CAGEfightR
analysis of 5’-end data using R/Bioconductor
Thodberg, Malte; Thieffry, Axel; Vitting-Seerup, Kristoffer; Andersson, Robin; Sandelin, Albin

Published in:
BMC Bioinformatics

DOI:
10.1186/s12859-019-3029-5

Publication date:
2019

Document version
Publisher’s PDF, also known as Version of record

Document license:
CC BY

Citation for published version (APA):
Thodberg, M., Thieffry, A., Vitting-Seerup, K., Andersson, R., & Sandelin, A. (2019). CAGEfightR: analysis of 5’-end data using R/Bioconductor. BMC Bioinformatics, 20, [487]. https://doi.org/10.1186/s12859-019-3029-5
CAGEfightR: analysis of 5′-end data using R/Bioconductor

Malte Thodberg1,2*, Axel Thieffry1,2, Kristoffer Vitting-Seerup1,2,3, Robin Andersson1 and Albin Sandelin1,2*

Abstract

Background: 5′-end sequencing assays, and Cap Analysis of Gene Expression (CAGE) in particular, have been instrumental in studying transcriptional regulation. 5′-end methods provide genome-wide maps of transcription start sites (TSSs) with base pair resolution. Because active enhancers often feature bidirectional TSSs, such data can also be used to predict enhancer candidates. The current availability of mature and comprehensive computational tools for the analysis of 5′-end data is limited, preventing efficient analysis of new and existing 5′-end data.

Results: We present CAGEfightR, a framework for analysis of CAGE and other 5′-end data implemented as an R/Bioconductor-package. CAGEfightR can import data from BigWig files and allows for fast and memory efficient prediction and analysis of TSSs and enhancers. Downstream analyses include quantification, normalization, annotation with transcript and gene models, TSS shape statistics, linking TSSs to enhancers via co-expression, identification of enhancer clusters, and genome-browser style visualization. While built to analyze CAGE data, we demonstrate the utility of CAGEfightR in analyzing nascent RNA 5′-data (PRO-Cap). CAGEfightR is implemented using standard Bioconductor classes, making it easy to learn, use and combine with other Bioconductor packages, for example popular differential expression tools such as limma, DESeq2 and edgeR.

Conclusions: CAGEfightR provides a single, scalable and easy-to-use framework for comprehensive downstream analysis of 5′-end data. CAGEfightR is designed to be interoperable with other Bioconductor packages, thereby unlocking hundreds of mature transcriptomic analysis tools for 5′-end data. CAGEfightR is freely available via Bioconductor: bioconductor.org/packages/CAGEfightR.

Keywords: Transcription start site, Promoter, Enhancer, Enhancer RNA, CAGE, 5′-end methods, R-package, Bioconductor

Background

Transcription start sites (TSSs) are central entities of transcriptional regulation, where a wide range of cues from surrounding factors such as core promoter elements, transcription factor binding sites, chromatin modifications, and distal elements such as enhancers and silencers are integrated to decide whether transcription initiation takes place, and with what rate [1–3]. Hence, accurate identification of TSSs and their activity is a prerequisite for understanding gene regulation.

Several genome-wide, high-throughput sequencing assays have been developed for identifying TSS activity, all based on the idea of capturing and sequencing only the 5′-end of RNAs (called tags), leading to the collective name of 5′-end methods [4]. In terms of TSS identification, such methods have distinct advantages over other assays, e.g. RNA-sequencing (RNA-Seq) and Chromatin Immunoprecipitation Sequencing (ChIP-Seq). While RNA-Seq is widely used for studying gene expression and splicing, it is ineffective for accurate detection of TSSs. This is due to the random fragmentation of RNA molecules, leading to a trail-off of sequencing reads near the end of transcripts. In contrast, 5′-end methods effectively pile up reads at TSSs, providing high local coverage for accurate prediction of TSSs. Similarly, ChIP-Seq targeting RNA polymerase II or pre-initiation complex proteins has low positional resolution due to the length of ChIP-Seq fragments, and does not explicitly measure TSS usage.

* Correspondence: malte.thodberg@bric.ku.dk; albin@bimf.ku.dk
1Department of Biology, University of Copenhagen, Ole Maaløes Vej 5, DK2100, Copenhagen N, Denmark
Full list of author information is available at the end of the article

© The Author(s). 2019 Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.

BMC Bioinformatics
The majority of available 5′-end methods capture steady-state capped RNAs (Table 1). This allows for identification of messenger RNA (mRNA) TSSs, including non-characterized alternative TSSs since the methods are not contingent on annotated transcript models. Alternative TSS usage is often tissue/cell-specific, and common in mammals [14, 15], plants [6], etc. and in clinical settings (inflammatory bowel disease [25], diabetes [26], cancer [27], Retts Syndrome [28]). Despite its wide usage, the current toolbase available for analyzing 5′-end data is not as developed as that for RNA-Seq or ChIP-Seq (Table 2). Most tools are either stand-alone pipelines (MOIRAI [29], RECLU [32], etc.) or focused on a single analysis problem, e.g. tag clustering (paraclu [30], CapFilter [6], etc., further discussed below), making it hard to combine different tools. An alternative to stand-alone tools is using R-packages from the Bioconductor [39] project, which allows easier interoperability between tools due to shared data representations. Bioconductor currently contains three packages (CAGEr [34], icetea [37], TSRchitect [38]) for analyzing 5′-end data in general and CAGE in particular. While these packages offer functionality for TSS identification, quantification and annotation, they lack any functions for predicting, quantifying and analyzing enhancer candidates, and are not efficiently scalable for large datasets.

To solve the above problems, we here introduce the CAGEfightR R/Bioconductor package for analyzing 5′-end data (including CAGE).

### Table 1: Examples of popular 5′-end methods

| Technology                                                                 | Acronym     | RNA state   | Ref. |
|---------------------------------------------------------------------------|-------------|-------------|------|
| Cap Analysis of Gene Expression                                           | CAGE        | Steady-state | [5]  |
| Nano CAGE                                                                 | NanoCAGE    | Steady-state | [6]  |
| Super Low Input Carrier CAGE                                              | SLIC-CAGE   | Steady-state | [7]  |
| no-Amplification non-Tagging CAGE                                         | nAnT-iCAGE  | Steady-state | [8]  |
| Transcription Start Site Sequencing                                       | TSS-Seq     | Steady-state | [9]  |
| RNA Annotation and Mapping of Promoters for the Analysis of Gene Expression | RAMPAGE     | Steady-state | [10] |
| Single-cell Tagged Reverse transcription                                  | STRT        | Steady-state | [11] |
| Precision Nuclear Run-on Sequencing for RNA Polymerase II Start Sites     | PRO-Cap     | Nascent     | [12] |
| 5′ Global Run-on Sequencing                                               | GRO-Cap/5′ GRO-Seq | Nascent | [13] |

### Table 2: Examples of software packages for analyzing 5′-end data (including CAGE)

| Tool            | Implementation      | Input data | Tag Clustering       | TSS candidate shape | Differential Expression | Gene-level analysis | Unique features                                      |
|-----------------|---------------------|------------|----------------------|---------------------|-------------------------|---------------------|------------------------------------------------------|
| MOIRAI [29]     | Graphical User Interface | FASTQ      | paraclu [30]         | None                | None                    | None                | RNAAdjust, TagDust [31]                             |
| RECLU [32]      | Bash                | BED        | modified paraclu [30]| None                | edgeR [33]             | None                | Hierarchical TSSs                                   |
| CAGEr [34]      | R/Bioconductor      | BAM        | distance or paraclu [30] | IQR                 | DESeq2 [35]            | Gene expression | G-bias correction [36], power-law normalization, TSS shifts |
| icetea [37]     | R/Bioconductor      | FASTQ      | Sliding window       | None                | edgeR [33]             | Gene expression | Paired-end methods, mapping via R                    |
| TSRchitect [38] | R/Bioconductor      | BAM        | X-means              | Shape Index [17]    | None                    | Gene expression | Enhancer calling, TSS-enhancer co-expression, super enhancers |
| CAGEfightR      | R/Bioconductor      | BigWig     | Slice-reduce         | IQR, entropy, etc.  | None                    | Gene expression and alternative TSS usage |                                                     |
end data. CAGEfightR is the first single framework that robustly detects, quantifies, annotates, links and visualizes TSSs and enhancer candidates in a manner that is highly compatible with other Bioconductor packages. The memory efficient and scalable implementation allows CAGEfightR to be used on datasets ranging from small-scale experiments to consortia-level projects. In this way, CAGEfightR unlocks hundreds of packages developed for RNA-Seq and ChIP-Seq for CAGE and similar types of 5′-end data.

Implementation
CAGEfightR is implemented purely in R making use of several R-packages from the Bioconductor project. It is based on standard Bioconductor S4-classes, primarily GRanges (GenomicRanges), RangedSummarizedExperiment (SummarizedExperiment) [40] and GInteractions (InteractionSet) [41] and visualization via Gviz [42] and GenomicInteractions [43]. This makes it easy to use CAGEfightR in conjunction with other Bioconductor packages.

5′-end data is conventionally stored, shared and analyzed by first mapping tags to the genome, followed by counting the number of 5′-ends of tags mapping to each individual base pair (bp), on each strand. In CAGE terminology, such data are referred to as CAGE-defined TSSs (CTSSs) [36], but we use the term generally for all 5′-end methods here. The processing of tags differs between 5′-methods due to distinctive protocols (5′-end isolation technique, single-end vs. paired-end sequencing, etc.), and for CAGE in particular specialized tools have been developed, e.g. rRNAAdust for removing contaminant ribosomal RNA (http://fantom.gsc.riken.jp/5/suppl/rRNAAdust/) and/or removing G’s added by reverse transcriptase at cDNA 5′-ends [36]. While filtering, mapping, and counting of tags can be done efficiently by dedicated tools a single library at a time, CAGEfightR is focused on analysis from the point when multiple libraries are jointly analyzed. To be as general as possible, CAGEfightR was designed to import and analyze 5′-end data after mapping and processing by starting from CTSSs from each library stored as BigWig files.

Most genome bps are not CTSSs (have no tags mapped to them), and only a small fraction of CTSSs have a high number of tags. CAGEfightR takes advantage of this sparsity by using sparse representations to efficiently store and analyze large CTSS datasets using little memory. This allows tens of samples to be analyzed on a typical laptop computer and hundreds of samples on a typical server. Most computationally heavy tasks can be parallelized, providing further speed increases when multiple cores or clusters are available.

As described below, CAGEfightR can analyze 5′-end data on three different levels: bp-accurate CTSSs (Fig. 1a, top), clusters of nearby CTSSs (Fig. 1a, middle) or expression summed over known gene models (Fig. 1a, bottom), where each analysis level is associated with a specific expression matrix (Fig. 1a, right). These expression matrices and other data structures used in CAGEfightR are designed to be readily usable by other Bioconductor packages, in particular popular differential expression packages such as limma [44], edgeR [33], DESeq2 [35], DEXSeq [45], DRIMSeq [46], etc.

Results and discussion
Below, we overview the core functionality of CAGEfightR with examples using previously published 5′-end data. Names of the main CAGEfightR functions for each analysis are indicated in Fig. 1, 2, 3, 4, 5. Genome-browser figures (Figs. 1b-c, 2d, 3b-c, 4c, 5d) are otherwise unedited output from R/CAGEfightR.

Analysis of 5′-end tags
CAGEfightR can import CTSSs from BigWig files and quantify their expression levels across all samples. The CTSSs can then be normalized to Tags-Per-Million (TPM) and summed across samples to yield a global or pooled CTSS signal (Fig. 1a, top). In case of a large number and/or low quality samples, CAGEfightR offers various strategies for calculating more robust pooled CTSS signals, chiefly by filtering CTSSs only observed in a single or few samples. The pooled CTSS signal can be visualized in genome-browser style along the genome (Fig. 1b-c, 2d, 3c, 4c, 5d).

Analysis of tag clusters
Pooled CTSSs can be used to identify clusters of closely spaced CTSSs on the same strand, referred to as unidirectional clusters, or conventionally Tag Clusters (TCs) in most CAGE papers. CAGEfightR identifies TCs using a slice-reduce approach: First, CTSSs with pooled CTSS signal below a chosen threshold are discarded (slice) and surviving CTSSs on the same strand are then merged into clusters (reduce) (Fig. 1a-b). CAGEfightR includes a host of functions for analyzing such TCs, including filtering on expression, hierarchical annotation of TCs with transcript models and analysis of TC shapes (see below).

TCs reflect the fact that when RNA-polymerase associates with the DNA, it rarely initiates from a single bp, but rather from an array of nearby bps. Such arrays are expected to produce nearly identical RNAs that are subject to the same regulatory cues, as they will share the same promoter sequence and genomic neighborhood. While genes are transcribed from multiple different CTSSs, these CTSSs are grouped in a single or multiple TCs corresponding to the major RNAs (or transcripts/isoforms in RNA-Seq terminology) produced from the gene.
Because of this, many studies use a simplification in which such TCs are referred to as TSSs for genes, even though technically TCs group several nearby bp-accurate TSS / CTSSs. To avoid confusion on terminology, and remain consistent with previous CAGE literature, we will primarily use the term `TC` to describe unidirectional clusters.

As an applied example, we analyzed three HeLa CAGE libraries from Andersson et al [47]. Using CAGEfightR,
we calculated the pooled CTSS signal across libraries, and from that defined 22,760 TCs with >1 TPM in at least two samples. As these TCs are defined de novo, it is useful to see how they relate to known transcript models. We therefore annotated the TCs according to University of California Santa Cruz (UCSC) transcripts using CAGEfightR’s hierarchical scheme (Fig. 2a). The hierarchical scheme accounts for the existence of multiple transcripts or isoform of genes, i.e. an annotated promoter of one transcript can be in the 5’-UTR of another transcript from the same gene. CAGEfightR can assign 9 different annotation categories (custom categories can be supplied by the user), based on the most likely association with known transcripts, e.g. a TC is more likely to correspond to an annotated promoter rather than a novel intragenic promoter.

Using this method, we plotted the number of TCs falling into the different annotation categories and their expression distributions (Fig. 2b). Most TCs candidates were found at annotated promoters and were generally highly expressed. However, a substantial number of novel (not overlapping annotated promoters) TCs were
identified, in particular in the promoter-proximal region and 5′-UTR, which on average had lower expression than those overlapping annotated promoters.

In vertebrates, the distribution of CTSSs within TCs is related to cell specificity and DNA sequence properties: sharp CTSS distributions have an overrepresentation of TATA-boxes and are more cell- or tissue-specific, while broad CTSS distributions are GC-rich and more ubiquitously expressed [36]. Classification is often based on the width of the CTSS distribution, expressed as the interquartile/interquantile range (IQR), as this measures the width of the bulk of CTSSs within a TC without being affected by a few straggler CTSSs potentially greatly extending the width of the TC. We used CAGEfightR to calculate the genomic region covering the 5′–95% IQR of total CTSS expression for each TC in the HeLa set (Similar results were obtained with tighter IQR intervals). This showed a clear bimodal distribution corresponding to sharp and broad CTSS distributions (Fig. 2c-d). Investigation of promoter regions using sequence logos also confirmed that sharp, but not broad distributions had a stronger TATA box (Fig. 2e).

Analysis of enhancer candidates

Active enhancers are characterized by bidirectional transcription initiation of eRNAs [48]. In 5′-end data, this manifests as bidirectional clusters (BCs) of CTSSs, which can be used to systematically identify enhancer candidates [21]. Similarly to above, CAGEfightR uses a slice-reduce approach to identify bidirectional clusters (BCs, as opposed to the previously discussed unidirectional TCs) to predict enhancers (Fig. 1a, c). First, the upstream and downstream pooled CTSSs are quantified for every genomic position. Second, the Bhattacharyya coefficient [49] is used to quantify the departure of the observed pooled CTSS signal from perfect bidirectionality, producing a bidirectionality or balance score for each bp (Additional file 1: Figure S1A). Third, locations with a balance score above a given threshold are identified, and nearby sites are merged into discrete BCs. This slice-reduce approach is conceptually similar to the original enhancer prediction method by Andersson et al. [21], but does not need an input seed of TCs used to find bidirectional pairs, and gives similar results while being more scalable (Additional file 1: Figure S1B-C). As BCs can be found at other genomic regions than active enhancers (e.g. bidirectional gene promoters), BCs in or near known exons can be filtered away to obtain a final set of enhancer candidates [21].

As an applied example, we used the same CAGE HeLa data set as above and identified BCs outside of exonic regions and more than 1 kbp upstream of annotated promoters (based on UCSC transcript models) as enhancer candidates. This resulted in a total of 6384 enhancer candidates (3780 intronic and 2604 intergenic).

As an initial validation step, we investigated whether enhancer candidates had the expected chromatin patterns compared to TCs, by overlapping with DNase I hypersensitive sites sequencing (DNase-Seq), H3K27ac, H3K4me3 and H3K4me1 ChIP-Seq signals from the same cell type. As expected, we observed high DNase sensitivity at enhancer midpoints and TC peaks, and higher levels of H3K27ac at TCs, compared to enhancer candidates. The ratio of H3K4me3 to H3K4me1 is often used to predict enhancers from ChIP-Seq signals, and consistent with this we observed low average H3K4me3 and high H3K4me1 signals around predicted enhancer candidates, and the opposite patterns around TCs (Fig. 3a) [50, 51].

Spatial prediction of enhancer-TSS links and enhancer clusters

An outstanding challenge in enhancer analysis is to link enhancers with target gene(s). Chromatin conformation capture data [52] is only available for a small set of cells, motivating computational prediction methods. A simple but popular linkage method is based on co-expression (correlation of expression) between enhancer candidates and genes across samples, assuming that true enhancer-gene pairs are co-expressed. This has been used for e.g. DNase and CAGE data, and serves as a reasonable hypothesis generator for physical interactions when the distance between enhancer and promoter is limited [21, 25, 53, 54]. CAGEfightR implements this approach by calculating the correlation of expression between enhancer candidates and TCs, with the option of supplying custom functions for calculating correlations in addition to the ones included in base R (Pearson, Spearman, Kendall).

As an example, we applied CAGEfightR to 50 CAGE samples obtained from colonic biopsies of ulcerative colitis and control subjects from Boyd et al. [25]. The reason for not using the HeLa set above was that correlations are more reliable if calculated across many samples. Specifically, we calculated a robust pooled CTSS signal (discarded CTSSs observed only in a single library), then selected TCs having >1 TPM (and >10% of total gene expression if they were assigned to genes, see below) in >6 samples, and predicted enhancer candidates >1 CAGE tag in >6 samples. This resulted in 31,480 TCs and 10,000 enhancer candidates. Using CAGEfightR, we computed the correlation (Kendall’s tau) between pairs of TCs and enhancer candidates within 10 kbp of each other, resulting in 19,271 positively correlated links between TCs and enhancers, where 978 were significant at FDR <0.05. CAGEfightR supports the visualization of multiple enhancer candidate-TC links across a genomic region. As an example, Fig. 3b shows...
Fig. 3 (See legend on next page.)
predicted enhancer-TC links around the TNFRSF1A gene, whose most dominant TC is most highly correlated with an intronic enhancer candidate in the neighboring gene.

Previously, large regions having enhancer-associated chromatin features were identified as drivers of central biological processes. Such regions are often referred to as “super”—or “stretch” enhancers [55]. Using CAGE data to predict enhancers, we have shown that many such chromatin-defined regions can be characterized as a group of bidirectionally transcribed loci, or a cluster of enhancer candidates [21, 25]. It follows that such larger regions can be predicted based on CAGE data, as genomic stretches with many enhancer candidates, and CAGEfightR implements methods for doing this. As an applied example, we used the enhancer candidates predicted based on the ulcerative colitis set above to identify enhancer clusters, defined as enhancers situated less than 12.5 kbp [25] from each other, resulting in 624 stretches with 4–24 enhancers per stretch. CAGEfightR can additionally calculate the average pairwise correlation between enhancer candidates in the stretch to reveal if they show concordant direction of change, indicative of joint activity, and provides methods to visualize these correlations (Fig. 3c).

Analysis in terms of known gene models

Although TCs can be identified de novo, it is useful to be able to analyze their expression across known gene models. Examples include the ability to compare 5′-end expression with RNA-Seq expression on gene level [56], or to link 5′-end expression estimates with gene-centric databases, such as Gene Ontology (GO) terms [57] or pathway/interaction annotation (Kyoto Encyclopedia of Genes and Genomes (KEGG) [58], STRING [51], etc.). CAGEfightR includes functions for annotating TCs to known genes and summarizing their expression within genes to obtain a gene-level expression matrix (Fig. 1a, bottom). This gene-level expression matrix can readily be used with other Bioconductor packages for gene-level analysis (e.g. limma [44], edgeR [33], DESeq2 [35]).

Another key use of gene models in relation to 5′-end methods is the analysis of alternative TSS or alternative promoter usage, which is a key contributor in generating transcript diversity (multiple different transcripts/isoforms from genes). This can be done by identifying genes harbouring several TCs on the same strand, with each TC giving rise to distinct RNAs. In this way, TCs can be seen as TSS candidates for the different transcripts/isoforms produced by a gene, phrasing the analysis in a similar way to alternative splicing or transcript usage for RNA-Seq. To be clear, this is different from analyzing changes in the distribution of CTSSs within a TC (see above), as different TCs in a gene will be widely spaced, have different promoter sequences and genomic neighbourhoods and produce different truncations of RNA, with potentially different regulation and function.

In addition to identifying such alternative TCs within gene models, CAGEfightR offers the option of filtering TCs within genes based on their contribution to overall gene expression: As 5′-end methods can detect very lowly expressed TCs, CAGEfightR can remove alternative TCs making up less than e.g. 10% of total gene expression in a given number of samples to focus only on the most highly expressed RNAs from a gene. This filtering approach is also useful when combining CAGEfightR with popular tools for differential transcript usage such as limma [44], edgeR [33], DESeq [45] and DRIMSeq [46], to investigate whether a given TC within a gene is preferentially used under certain conditions.

To illustrate these features in CAGEfightR, we used the ulcerative colitis set, and assigned TCs to genes using UCSC gene models and determined how much each such TCs contributed to overall gene expression. Without any composition filtering, 40% of all genes used more than one TC, falling to 23% when only considering TCs contributing more than 10% of total gene expression in more than 6 samples (Fig. 4a). The majority of discarded TCs were found in protein-coding, intronic and 3′-UTR regions (Fig. 4b), but several interesting cases remained, for example in the SLC16A5 gene, where we identified a highly expressed novel intronic TC (Fig. 4c).

Example of PRO-cap data analysis using CAGEfightR

While conceived as a tool for analyzing CAGE data, CAGEfightR can analyze any 5′-end data similar to CAGE, including nascent RNA 5′-end methods (Table 1).
This is highly relevant since nascent 5′-end methods may be more sensitive in terms of enhancer detection, and hence are often used specifically for this purpose. To illustrate the usefulness of CAGEfightR for analyzing such data, we applied CAGEfightR to 59 lymphoblastoid cell line Precision Nuclear Run-on Sequencing for RNA Polymerase II Start Sites (PRO-Cap) libraries from Katla et al [59]. Similarly to the analysis of ulcerative colitis CAGE data above, robust pooled CTSSs (CTSSs observed in > 2 samples) were used to identify TCs (> 1 TPM in > 5 samples) and enhancer candidates (> 0 tags in > 5 samples), and annotated these using UCSC transcript models. Compared to the CAGE libraries above, a larger number of antisense TCs, intergenic TCs and enhancer candidates were detected (Fig. 5a). This is expected as these RNAs are subject to nuclear degradation and thus are more difficult to detect with steady-state RNA methods. PRO-Cap TCs showed the expected pyrimidine-purine di-nucleotide at positions −1 +1 (Fig. 5b), and enhancer candidates exhibited the characteristic H3K4me3 to H3K4me1 ratio (Fig. 5c). Figure 5d shows an example of an enhancer candidate detected by PRO-Cap.
Conclusions

CAGEfightR is a user-friendly R-Bioconductor package for analyzing CAGE and 5′-end data. It includes a wide range of functions for analyzing TSSs and enhancers and generating publication-ready visualizations. CAGEfightR was designed from the ground up to adhere closely to Bioconductor standards, making it easy to learn, use and combine CAGEfightR with other Bioconductor packages for transcriptomic analyses. CAGEfightR is extensively documented, including both a vignette describing the core functionality (http://bioconductor.org/packages/CAGEfightR/) and a case-study based workflow discussing common analysis tasks on a full dataset [60] (http://bioconductor.org/packages/CAGEWorkflow/).

Materials and methods

Datasets

Analysis of all 5′-end data was based on supplied CTSSs from the GEO repositories of the respective papers. The Hela set [47] was obtained from GSE62047, using only EGFP samples. The ulcerative colitis set [25] was obtained from GSE95437, using only active ulcerative colitis (UCa) samples from the largest batch. The PRO-Cap set [59] was obtained from GSE110638, using only non-replicated samples.

Transcript and gene models were obtained from UCSC via the R/Bioconductor packages TxDb.Hsapiens.UCSC.hg19.knownGene and TxDb.Mmusculus.UCSC.mm9.knownGene. Chromatin data was obtained from the Roadmap Epigenomics project [61] via the AnnotationHub R/Bioconductor package, for Hela cells AH32877, AH32879, AH32881 and AH32884 and for lymphoblastoid cell lines AH32865, AH33899, AH33901 and AH33904.

Analysis

All analyses of 5′-end data were done using CAGEfightR as indicated in the main text. Average meta profiles were
made using the TeMPO R-package (https://github.com/MalteThodberg/TeMPO), removing the top 1% highest scoring features to dampen the effect of outliers. Sequence logos were done using the ggseqlogo R-package [62], genome-browser figures using Gviz [42] and remaining figures using ggplot2 (https://www.tidyverse.org/).

Andersson enhancers were predicted using scripts from the original publication [21] (https://github.com/anderssonrobin/enhancers). TCs used as input were defined by CAGEfightR with default settings. A balance cutoff of 0.6 was used, as this corresponds to the 0.95 balance cutoff used in CAGEfightR in the case of a BC with only divergent signal (PD and MD in Additional file 1: Figure S1A).

Availability and requirements

Project name: CAGEfightR.

Project home page: https://bioconductor.org/packages/release/bioc/html/CAGEfightR.html

Operating system(s): Platform independent (BigWig I/O only available on Windows).

Programming language: R.

Other requirements: Bioconductor.

License: GPL-3.

Any restrictions to use by non-academics: GPL-3.

Acknowledgements

We would like to acknowledge all members of the Sandelin and Andersson laboratories for discussions of countless aspects of 5′-end data analysis.

Authors’ contributions

MT and AS conceived the project and wrote the paper. MT implemented the package with input from AT. KV and RA provided input on statistics and validations. All authors read, commented on and approved the final manuscript.

Funding

This work was supported by the Novo Nordisk Foundation, the Lundbeck Foundation, Danish Cancer Society, Innovation Fund Denmark and Independent Research Fund Denmark. The funders played no role in the design of the study and collection, analysis, and interpretation of data or in writing the manuscript.

Availability of data and materials

The CAGEfightR R/Bioconductor package is freely available from http://bioconductor.org/packages/CAGEfightR/. All analyses are based on publicly available data obtained via GEO and AnnotationHub (See Materials and Methods for accession numbers).

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

Author details

1Department of Biology, University of Copenhagen, Ole Maaløes Vej 5, DK2100, Copenhagen N, Denmark. 2Biotech Research and Innovation Centre, University of Copenhagen, Ole Maaløes Vej 5, DK2100, Copenhagen N, Denmark. 3Danish Cancer Society, Strandboulevarden 49 DK2100, Copenhagen Ø, Denmark.

Received: 15 March 2019 Accepted: 15 August 2019

Published online: 04 October 2019

References

1. Lenhard B, Sandelin A, Caminici P. Metaazoan promoters: emerging characteristics and insights into transcriptional regulation. Nat Rev Genet. 2012;13(4):233–45.
2. Haberle V, Stark A. Eukaryotic core promoters and the functional basis of transcription initiation. Nat Rev Mol Cell Biol. 2018;19(10):621–37.
3. Sandelin A, Caminici P, Lenhard B, Ponjavic J, Hayashizaki Y, Hume DA. Mammalian RNA polymerase II core promoters: insights from genome-wide studies. Nat Rev Genet. 2007;8(6):424–36.
4. Adiconis X, Haber AL, Simmons SK, Levy Moonshine A, Ji Z, Busby MA, et al. Comprehensive comparative analysis of 5′-end RNA-sequencing methods. Nat Methods. 2018;15(7):505–11.
5. Takahashi H, Kato S, Murata M, Caminici P CAGE. (Cap analysis of gene expression): a protocol for the detection of promoter and transcriptional networks. Methods in molecular biology (Clifton, NJ). 2012;786:181–200.
6. Cumbie JS, Ivanchenko MG, Megraw M. NaroCAGE-XL and CapFilter: an approach to genome wide identification of high confidence transcription start sites. BMC Genomics. 2015;16:597.
7. Cvetesic N, Leitch HG, Borkowska M, Müller F, Caminici P, Hajkova P, et al. SLIC-CAGE: high-resolution transcription start site mapping using nanogram-levels of total RNA. Genome Res. 2018;28(12):1943–56.
8. Murata M, Nishiyori-Sueki H, Kojima-Ihijama M, Caminici P, Hayashizaki Y, Itoh M. Detecting expressed genes using CAGE. Methods in molecular biology (Clifton, NJ). 2014;1164:67–85.
9. Tsuchihara K, Suzuki Y, Wakaguri H, Irie T, Tanimoto K, Hashimoto S, et al. Massive transcriptional start site analysis of human genes in hypoxia cells. Nucleic Acids Res. 2009;37(7):2249–63.

Abbreviations

5′-3′ UTR: 5 prime / 3 prime untranslated region; BC: Bidirectional Cluster; bp / kb: Base pair / kilo base pairs; CAGE: Cap Analysis of Gene Expression; CDS: Coding sequencing; ChIP-Seq: Chromatin Immunoprecipitation Sequencing; CTSS: CAGE Transcription Start Sites; DNase-Seq: DNase I hypersensitive sites sequencing; GO: Gene Ontology; IQR: Interquartile Range; KEGG: Kyoto Encyclopedia of Genes and Genomes; PROCAP: Precision Nuclear Run-on Sequencing for RNA Polymerase II Start Sites; RNA-Seq: RNA-sequencing; TC: Tag Cluster or Unidirectional Cluster (closely spaced array of tags on the same strand); TPM: Tags-Per-Million; TSS: Transcription Start Site; UCSC: University of California Santa Cruz.

Additional file

Additional file 1: Figure S1. Details on finding Bidirectional Clusters (BCs). A: Calculating balance score using the Bhattacharyya coefficient. For a potential BC midpoint, pooled CTSS signal is summed within a certain distance (200 bp by default) on both strands, yielding four values (left). The “ideal” bidirectional cluster would have only perfect divergent signal (50% PD and 50% MD). The Bhattacharyya Coefficient quantifies the difference between the observed signal to this ideal enhancer (right), with a balance score of 1 indicating perfect agreement. The balance score is calculated for every bp in the genome (Fig. 1c). B: Overlap between CAGEfightR and Andersson enhancer predictions. The original enhancer prediction method from Andersson et al were applied to the Hela set. The venn diagram shows overlap in predictions between CAGEfightR, Andersson et al and DNase hypersensitive sites. CAGEfightR predicts all enhancers candidates form Andersson et al that are also supported by DNase hypersensitive sites. C: Chromatin modifications at CAGEfightR and Andersson predicted enhancer from the Hela set. X-axis shows distance to enhancer midpoint. Y-axis are average signal of respective DNase-Seq or ChIP-Seq data in the given panel row. Color indicates the enhancer candidate sets (Andersson in gold and CAGEfightR in grey), with panel columns indicating whether enhancer are shared between sets (left) or uniquely predicted (right). All sets exhibit the characteristic DNase hypersensitivity and H3K4me1/H3K4me3 ratio, despite the CAGEfightR enhancer candidate set is much larger. (PDF 840 kb)
61. Roadmap Epigenomics Consortium, Kundaje A, Meuleman W, Ernst J, Bilenky M, Yen A, et al. Integrative analysis of 111 reference human epigenomes. Nature. 2015;518(7539):317–30.
62. Wagih O. ggseqlogo: a versatile R package for drawing sequence logos. Bioinformatics (Oxford, England). 2017;33(22):3645–7.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.