




periodicDNA: an R package to investigate oligonucleotide periodicity

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Statement of Need

periodicDNA provides a framework to quantify oligonucleotide periodicity over individual or multiple DNA genomic loci.

Summary

periodicDNA is an R package offering a set of functions to identify local periodic elements in short sequences such as regulatory elements. Many oligonucleotides are periodically occurring in genomes across eukaryotes, and some are impacting the physical properties of DNA. Notably, DNA bendability is modulated by 10-bp periodic occurrences of WW (W = A/T) dinucleotides. The package relies on Biostrings and GenomicRanges packages to handle DNA sequences and genome assemblies. It uses the Fourier Transform to measure the periodicity of a given oligonucleotide in sets of sequences. It also provides methods to generate continuous tracks of oligonucleotide periodicity over genomic loci, as well as visualization tools to interpret these tracks. The use of periodicDNA has already shed light on fundamental differences in sequence features in functional classes of promoters (Serizay et al. (2020)). We hope that the integration of this open-source package into genomic assay analysis workflows will help further improve our understanding of chromatin organization.

Methodology

periodicDNA can be used to estimate the power spectral density (PSD) of a given dinucleotide (motif argument) at specific periods (period argument) in a set of sequences of interest (seqs argument), using a simple wrapper function:

```
{r eval = FALSE} library(periodicDNA) library(magrittr) library(ggplot2)
## The periodicity can be calculated from DNASTringSet objects: data(ce_proms_seqs)
score <- getPeriodicity(      seqs = ce_proms_seqs,      motif = 'TT',
cores = 4 ) ## Alternatively, the periodicity can be calculated  ## from
a GRanges object in combination with a genome: data(ce_proms) score <-
getPeriodicity(      granges = ce_proms[ce_proms$which.tissues == 'Ubiqu.'],
%>%      '['(strand(.) == '+') %>%      resize(width = 1, fix
= 'end') %>%      resize(width = 300, fix = 'start'),      genome =
'cell1',      motif = 'TT',      cores = 120 ) ## Results can be plotted
```

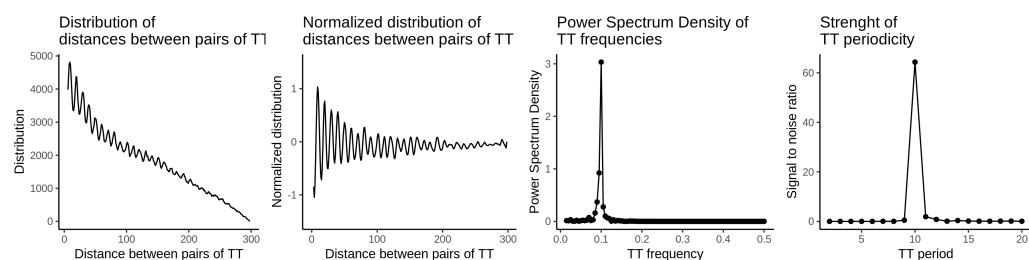


Figure 1: Output of `plotPeriodicityResults()` function. A- Distribution of all the TT pairwise distances. B- Normalized TT pairwise distance frequency. C- Power Spectral Density (PSD) of TT occurrences. D- PSD signal-to-noise ratio.

```
using the plotPeriodicityResults() function: ## See Figure 1 plots <-
plotPeriodicityResults(score) %>%      cowplot::plot_grid(plotlist = .,
nrow = 1)
```

The intermediate steps internally performed when calling this function are the following (Figure 2):

1. In each sequence of a set of `n` sequences (the `seqs` argument), all the pairs of the dinucleotide of interest (the `motif` argument, e.g. TT) are identified and their pairwise distances are measured.
2. The distribution of the all the resulting pairwise distances (also called “distogram”) is generated.
3. The following normalization steps are then performed:
 - The distogram is transformed into a frequency histogram and then normalized by the following steps:
 - The frequency histogram follows a marked overall decrease of frequencies with increased pairwise distances. Indeed, for a 200-bp long sequence containing 20 WW dinucleotides exactly distant from each other by 10 base pairs, there are 19 pairs with a pairwise distance of 10 but only 1 pair of dinucleotides with a pairwise distance of 190. To overcome this distance decay, the frequency histogram is smoothed using a moving average window of 10 and the resulting smoothed frequency histogram is subtracted from the frequency histogram. This effectively transforms the decreasing frequency histogram into a dampened oscillating signal and improves the PSD estimation by Fourier Transform.
 - The dampened oscillating signal is then scaled (i.e. mean-centered and normalized) and smoothed using a moving average window of 3. This effectively removes the effect of the latent 3-bp periodicity of most dinucleotides found in eukaryote genomes (Gutiérrez et al., 1994).
4. A Fast Fourier Transform (FFT) is then used to estimate the power spectral density (PSD) of the normalized oscillating distribution at different periods (the `period` argument).

The PSD can be used in itself to identify which dinucleotide frequencies are enriched in the provided set of sequences. Its amplitude at a given frequency can also be used to compare dinucleotide frequencies across samples.

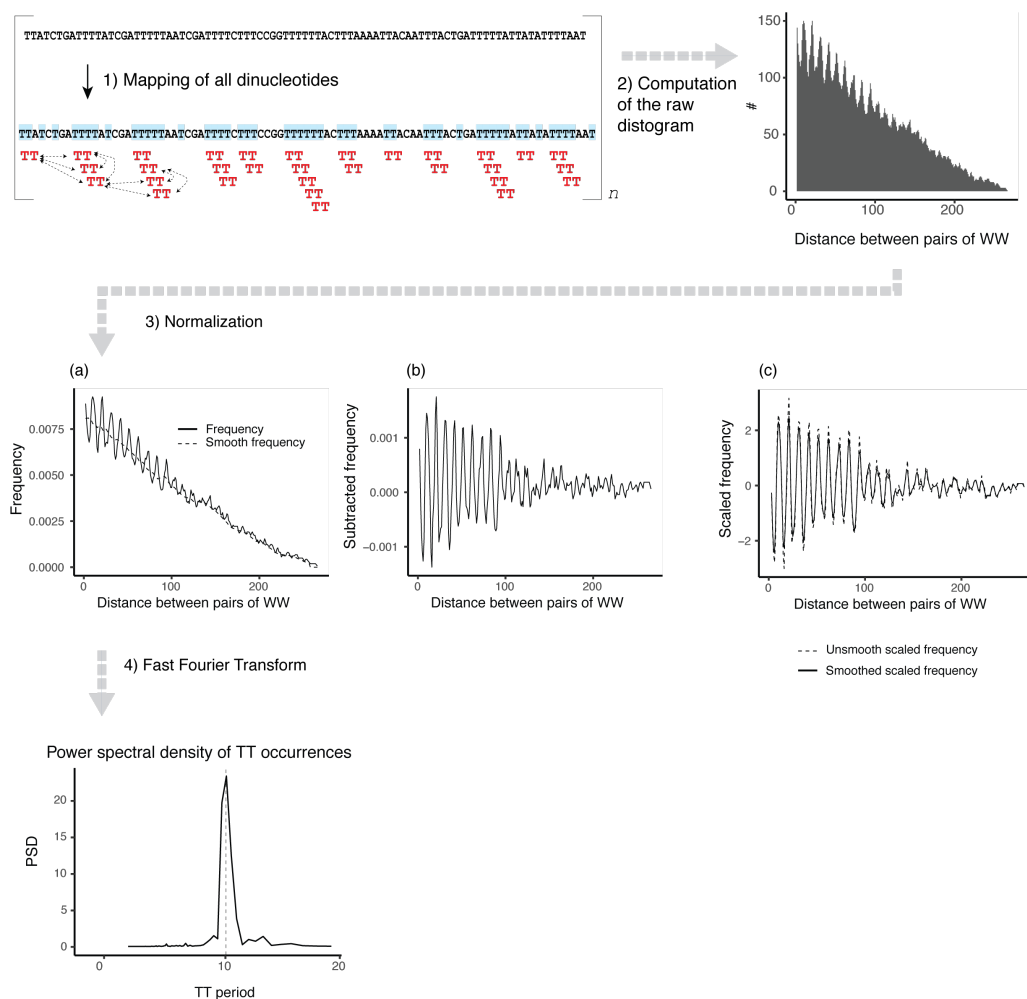


Figure 2: Internal steps of the `getPeriodicity()` function. Each step is further described in the main text. The dotted double-arrows in the first step represent the distances measured by `periodicDNA` between some of the pairs of TT. For the single sequence shown here, there are 31 individual TT dinucleotides, resulting in $\binom{31}{2} = 465$ different pairs in total.

Leveraging periodicDNA to understand chromatin organization

Soon after solving the structure of nucleosomes, Kornberg raised a fundamental question: whether the positioning of nucleosomes *in vivo* in regard to a DNA locus was “specific” or “statistical” (Kornberg (1981)). Nucleosome “specific” positioning implies that the physicochemical properties of DNA sequences are enough to explain how nucleosomes are arranged along a DNA double-helix (e.g. described in Segal et al. (2006)). On the contrary, a “statistical” positioning postulates the presence of a “boundary” nucleosome (either a protein or a strong intrinsic positioning sequence, or both) which specifies one end of a nucleosomal array not determined by the physicochemical properties of DNA sequence (e.g. described in Mavrich et al. (2008)). Later on, biochemists and computational biologists found out that periodic dinucleotide sequences were associated with positioned nucleosomes, suggesting that the “specific” model is contributing to nucleosome positioning - at least to a certain extent (Jiang & Pugh (2009); Struhl & Segal (2013) for review). To test whether specific periodic sequences were associated with the positioning of nucleosomes directly flanking regulatory elements, I leveraged periodicDNA. I focused on ubiquitous and tissues-specific promoters and enhancers, splitting each element into core (-70 to +70 base pairs around the center of the regulatory element), flanking (-210 to -70 base pairs and +70 to +210 base pairs) and distal sequences (-350 to -210 base pairs and +210 to +350 base pairs) (Figure 3A). Ubiquitous and germline-specific promoters exhibit a high TT 10-bp periodicity in the flanking sequences which decreases remarkably in the neighboring distal sequences (Figure 3B). In contrast, ubiquitous and germline enhancers both show a mild increase of TT 10-bp periodicity in flanking sequences as well as in distal sequences of enhancers (Figure 3B). This suggests that the 10bp TT periodicity can act as a local positioning signal at ubiquitous and germline-specific promoters, but not at ubiquitous and germline-specific enhancers (Figure 3C). This 10-bp TT periodicity is absent in other somatic tissue-specific promoters, as expected from previous results (Serizay et al. (2020)), as well as in somatic tissue-specific enhancers (Figure 3B). This supports a model whereby nucleosome organization at soma-restricted regulatory elements does not primarily depend on the underlying DNA sequence (Figure 3C).

Acknowledgments

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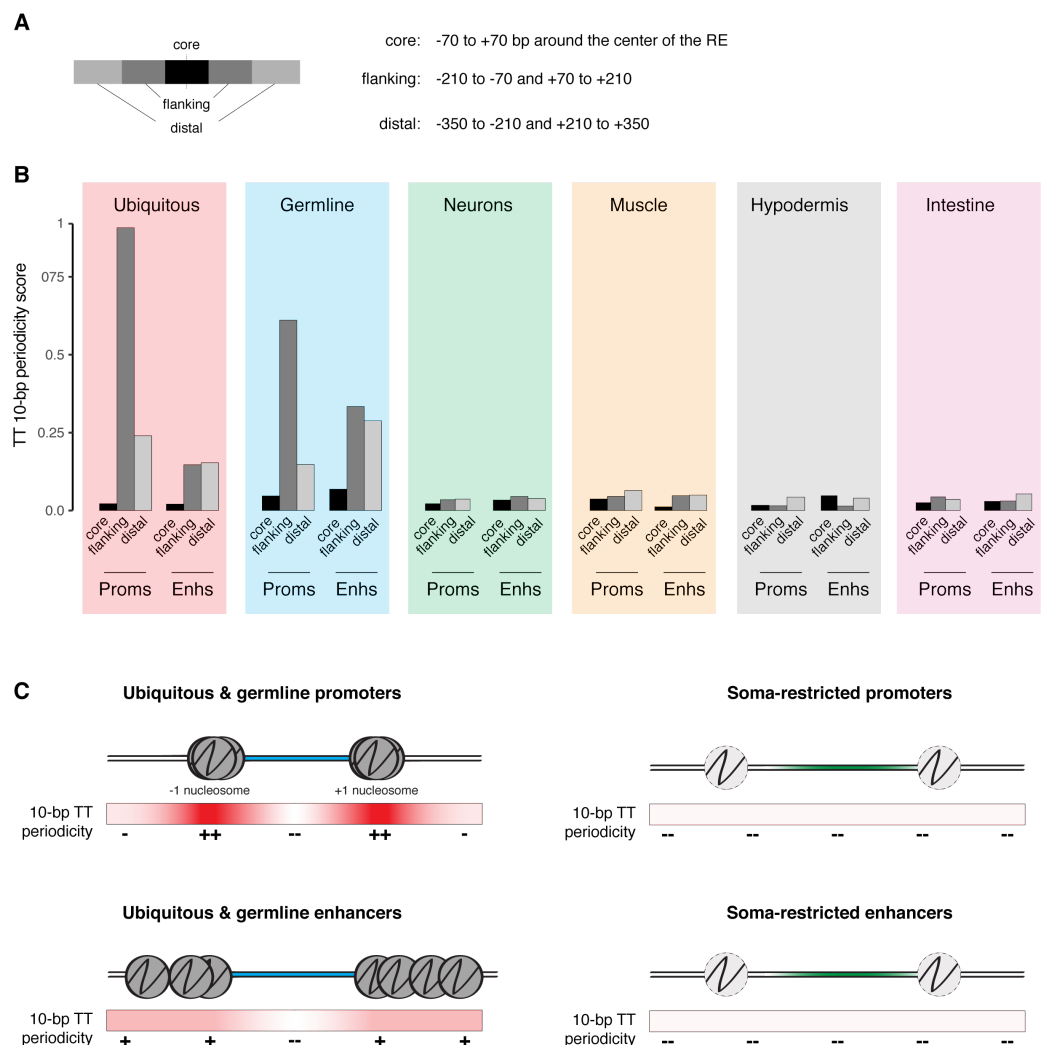


Figure 3: TT periodicity in promoters and enhancers. A- Pictogram representing how regulatory elements were divided into core, flanking and distal regions. The core sequence is the 140-bp long sequence at the center of the regulatory element; the flanking sequences range from -210 to -70 and from +70 to +210; the distal sequences range from -350 to -210 and from +210 to +350 (with the center of the regulatory element being the reference). B- TT 10-bp periodicity scores obtained from periodicDNA. C- Model of sequence-driven nucleosome positioning at different sets of promoters or enhancers. Three different situations are observed: (1) a decrease of TT periodicity on both sides of the flanking nucleosomes favors their precise positioning, (2) a weaker widespread TT periodicity favors nucleosome positioning without local enrichment and (3) absence of TT periodicity does not favor nucleosome positioning. Note that these models do not illustrate the role of other factors such as chromatin remodelers.

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