1	2	1	6	1	7	1	1	1	2	8	3	17	3	2	2	2	1
53	54	55	56	57	58	59	60	61	62	63	64	65	66	67	68	69	70	71	72	73	74	75	76	77	78
1	1	1	3	3	12	1	75	4	1	1	11	3	1	1	1	1	1	1	5	6	3	3	2	1	1
79	80	81	82	83	84	85	86	87	88	89	90	91													
1	1	1	17	13	6	3	1	1	1	1	1	1	1												

7 Clustering both nucleotide strands

Sometimes the input sequences are present in different orientations and it is necessary to harmonize the clusterings from both strands. Without trying to hyperbolize how easy this is to do, here's an example of clustering both strands:

```
> # simulate half of strands having opposite orientation
> s <- sample(c(TRUE, FALSE), length(dna), replace=TRUE)
> dna[s] <- reverseComplement(dna[s])
> # cluster both strands at the same time
> clus <- Clusterize(c(dna, reverseComplement(dna)), cutoff=0.2, processors=1)
Partitioning sequences by 5-mer similarity:
```

```
=====
```

```
Time difference of 0.28 secs
```

```
Sorting by relatedness within 142 groups:
```

```
iteration 28 of up to 50 (100.0% stability)
```

```
Time difference of 5.12 secs
```

```
Clustering sequences by 10-mer similarity:
```

```
=====
```

```
Time difference of 1.23 secs
```

```
Clusters via relatedness sorting: 100% (0% exclusively)
Clusters via rare 5-mers: 100% (0% exclusively)
Estimated clustering effectiveness: 100%
```

```

> clus <- match(clus[[1]], clus[[1]]) # renumber clusters ascending
> # if needed, reorient all clustered sequences to have the same orientation
> strand <- clus[seq_len(length(clus)/2)] >= clus[-seq_len(length(clus)/2)]
> dna[strand] <- reverseComplement(dna[strand])
> # renumber clusters across both strands and compare to original clustering
> clus <- pmin(clus[seq_len(length(clus)/2)], clus[-seq_len(length(clus)/2)])
> org <- match(abs(c1[[1]]), abs(c1[[1]])) # renumber original clustering
> mean(clus == org) # some differences expected due to algorithm stochasticity
[1] 0.9842271
> # verify the largest cluster is now back in the same orientation
> dna[clus == which.max(tabulate(clus))]
DNAStrngSet object of length 75:
      width seq
[1] 828 ATGGCGATTAAACTTATAAGCC...CATTTCCAGAAAGAAACACAAA Helicobacter pylo...
[2] 828 ATGGCGATTAAACTTATAAGCC...CATTTCCAGAAAGAAACATAAA Helicobacter pylo...
[3] 828 ATGGCGATTAAACTTATAAGCC...CATTTCCAGAAAGAAACATAAA Helicobacter pylo...
[4] 828 ATGGCGATTAAACTTATAAGCC...CATTTCCAGAAAGAAACATAAA Helicobacter pylo...
[5] 828 ATGGCGATTAAACTTATAAGCC...CATTTCCAGAAAGAAACATAAA Helicobacter pylo...
...
[71] 828 ATGGCGATTAAACTTATAAGCC...CATTTCCAGAAAGAAACATAAA Helicobacter pylo...
[72] 828 ATGGCGATTAAACTTATAAGCC...CATTTCCAGAAAGAAACATAAA Helicobacter pylo...
[73] 828 ATGGCGATTAAACTTATAAGCC...CATTTCCAGAAAGAAACATAAA Helicobacter pylo...
[74] 828 ATGGCGATTAAACTTATAAGCC...CATTTCCAGAAAGAAACATAAA Helicobacter pylo...
[75] 828 ATGGCGATTAAACTTATAAGCC...CATTTCCAGAAAGAAACACAAA Helicobacter pylo...

```

8 Finalize your use of Clusterize

Notably, Clusterize is a stochastic algorithm, meaning it will *randomize* which sequences are selected during pre-sorting. Even though the clusters will typically *stabilize* with enough iterations, you can set the random number seed (before every run) to guarantee reproducibility of the clusters:

```

> set.seed(123) # initialize the random number generator
> clusters <- Clusterize(seqs, cutoff=0.1, processors=1)
Partitioning sequences by 3-mer similarity:
=====

```

Time difference of 0.04 secs

Sorting by relatedness within 35 groups:

iteration 1 of up to 34 (100.0% stability)

Time difference of 0.02 secs

Clustering sequences by 5-mer similarity:

```

=====
Time difference of 0.2 secs

```

```

Clusters via relatedness sorting: 100% (0% exclusively)
Clusters via rare 3-mers: 100% (0% exclusively)
Estimated clustering effectiveness: 100%
> set.seed(NULL) # reset the seed

```

Now you know how to utilize `Clusterize` to cluster sequences. To publicize your results for others to reproduce, make sure to provide your random number seed and version number:

- R Under development (unstable) (2024-11-20 r87352), x86_64-apple-darwin20
- Running under: macOS Monterey 12.7.6
- Matrix products: default
- BLAS:
/Library/Frameworks/R.framework/Versions/4.5-x86_64/Resources/lib/libRblas.0.dylib
- LAPACK:
/Library/Frameworks/R.framework/Versions/4.5-x86_64/Resources/lib/libRlapack.dylib
; LAPACK version 3.12.0
- Base packages: base, datasets, grDevices, graphics, methods, stats, stats4, utils
- Other packages: BiocGenerics 0.53.3, Biostrings 2.75.3, DECIPHER 3.3.2, GenomeInfoDb 1.43.2, IRanges 2.41.2, S4Vectors 0.45.2, XVector 0.47.1, generics 0.1.3
- Loaded via a namespace (and not attached): DBI 1.2.3, GenomeInfoDbData 1.2.13, R6 2.5.1, UCSC.utils 1.3.0, compiler 4.5.0, crayon 1.5.3, httr 1.4.7, jsonlite 1.8.9, tools 4.5.0