# DiffLogo user guide

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#### 1 Introduction

The DiffLogo tool is a R package for the visualization of differences between multiple motifs for different alphabets. The user supplies a set of motifs each represented as position weight matrices (PWMs) [1]. The DiffLogo package supports the comparison of two motifs by a single difference logo and the comparison of multiple motifs by a table of difference logos. Difference logos are based on the idea behind the well-known sequence logo [2], i.e. motifs are visualized position-wise based on two functions. First, the stackHeight function computes the height of each stack. Second, the baseDistribution function breaks down the stack height on the individual characters. The user is able to parametrise the individual functions with arbitrary functions stackHeight and baseDistribution. Default implementations of both functions are provided.

# 2 Download and import library

After installing the package, the user is able to import DiffLogo.

> library(DiffLogo)

# 3 Import PWMs

PWMs can be represented as object of type pwm, data.frame, or matrix. The user is able to import motifs from any source in one of these formats.

Here, we import two motifs from the transcription factor CTCF from package MotifDb [3]. Alternatively, there are example PWMs in folder ext-data/pwm and extdata/alignments shipped with the package DiffLogo. (CTCF motifs extracted from [4], E-Box transcription factor binding sites extracted from [5], and F-Box protein domains extracted from [6]).

```
> ## import nine DNA motifs for transcription factor CTCF from matrix
> motif_folder <- "extdata/pwm"</pre>
> motif_names_dna = c(
    "GM12878", "H1-hESC", "HeLa-S3", "HepG2",
                                                    "HUVEC".
                "MCF7",
                            "NHEK",
    "K562",
                                        "ProgFib")
> motifs_dna = list()
> for (name in motif_names_dna) {
    fileName <- paste(motif_folder,"/",name,".txt",sep="")</pre>
    file <- system.file(fileName, package = "DiffLogo")
    motifs_dna[[name]] <- as.matrix(read.delim(file, FALSE))</pre>
+ }
> ## import DNA motifs for three transcription factors from table
> motif_folder <- "extdata/alignments"
> motif_names_dna2 <- c("Mad", "Max", "Myc")</pre>
> motifs_dna2 <- list()</pre>
> for (name in motif_names_dna2) {
    fileName <- paste(motif_folder,"/",name,".txt",sep="")</pre>
    file <- system.file(fileName, package = "DiffLogo")</pre>
    fileContent <- readLines(file)</pre>
    fileContent <- unlist(lapply(</pre>
      X = fileContent,
```

```
FUN = function(x) \{ strsplit(x = x, split = "\t")[[1]][[1]] \}))
    motifs_dna2[[name]] <- getPwmFromAlignment(fileContent, DNA, 1)</pre>
+ }
> ## import three ASN motifs for one protein domain from fasta files
> motif_folder = "extdata/alignments"
> motif_names_asn = c("F-box_fungi.seq", "F-box_metazoa.seq",
                       "F-box_viridiplantae.seq")
> motifs_asn = list()
> for (name in motif_names_asn) {
    fileName = paste(motif_folder,"/",name,".fa",sep="")
    file = system.file(fileName, package = "DiffLogo")
    fileContent <- readLines(file)</pre>
    fileContent <- fileContent[seq(from = 2, by = 2,
                      length.out = floor(length(fileContent)/2))]
    motifs_asn[[name]] <- getPwmFromAlignment(fileContent, ASN, 1)</pre>
+ }
```

Here, we import a set of nine DNA motifs for transcription factor CTCF from matrix files, a set of DNA motifs for three different E-Box transcription factors from sequences in tabular files, and a set of three ASN motifs for the F-Box protein domain from FASTA files.

# 4 Plot sequence logo

The user is able to examine motifs using the classical sequence logo from package seqLogo [7].

```
> ## plot classic sequence logo
> library(seqLogo)
> seqLogo::seqLogo(pwm = pwm1)
```

The user is also able to plot sequence logos with custom functions for stack height and base distribution using the package DiffLogo. In case of stack-Height=informationContent and baseDistribution=probabilities, the result is equivalent to the result of package seqLogo

```
> ## plot custom sequence logo
> par(mfrow=c(2,1), pin=c(3, 1), mar = c(2, 4, 1, 1))
```

```
> DiffLogo::seqLogo(pwm = pwm1)
> DiffLogo::seqLogo(pwm = pwm2, stackHeight = sumProbabilities)
> par(mfrow=c(1,1), pin=c(1, 1), mar=c(5.1, 4.1, 4.1, 2.1))
```

## 5 Plot difference logo

The user is easily able to plot a difference logo for a pair of motifs.

```
> ## plot DiffLogo
> diffLogoFromPwm(pwm1 = pwm1, pwm2 = pwm2)
> ## diffLogoFromPwm is a convenience function for
> diffLogoObj = createDiffLogoObject(pwm1 = pwm1, pwm2 = pwm2)
> diffLogo(diffLogoObj)
```

## 6 Plot table of difference logos

The user is easily able to plot a table of difference logos for a set of motifs.

```
> ## plot table of difference logos for CTFC motifs (DNA)
> diffLogoTable(PWMs = motifs_dna, )
> ## plot table of difference logos for E-Box motifs (DNA)
> diffLogoTable(PWMs = motifs_dna2)
> ## plot table of difference logos for F-Box motifs (ASN)
> diffLogoTable(PWMs = motifs_asn, alphabet = ASN)
```

### 7 Export visualization

The user is able to export the generated visualizations in various formats. Please find two examples below.

```
> ## parameters
> widthToHeightRatio = 16/10;
> size = length(motifs_dna) * 2
> resolution <- 300
> width <- size * widthToHeightRatio
> height <- size
> ## export single DiffLogo as pdf document
```

```
> fileName <- "Comparison_of_two_motifs.pdf"
> pdf(file = fileName, width = width, height = height)
> diffLogoFromPwm(pwm1 = pwm1, pwm2 = pwm2)
> dev.off()

pdf
   2
> ## export DiffLogo table as png image
> fileName <- "Comparison_of_multiple_motifs.png"
> png(
+ filename = fileName, res = resolution,
+ width = width * resolution, height = height * resolution)
> diffLogoTable(PWMs = motifs_dna)
> dev.off()

pdf
   2
```

#### Literature

- [1] http://en.wikipedia.org/wiki/Position\_weight\_matrix
- [2] Schneider TD, Stephens RM. 1990. Sequence Logos: A New Way to Display Consensus Sequences. Nucleic Acids Res. 18:6097-6100
- [3] Shannon P (2014). MotifDb: An Annotated Collection of Protein-DNA Binding Sequence Motifs. R package version 1.10.0.
- [4] Eggeling, R., Gohr, A., Keilwagen, J., Mohr, M., Posch, S., Smith, A.D., Grosse, I.: On the value of intra-motifdependencies of human insulator protein ctcf. PLoS ONE 9(1), 85629 (2014). doi:10.1371/journal.pone.0085629
- [5] Mordelet, Fantine and Horton, John and Hartemink, Alexander J and Engelhardt, Barbara E and Gordân, Raluca: Stability selection for regression-based models of transcription factor-DNA binding specificity. Bioinformatics 29(13), 11725 (2013). doi:10.1093/bioinformatics/btt221
- [6] Finn, R.D., Bateman, A., Clements, J., Coggill, P., Eberhardt, R.Y., Eddy, S.R., Heger, A., Hetherington, K., Holm, L., Mistry, J., Sonnhammer, E.L.L., Tate, J., Punta, M.: Pfam: the protein families database. Nucleic Acids Research 42(D1), 222230 (2014). doi:10.1093/nar/gkt1223
- [7] Bembom O. seqLogo: Sequence logos for DNA sequence alignments. R package version 1.34.0.