An introduction to the nuCpos package

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1 About nuCpos

nuCpos, a derivative of NuPoP, is an R package for predicting nucleosome positions. nuCpos calculates local and whole nucleosomal HBA scores for a given 147-bp sequence. This package was designed to demonstrate the use of chemical maps in prediction. As the parental package NuPoP now provides chemical-map-based prediction, the function for dHMM-based prediction was removed from this package. nuCpos continues to provide functions for HBA calculation. The models are based on chemical maps of nucleosomes from budding yeast Saccharomyces cerevisiae (Brogaard et al. (2012)), fission yeast Schizosaccharomyces pombe (Moyle-Heyrman et al. (2012)), or embryonic stem cells of house mouse Mus musculus (Voong et al. (2016)).

The parental package NuPoP, licensed under GPL-2, was developed by Ji-Ping Wang and Liqun Xi. Please refer to Xi et al. (2010) and Wang et al. (2008) for technical details of NuPoP. Their excellent codes were adapted in nuCpos to demonstrate the usefulness of chemical maps in prediction.

Note that when nuCpos was released, NuPoP only used an MNase-seq-based map of budding yeast nucleosomes to train a duration hidden Markov model. However, as NuPoP now provides chemical mapbased prediction, users are encouraged to use NuPoP functions to conduct dHMM-based prediction in their original way.

2 nuCpos functions

nuCpos has two functions: HBA, and localHBA.

The functions HBA and localHBA receive a sequence of 147-bp DNA and calculate whole nucleosomal and local HBA scores. These functions invoke core Fortran codes for HBA calculation that were adapted from the excellent dHMM code of *NuPoP*.

nuCpos requires the Biostrings package, especially when DNA sequences are given as DNAString objects to the functions HBA, and localHBA. These functions can also receive DNA sequences as simple character string objects without loading the Biostrings package. Note: nuCpos requires the NuPoP package to perform some example runs.

Load the nuCpos package as follows:

> library(nuCpos)

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3 Histone binding affinity score calculation with HBA

HBA score can be calculated for a given 147-bp sequence with the HBA function. In the examples bellow, a character string object inseq and a DNAString object INSEQ with the same 147-bp DNA sequences are given to HBA. Note: the *Biostrings* package is required for the latter case.

```
> load(system.file("extdata", "inseq.RData", package = "nuCpos"))
> HBA(inseq = inseq, species = "sc")
      HBA
-2.460025
> for(i in 1:3) cat(substr(inseq, start = (i-1)*60+1,
      stop = (i-1)*60+60), "\n"
ATCGAGAATCCCGGTGCCGAGGCCGCTCAATTGGTCGTAGACAGCTCTAGCACCGCTTAA
ACGCACGTACGCGCTGTCCCCCGCGTTTTAACCGCCAAGGGGATTACTCCCTAGTCTCCA
GGCACGTGTCAGATATATACATCCGAT
> load(system.file("extdata", "INSEQ_DNAString.RData",
      package = "nuCpos"))
+
> INSEQ
147-letter DNAString object
seq: ATCGAGAATCCCGGTGCCGAGGCCGCTCAATTGGTC...TAGTCTCCAGGCACGTGTCAGATATATACATCCGAT
> HBA(inseq = INSEQ, species = "sc")
      HBA
-2.460025
```

The argument *inseq* is the character string object to be given. Alternatively, a DNAString object can be used here. The length of DNA must be 147 bp. The argument *species* can be specified as follows: mm = M. musculus; sc = S. cerevisiae; sp = S. pombe.

4 Local histone binding affinity score calculation with localHBA

Local HBA scores are defined as HBA scores for 13 overlapping subnucleosomal segments named A to M. They can be calculated for a given 147-bp sequence with the localHBA function. Like HBA, this function can receive either a character string object or a DNAString object. The segment G corresponds to the central 21 bp region, in which the dyad axis passes through the 11th base position. This means that the local HBA score for the G segment implies the relationship between DNA and histone proteins at around superhelical locations -0.5 and +0.5. The neighboring F segment, which is 20 bp in length, is for SHLs -1.5 and -0.5. The result of example run shown below suggests that subsequence of inseq around SHL -3.5 and -2.5 is suitable for nucleosome formation.

```
> localHBA(inseq = inseq, species = "sc")
```

```
1HBA_A
                 1HBA_B
                              1HBA_C
                                          1HBA_D
                                                       1HBA_E
                                                                   1HBA_F
-1.56140949 -1.62502354
                         0.48885990
                                      2.37615568
                                                  2.90458625 -1.35195919
     1HBA_G
                 1HBA_H
                              1HBA_I
                                          1HBA_J
                                                       1HBA_K
                                                                   1HBA_L
-3.13228907 -0.32208031
                         0.27650871
                                      0.01922002
                                                  0.49787625 -0.17151500
     1HBA_M
-1.27186158
> barplot(localHBA(inseq = inseq, species = "sc"),
      names.arg = LETTERS[1:13], xlab = "Nucleosomal subsegments",
      ylab = "local HBA", main = "Local HBA scores for inseq")
```

5 Acknowledgements

We would like to thank Drs. Shimizu, Fuse and Ichikawa for sharing DNA sequences and *in vivo* data, and giving fruitful comments. We would like to thank Dr. Ji-Ping Wang and his colleagues for distributing NuPoP under the GPL-2 license. In this package, their excellent code for dHMM-based prediction was adapted for chemical map-based prediction to demonstrate the usefulness of chemical maps in prediction. As we noticed that canceling of HBA smoothing helps predicting rotational settings, predNuCpos in the earlier version provided this option. However, for those who want to predict nucleosome occupancy in the original way with chemical maps, we encourage users to use NuPoP functions as it now provides chemical map-based predictions. In our functions HBA and localHBA, their excellent code was also adapted to calculate the scores of given 147-bp sequences independently of the genomic context. The function HBA now runs without invoking a fortran subroutine.

References

- Wang JP, Fondufe-Mittendorf Y, Xi L, Tsai GF, Segal E and Widom J (2008). Preferentially quantized linker DNA lengths in Saccharomyces cerevisiae. PLoS Computational Biology, 4(9):e1000175.
- Xi L, Fondufe-Mittendorf Y, Xia L, Flatow J, Widom J and Wang JP (2010). Predicting nucleosome positioning using a duration hidden markov model. *BMC Bioinformatics*, 11:346.
- Brogaard K, Xi L, and Widom J (2012). A map of nucleosome positions in yeast at base-pair resolution. *Nature*, 486(7404):496-501.
- Moyle-Heyrman G, Zaichuk T, Xi L, Zhang Q, Uhlenbeck OC, Holmgren R, Widom J and Wang JP (2013). Chemical map of *Schizosaccharomyces pombe* reveals species-specific features in nucleosome positioning. *Proc. Natl. Acad. Sci. U. S. A.*, 110(50):20158-63.
- Ichikawa Y, Morohoshi K, Nishimura Y, Kurumizaka H and Shimizu M (2014). Telomeric repeats act as nucleosome-disfavouring sequences in vivo. *Nucleic Acids Res.*, 42(3):1541-1552.
- Voong LN, Xi L, Sebeson AC, Xiong B, Wang JP and Wang X (2016). Insights into Nucleosome Organization in Mouse Embryonic Stem Cells through Chemical Mapping. Cell, 167(6):1555-1570.
- Fuse T, Katsumata K, Morohoshi K, Mukai Y, Ichikawa Y, Kurumizaka H, Yanagida A, Urano T, Kato H, and Shimizu M (2017). Parallel mapping with site-directed hydroxyl radicals and micrococcal nuclease reveals structural features of positioned nucleosomes in vivo. *Plos One*, 12(10):e0186974.