Genome-wide circadian rhythm detection methods: systematic evaluations and practical guidelines

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Abstract
Circadian rhythms are oscillations of behavior, physiology and metabolism in many organisms. Recent advancements in omics technology make it possible for genome-wide profiling of circadian rhythms. Here, we conducted a comprehensive analysis of seven existing algorithms commonly used for circadian rhythm detection. Using gold-standard circadian and non-circadian genes, we systematically evaluated the accuracy and reproducibility of the algorithms on empirical datasets generated from various omics platforms under different experimental designs. We also carried out extensive simulation studies to test each algorithm’s robustness to key variables, including sampling patterns, replicates, waveforms, signal-to-noise ratios, uneven samplings and missing values. Furthermore, we examined the distributions of the nominal P-values under the null and raised issues with multiple testing corrections using traditional approaches. With our assessment, we provide method selection guidelines for circadian rhythm detection, which are applicable to different types of high-throughput omics data.

Key words: biological rhythm; circadian rhythm detection; benchmarking; omics; precision and recall; reproducibility

Background
Circadian rhythms are approximately 24-h oscillations of behavior, physiology and metabolism that exist in almost all living organisms ranging from prokaryotes to mammals [1, 2]. Circadian rhythm is regulated by the circadian system, which consists of many ‘clock-controlled genes’ that exhibit oscillatory patterns [1]. These oscillations provide organisms with an adaptive advantage by enabling them to predict and adjust to the variations within their environments [3]. Additionally, and perhaps more importantly, disruptions of circadian rhythms have shown to contribute to numerous diseases, including metabolic...
disorders, heart disease and aging [4–7]. It is, therefore, of great importance and interest to perform genome-scale analysis of biological rhythms.

Recent advances in omics technologies, including both microarrays and next-generation sequencing, offer appealing platforms to identify circadian genes on a genome-wide scale. These have, indeed, led to the proposal of multifarious methodologies adopted from various fields including mathematics, statistics, astrophysics, etc. The earliest of the selected methods is Lomb-Scargle (LS) periodogram [8], an algorithm adapted from astrophysics that detects oscillations by comparing the data to sinusoidal reference curves of varying periods and phases [9, 10]. ARSER is an algorithm that employs autoregressive spectral estimation to predict periodicity and applies a harmonic regression model to fit the time series [11]. Unlike the model-based LS and ARSER, JTK_CYCLE is a non-parametric method that detects oscillations by comparing the ranks of the measured values to a set of prespecified symmetric reference curves [3]. Both RAIN and eJTK_CYCLE build on the strengths of JTK_CYCLE: RAIN includes an additional set of asymmetric waveforms and examines the increasing and decreasing portions of the curve separately [12]; eJTK_CYCLE improves JTK_CYCLE by explicitly calculating the null distribution such that it accounts for multiple hypothesis testing and by including non-sinusoidal reference waveforms [13]. Based on the successes of the aforementioned methods, MetaCycle proposes an ensemble framework that integrates results from three different algorithms: LS, ARSER and JTK_CYCLE [14]. Specifically, MetaCycle detects periodicity using the best of breed methods: its P-values are generated using Fisher’s method, its periods and phase estimations are integrated using arithmetic and circular means and a new periodic model, formulated from ordinary least squares method, is applied to recalculate the amplitude. The most recent method, BIO_CYCLE, is a deep neural network trained on both simulated and empirical circadian and non-circadian time series [15]. More general information and characteristics of each method are summarized in Table 1.

Multiple studies [10, 16, 17] have evaluated the performance of different methods for circadian rhythm detection, showing discrepancies among the methods, whose performances depend on multiple factors including experimental designs, waveforms of interest, etc. However, there has not been, to our best knowledge, a comprehensive summary and evaluation of all existing methods to date. Here, we systematically assess the performance of the seven aforementioned algorithms for circadian rhythm detection: LS, ARSER, JTK_CYCLE, RAIN, eJTK_CYCLE, MetaCycle and BIO_CYCLE.

Specifically, we demonstrated and benchmarked the algorithms using real datasets with gold-standard circadian and non-circadian genes. All empirical data were generated using the liver tissue from Mus musculus that had undergone two different experimental designs. Under the dark–dark experimental design (24-h darkness), we focused on using data from gene expression microarrays to assess the accuracy and reproducibility of each algorithm; under the light–dark experimental design (12-h light followed by 12-h darkness), we adopted four different next-generation sequencing platforms and explored the robustness of each method in identifying circadian genes. Furthermore, to extend our assessment to non-transcriptomic datasets, we included a proteomic dataset in our evaluation. In addition, we carried out extensive simulation studies to study how key variables, including sampling patterns, replicates, waveforms, signal-to-noise ratios (SNRs), uneven samplings, missing values, affect the performance of each method. Lastly, we point out the flaw with using the Benjamini–Hochberg procedure to control for false discovery rate (FDR). Through these, we offer guidelines on experimental designs as well as best practices and methods of choice to increase the rigor and reproducibility in the analysis of large-scale circadian rhythms. To assist with the comparison of future methods and datasets using our framework, we provide detailed vignettes on applications of existing methods and performance evaluations with source code available at https://github.com/wenwenm183/Circadian_Genes_Benchmark.

**Results**

**Performance assessment using empirical datasets with dark–dark design**

We first adopted three gene expression microarray datasets from Hughes et al. [18], Hughes et al. [19] and Zhang et al. [20]. For all three studies, mouse liver samples were collected in every hour or every 2 h under the dark–dark experimental design for 48 h. We named these three datasets after the first author’s last name and the year of publication as Hughes 2009, Hughes 2012 and Zhang 2014. In addition, we generated a new downsampled dataset from the Hughes 2009 dataset by keeping the even time points only, and named it ‘Downsampled Hughes 2009.’ Refer to Table 2A for details of the data. Figure 1 shows the scaled gene expression levels of four known circadian and four non-circadian genes. The circadian genes, including the well-studied Clock, Cry1, Npas2 and Per1 [10], show oscillatory patterns that can be well reproduced across studies, while the non-circadian genes exhibit only noisy signals.

We set out to apply the seven algorithms to these four datasets to detect significantly cyclic genes and evaluate their performances using 104 circadian [10] and 113 non-circadian genes [21] from previous studies (Supplementary Table 1). The accuracy of each method in Hughes 2009, Downsized Hughes 2009, Hughes 2012 and Zhang 2014 was the first assayed with the precision and recall rates for each algorithm given three P-value thresholds, 0.000005 (Bonferroni), 0.00005, 0.0005 and one q-value threshold 0.05 (Benjamini–Hochberg). Due to the trade-off between sensitivity and specificity, with more relaxed thresholds of significance, the precision rates of all methods decrease while the recall rates increase—the 0.05 q-value threshold achieves the lowest precision rate yet the highest recall rate for any given method (Figure 2A). While there does not exist a single method that consistently achieves the highest precision or recall rate, JTK_CYCLE and BIO_CYCLE are more effective in controlling for false positives while still detecting true circadian genes. For the other methods, however, there is a much higher variability in precision, especially in the Zhang 2014 dataset (Figure 2A). RAIN and MetaCycle tend to have the highest sensitivity/recall, but this can come with significant sacrifice on precision (Figure 2A).

In addition, we find that higher sampling frequency can significantly improve the recall rates of all methods. While MetaCycle and RAIN achieve the apparently higher recall rate under different thresholds in dataset sampled at a lower frequency (2 h/2 days), all methods, except for LS, produce comparable recall rates when applied to the Hughes 2009 dataset, which is sampled at 1 h/2 days (Figure 2A). Notably, when analyzing the three datasets with lower sampling frequencies, LS failed under all circumstances with recall rates less than 0.1 (Figure 2A). This is due to the extreme P-value distribution of the method with a spike at one, which we will discuss in more detail under
Table 1. Summary of seven existing methods for circadian rhythm detection

| Package | Method key words | Method Type | Reference | Availability | Language | Replicates | Missing Values | Uneven Sampling |
|---------|------------------|-------------|-----------|--------------|----------|------------|----------------|-----------------|
| LS      | Periodogram      | Parametric  | Bioinformatics (2006) | https://www.iiap.res.in/astrostat/tuts/Lomb-Scargle.html | R | ✓✓✓ | ✓ | ✓ |
| ARSER (ARS) | Harmonic Regression | Parametric  | Bioinformatics (2010) | http://bioinformatics.cau.edu.cn/ARSE R | Python & R | ××× | × | × |
| JTK_CYCLE (JTK) | Kendall's Tau | Non-parametric | J Biol Rhythms (2010) | https://openwetware.org/wiki/HughesLab:JTK_Cycle | R | ✓✓✓ | ✓ | × |
| RAIN    | Asymmetric waveforms | Non-parametric | J Biol Rhythms (2014) | http://bioconductor.org/packages/rain | R | ✓✓✓ | ✓ | ✓ |
| eJTK_CYCLE (eJTK) | Empirical p-values | Non-parametric | PLOS Comp. Bio. (2015) | https://github.com/alanlhutchison/empirical-JTK_CYCLE-with-asymmetry | Python | ✓✓✓ | ✓ | ✓ |
| MetaCycle (MC) | Integration | Parametric  | Bioinformatics (2016) | https://cran.r-project.org/package=MetaCycle | R | ✓✓✓ | ✓ | ✓ |
| BIO_CYCLE (BC) | Deep Neural Network | Parametric  | Bioinformatics (2016) | http://circadiomics.igb.uci.edu | R | ✓✓✓ | ✓ | ✓ |

*BIO_CYCLE can be applied to datasets with missing values only if there are replicates and the missingness only pertains to part of the replicates.

Figure 1. Examples of circadian and non-circadian benchmark gene expressions among three datasets with dark–dark experimental design. Scaled gene expressions from selected (A) circadian genes including Clock, Cry1, Npas2 and Per1 and (B) non-circadian genes including Utp6, Mtf1, Cln3, Abcd4.

‘Correlated multiple testing and non-uniform distribution of P-values under the null’.

We further computed with the receiver operating characteristic (ROC) curves with a varying threshold on the nominal P-values returned by each method (Figure 2B). The areas under the curve (AUC) values serve as a joint measure of sensitivity and specificity and are above 0.80 across all benchmark results, suggesting that all methods achieve good sensitivities while controlling for false positive rates. BIO_CYCLE, the deep-learning-based method, achieves the best performance with the highest AUC across all datasets (Figure 2B).

Reproducibility assessment using empirical datasets with dark–dark design

Reproducibility is one of the core principles for any bioinformatic tools and yet it remains a challenge in the field of circadian rhythm detection, which has not been fully explored. To evaluate the reproducibility of the methods, we first compared and contrasted the significantly cyclic genes returned by each method across the four datasets. To make the input dimensions compatible, we selected a total of 7570 common genes that are shared across datasets and adopted a q-value threshold of 0.05 for significance. The Venn diagrams in Figure 3A show the overlapping...
relationships of the significant genes returned by each method. While the experimental designs are the same and the observed gene expression measurements are highly concordant (Figure 1), significant discrepancies of the calling results are observed. Of the seven benchmarked methods, ARSER resulted in 721 overlapping significant genes, which is the highest. This is followed by RAIN, eJTK_CYCLE, MetaCycle, BIO_CYCLE, JTK_CYCLE and LS with 613, 528, 485, 296, 204 and 0 mutually identified positives, respectively. As mentioned previously, LS failed in detecting any significant oscillations for three out of the four datasets.

To further assess the reproducibility of the methods, we computed the Jaccard index and the Sorensen index to measure the similarities among the results from each method. Details of these metrics are included in the Materials and Methods section. As a result, RAIN achieves one of the highest Jaccard indices for any pair of comparisons and ARSER achieves the highest overall Sorensen index across all datasets (Figure 3B). On the other hand, our results indicate that JTK_CYCLE and BIO_CYCLE produce the lowest similarity metrics across all comparisons (Figure 3B).

Performance assessment using empirical datasets with light–dark design

Next, we adopted four datasets that underwent light–dark experimental design using different next-generation sequencing platforms (i.e. RNA-seq [22], Nascent-seq [22], GRO-seq [23] and XR-seq [24]) and named each one after its sequencing protocol (Table 2B). The four datasets have much fewer numbers of time points compared to the datasets from the dark–dark design, yet three of the four datasets have technical replicates (Table 2B). More details of the data can be found in the Materials and Methods section. The oscillatory patterns of known circadian genes are apparent and similar among the various sequencing technologies (Figure 4A), indicating good data quality.

ARSER, despite its high reproducibility, cannot handle replicates, and previous studies have shown that data should never be concatenated [17]. Therefore, we focused on assessing the performance of the other six methods. We first examined the distribution of the nominal P-values of the 104 gold-standard circadian genes returned by each method, visualized as bee-hive plots in Figure 4B, where LS is significantly underpowered in the detection of circadian genes compared to the other methods, given any of the sequencing platforms. This result can be attributed to LS’s inability to effectively detect circadian rhythms in datasets with low sampling resolution, which is concordant with our previous results. We observe that JTK_CYCLE, RAIN, eJTK_CYCLE, MetaCycle and BIO_CYCLE can withstand the sparse sampling and result in overall good performance.

To further assess the performance of the methods, we examined the number of significant genes identified by each method with a FDR of 0.05. Of the 9481 mutual genes in the four datasets, LS did not identify any significant genes in any of the datasets. This result aligns with the results from the previous analysis, where we observed LS as being underpowered. JTK_CYCLE and MetaCycle detected a relatively small number of significant genes by RNA-seq and XR-seq. eJTK_CYCLE identified 2623 significant genes by RNA-seq, and RAIN and BIO_CYCLE identified 2262 and 1970 significant genes by XR-seq, respectively. When comparing across different sequencing
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Figure 3. Evaluation of method reproducibility. (A) Venn diagrams display the number of cyclic genes that are significant by each method among the four dark–dark datasets. (B) Jaccard index and the Sorensen index are used as metrics for reproducibility for each method across the four datasets with the same experimental design.

Performance assessment using empirical proteomic dataset of dark–dark design

To assess performance of the various methods on non-transcriptomic data, we adopted a proteomic dataset of mouse livers under dark–dark experimental design from Robles et al. [27]. Refer to the Materials and Methods section for details. Since this dataset consists of replicates and missing values, only LS, JTK_CYCLE, RAIN and MetaCycle were directly applicable. eJTK_CYCLE was not included due to its inefficiency in handling random missing values across different genes/proteins. We calculated the number of significant proteins identified by each method using an FDR threshold of 0.05 (Supplementary Figure 1A). LS identified the least number of oscillatory proteins. JTK_CYCLE and MetaCycle returned a moderate number of significant proteins. RAIN identified the largest number of oscillatory proteins, 582, exceeding that of other methods by more than 300. Heatmaps of scaled measurements of oscillatory
proteins identified by at least two methods are shown in Supplementary Figure 1B, where the proteins are ordered based on their inferred phases. With the identified oscillatory proteins, we conducted a gene set enrichment analysis using the DAVID web server. While the results did not indicate that circadian rhythm was significantly enriched by any of the algorithms, KEGG metabolic pathways were significantly enriched by all algorithms but LS (Supplementary Table 3).

**Performance assessment using synthetic datasets**

To provide guidelines for method selection, we evaluated the performance of the seven methods in detecting circadian rhythm by simulations with known ground truths. Examples of waveforms generated for the simulated datasets are shown in Supplementary Table 4. We generated six groups of simulated datasets to investigate how key factors affect the performance, including sampling patterns, replicates, waveforms, SNRs, uneven samplings and missing values. Supplementary Table 5 outlines the six groups of simulations and we leave the detailed setup in the Materials and Methods section. Within each simulation group, we repeated each assessment with three different sampling frequencies to determine whether increasing sampling frequency may have an effect on the aforementioned factors. The three sampling frequencies include 4 h/1 (six time points), 3 h/1 (eight time points) and 2 h/1 day (12 time points) and the results are shown in Figure 5A–C, respectively.

**Sampling patterns**

To determine whether increasing the sampling frequency or lengthening the time-window is more important for each method, we first evaluated the results under the sampling pattern of 4 h/1 day versus 8 h/2 days, 3 h/1 day versus 6 h/2 days and 2 h/1 day versus 4 h/2 days. We did not find strikingly different results within each pair of comparison, indicating that when the total number of data points are fixed, having a denser sampling density and enlarging the sampling time-window tend to have similar impact on performance. However, when we increase the number of data points, the performances of all methods are improved, which is concordant with existing studies [16, 17]. BIO_CYCLE generally outperforms the other methods, especially in datasets with lower sampling frequency and shorter time-window, while JTK_CYCLE is the most sensitive to fewer observations.

**Replicates**

To investigate the trade-off between replicates and sampling frequency, we compared the results of higher sampling frequency without replicates to those of lower sampling frequency with replicates. We first compared the dataset sampled at 4 h/1 day X1 to the dataset sampled at 8 h/1 day X2. LS, JTK_CYCLE, RAIN, eJTK_CYCLE and MetaCycle show better performance with replicates, while BIO_CYCLE performs significantly better on densely sampled datasets without replicates. Similar results are seen when we applied the methods to the dataset at 3 h/1 day without replicates and the dataset at 6 h/1 day with replicates. As expected, further increasing the sampling resolution offsets the existing preferences that the methods have for inclusion of replicates or higher sampling density.

**Waveforms**

Supplementary Table 4 outlines the different types of periodic waveforms that we generated in silico in three broad categories: stationary, non-stationary and asymmetric ones. Through our simulations, we find that all of the algorithms perform the best in detecting non-stationary waveforms. Additionally, all methods, with the exception of eJTK_CYCLE, perform better on stationary waveforms, compared to asymmetric waveforms. eJTK_CYCLE and RAIN are the top two methods for identifying asymmetric waveforms, which are expected due to their design. This is followed by LS, BIO_CYCLE, MetaCycle and ARSER. JTK_CYCLE is the least effective in identifying asymmetric waveforms regardless of sampling frequency.
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Figure 5. Performance assessment via simulation studies. Seven circadian rhythm detection methods are evaluated under different experimental designs to explore how sampling patterns, replicates, waveforms, SNRs, uneven samplings and missing values affect performance. Simulations under each design are carried out with different sampling frequencies: (A) 4 h/1 day, (B) 3 h/1 day and (C) 2 h/1 day. AUC values calculated from ground truths are used as metrics.

Signal-to-noise ratios (SNRs)

To test the effects of different noise levels on method performance, we generated various datasets with SNRs of 3, 2, 1 and 0.5. For all methods, our results suggest that the larger the SNRs, the higher the accuracy, as expected. LS, MetaCycle and BIO_CYCLE are overall the most robust to noises regardless of sampling frequency, while JTK_CYCLE has the poorest performance given high noise levels.

Uneven samplings

To understand how well the methods deal with uneven samplings, we focus on the results of datasets with one or more uneven time points. Our results suggest that BIO_CYCLE and LS/MetaCycle outperform the other two compatible methods. Under a sparse sampling design, RAIN and eJTK_CYCLE suffer significantly from an increasing number of uneven samplings; a dense sampling design, on the other hand, rescues the aforementioned methods.

Missing values

We generated datasets that contain 1, 5 and 10% missing data, and benchmarked the four methods that allow missing values. The performances of eJTK_CYCLE and RAIN degrade with an increasing proportion of missing values, while the performances of LS, JTK_CYCLE and MetaCycle are comparably invariant, especially under dense sampling design. We note that eJTK_CYCLE does not handle missing values efficiently, unless the same sampling time points are missing across all genes, which reduces to uneven sampling. When there is not a shared missing pattern across different genes, the dataset needs to be split into multiple uneven sampling cases, and eJTK_CYCLE needs to be applied separately, followed by results integration. Note that BIO_CYCLE can be applied to datasets with missing values only if there are replicates and the missingness only pertains to part of the replicates. We therefore did not include it in the benchmark.

Computational efficiency

Last but not least, we evaluated the computational efficiency across all benchmarked methods. For dataset with low sampling resolution, the execution times among the methods are approximately the same (Supplementary Table 6). However, when analyzing data of larger sizes, RAIN requires significantly more time compared to the other methods. The running time for LS, ARSER and MetaCycle, which integrates results from LS, JTK_CYCLE and ARSER, is calculated as the total running time of the three methods.

Correlated multiple testing and non-uniform distribution of $P$-values under the null

To detect circadian rhythm across thousands of genes, multiple hypothesis testing corrections are needed [28]. A common FDR threshold of 0.05 is recommended by most methods and adjusted $P$-values ($q$-values) are returned by all methods except for RAIN. In the previous sections, we adopted both Bonferroni and Benjamini–Hochberg procedures for corrections. Here, we more carefully examine such procedures and point out a potential drawback resulted from both correlated multiple testing and non-uniform distributions of the nominal $P$-values under the null. We started with the observed expression measurements from the Hughes 2009 dataset and generated a 'null' dataset by randomly permuting the time labels for each gene (Figure 6A).
Table 2. High-throughput mouse liver datasets adopted for circadian rhythm detection

| Design          | Name                        | Reference          | Accession number | Tissue type | Sequencing platform | Number of time points & replicates |
|-----------------|-----------------------------|--------------------|------------------|-------------|---------------------|-----------------------------------|
| (A) Dark-Dark   | Hughes et al.               | PLOS Genetics (2009) | GSE11923         | Liver       | Microarray          | 48 x 1                            |
| (B) Light-Dark  | Hughes et al.               | PLOS Genetics (2012) | GSE29411         | Liver       | Microarray          | 24 x 1                            |
|                 | Zhang et al.                | CAL (2014)         | GSE29405         | Liver       | Microarray          | 24 x 1                            |
|                 | Pang et al.                 | eLIFE (2012)       | GSE29467         | Liver       | Microarray          | 6 x 2                             |
|                 | Menet et al.                | eLIFE (2012)       | GSE29487         | Liver       | Nascent-seq         | 6 x 2                             |
|                 | Yang et al.                 | PNAS (2018)        | GSE109938        | Liver       | XR-seq (TS)         | 6 x 2                             |

Discussion

Here, we propose a benchmark framework to systematically evaluate the performance of seven circadian rhythm detection methods, using high-throughput omics data. The empirical datasets that we adopted in this paper were from microarray [18–20] and RNA-seq [22] to measure gene expression, Nascent-seq [22] and GRO-seq [23] to measure nascent RNA and XR-seq [24] to measure transcription-coupled repair. While these omics data were generated from different platforms, they focus on directly or indirectly profiling transcription. It has been well studied that biological rhythm goes beyond the transcriptomic transcript-level oscillations [34]. For example, posttranslational protein acetylation has been linked to circadian rhythm via mass spectrometry [35, 36]. Moreover, it has been shown that a large number of metabolites and proteins exhibit circadian oscillations [27, 37, 38]. The methods and the evaluation procedures are not limited to transcriptomic studies, but can also be applied to acetylic, metabolomic and proteomic experiments.

Given the assessment results from both simulations and empirical dataset analysis, as well as literature review of the seven methods, we have summarized the strengths and weaknesses of each method in Table 3. In general, LS, RAIN, eJTK_CYCLE and MetaCycle are more versatile in that they can be applied to datasets with replicates, uneven samplings or missing values. eJTK_CYCLE and BIO_CYCLE generally outperform the other methods under most situations except for handling missing values. On the other hand, JTK is sensitive to high noise levels and low sampling resolutions, and LS cannot detect any significant genes when sampling resolution is lower than...
Figure 6. Existing methods return non-uniformly distributed $P$-values under the null, partially due to non-independent testing due to gene-gene correlations. (A) Gene expression values for the benchmark circadian gene Cry1 before and after random permutations of the time labels. (B) Heatmaps of pairwise correlation coefficients among the top 200 highly variable genes from the Hughes 2009 dataset. The top illustrates the gene-gene correlation coefficients calculated from raw data input, and the bottom shows the gene-gene correlations after permutation. (C) The distributions of nominal $P$-values for each method when applied to the dataset before and after permutation. Gene-gene correlations, which are accounted for by eJTK_CYCLE, partially lead to the systematic deviations from the null distributions. The hypothesis testing by LS, JTK_CYCLE, RAIN and MetaCycle are overly conservative, while ARSER’s and BIO_CYCLE’s testing procedures are biased with an overabundance of $P$-values around 0.3 and 0.1, respectively, under the null.

2 h/2 days with an FDR threshold of 0.05. The best detection algorithm depends on experimental designs and characteristics of the input data. Therefore, we have created two decision trees, one for low sampling resolution and the other for high sampling resolution, that outline the recommended method(s) under different scenarios (Supplementary Figure 2).

Recent advances of high-throughput technologies enable circadian rhythm detection on the genome-wide scale. As with all genomic data, the multi-time-point omics data for circadian rhythm detection bear both technical and biological variability, which can bias the analysis if not properly accounted. Data normalization and batch effect correction are crucial to remove technical biases and artifacts [39]. Cross-subject variability in rhythmic profiles, especially for human subjects, is a non-negligible source of genetic variation that needs to be adjusted [14]. This is especially important in the case-control setting where multiple subjects are involved. While we did not particularly focus on differential analysis since it is outside the scope of this paper, a few methods, including LimoRhyde [40] and DODR [12] have been made available for differential rhythmicity analysis under different conditions.

Increasingly more circadian omics data are being made available through existing studies and databases [34, 41]. We showed from our empirical studies, that the rhythmic signals can be well recapitulated across different studies and/or different platforms (Figures 1 and 4A). Meta-analysis and multi-omics data integration remain an open-ended question in circadian rhythm detection [42]. In addition, transfer learning has been applied to multiple genomic research domains in genomics [43]—to borrow information and to transfer knowledge from existing data deposited in public repositories remain one of the future directions. Similarly, across different methods, an ensemble framework, as implemented by MetaCycle, can potentially boost performance. However, as we have pointed out earlier, the instability issue needs to be addressed, especially when multiple drastically distinct results are to be integrated.

To our best knowledge, all existing studies for circadian rhythm detection resort to bulk-tissue omics data, which characterize an averaged profile across different cell types in a tissue. The inherent heterogeneity can bias the analysis with reduced power and/or inflated FDR. Single-cell sequencing circumvents the averaging artifacts associated with traditional bulk population data and has seen rapid technological developments over the past few years. To assess the feasibility of single-cell circadian rhythm detection, we in silico generated single-cell RNA sequencing profiles by downsampling bulk RNA-seq read counts. Gold-standard circadian and non-circadian genes were used to calculate the associated AUC values (Supplementary Table 7). All methods suffer from low sequencing depth—a characteristic of the single-cell data. With the decreasing cost and the increasing popularity of single-cell omics techniques, to profile circadian rhythmicity at the cellular level and to disentangle within tissue heterogeneity with regard to biological rhythm can be of great impact.

Materials and Methods

Empirical transcriptomic datasets

Three datasets under the dark-dark experimental design including Hughes [18], Hughes [19] and Zhang [20] were
assays exons [44]. XR-seq profiles DNA excision repair on the sequence and Nascent-seq differ from traditional RNA-seq, in which and evaluating transcriptionally engaged polymerase [45]. GRO-seq measures nascent RNAs by mapping, characterizing, genome-wide scale with single-nucleotide resolution [46]. Here, gene expressions (Table 2A). Additionally, we obtained four datasets under the light–dark design from the different sequencing platforms, including Nascent-sequencing (Nascent-seq) [22], RNA-sequencing (RNA-seq) [22], Global Run-On sequencing (GRO-seq) [23] and eXcision Repair-sequencing (XR-seq) [24] (Table 2B). Nascent-seq sequence transcribed RNAs, obtained from the nuclei without formation of the 3′ end [44]. GRO-seq measures nascent RNAs by mapping, characterizing, and evaluating transcriptionally engaged polymerase [45]. GRO-seq and Nascent-seq differ from traditional RNA-seq, in which the reads map to predominantly introns, while RNA-seq mainly assesses exons [44]. XR-seq profiles DNA excision repair on the genome-wide scale with single-nucleotide resolution [46]. Here, we focus on XR-seq data from the transcribed strand only—it has been shown that the transcription-coupled repair from the transcribed strand is positively correlated with expression [47].

For quality control, we removed genes that had constant gene expression measurements in all datasets and further removed genes with more than half zero gene expression values in the light–dark datasets. In cases where multiple probes got mapped to the same RefSeq loci, we averaged the gene expression of the probes using the limma package [48], available in Bioconductor. For data normalization, robust multi-array average (RMA) [49] and genechip RMA (GC-RMA) [50] were used to normalize the array data; transcript per million and reads per kilobase per million reads (RPKM) [51] were used to normalize the transcriptomic sequencing data. We scaled the normalized data within each gene to make them compatible for visualization only, as shown in Figures 1 and 4A.

### Table 3. Pros and cons of circadian rhythm detection methods

| Methods | Pros | Cons |
|---------|------|------|
| LS      | • Effective in handling missing data • Can be applied to datasets with replicates, uneven samplings, and missing data | • Highly sensitive to low sampling resolution • U-shaped P-values distribution • Sensitive to outliers |
| ARSER  | • High reproducibility | • Cannot handle replicates, uneven samplings, or missing data |
| JTK_CYCLE | • High precision • Robust to outliers | • Incapable of detecting asymmetric waveforms • U-shaped P-values distribution • Low reproducibility |
| RAIN    | • High recall • Effective in detecting asymmetric waveforms • High reproducibility • Can be applied to datasets with replicates, uneven samplings, and missing data | • High false positive rates U-shaped P-values distribution Computationally intensive with increasing sampling resolution |
| eJTK_CYCLE | • Uniform distribution of nominal P-values • Most effective in detecting asymmetric waveforms | • Unable to test different periods simultaneously • Inefficient in handling missing data • Sensitive to high level of uneven samplings |
| MetaCycle | • High recall • Offset the disadvantages of one method with the other two among LS, ARSER and JTK_CYCLE via ensemble • Directly return calling results from three perspective methods | • P-values generated with Fisher’s integration require independence assumption U-shaped P-values distribution |
| BIO_CYCLE | • Most effective in controlling for false positive rates • Most robust to data with uneven samplings, and low sampling resolutions. • High precision | • Require extensive time and in silico generated data to train the DNN model • Handle missingness only if data have replicates and the missingness only pertains to part of the replicates |

### Empirical proteomic dataset

A proteomic dataset of *Mus musculus* liver tissues from Robles et al. [27] was adopted to detect oscillatory proteins. Mouse liver samples were collected from a total of 64 mice that were released into constant darkness for 1 day after being entrained to a 12–12 h light–dark schedule for 10 days. Four mice were sacrificed every 3 h for 2 days. Then, in vivo stable isotope labeling by amino acids in cell culture (SILAC) [52, 53] in combination with mass spectrometry was performed to profile the proteome. For each time point, equal amount of protein liver extracts from the four mice were mixed together with equal amount of protein lysates, collected in antiphase, from the liver samples of two SILAC mice. The pooled protein extracts were measured with Orbitrap mass spectrometry. The protein abundance was calculated by taking the ratio of the signal for the mice and the signal for the heavy SILAC mix. After assessing quantification values, a total of 3132 proteins remained for downstream circadian rhythm analysis.

### Downsampling RNA-seq dataset

We generated several downsampling RNA-seq datasets from the original RNA-seq dataset under the light–dark design to assess the robustness of the various methods to low sequencing depths. We obtained the raw sequencing data from GEO, performed read alignment to the mouse reference genome (mm10) using STAR [54], carried out quality control procedures on the aligned reads, and obtained integer-valued read counts using featureCounts [55]. We then generated downsampling RNA-seq data by multinomial sampling with index 5 K, 10 K, 50 K, 100 K and 500 K, and gene-specific probability parameters calculated from the raw
data. RPKM was used to normalize the downsampled RNA-seq read counts, followed by circadian rhythm detection.

**Evaluation metrics**

To evaluate the performance of the benchmarked methods, we adopted a list of 104 circadian [10] and 113 non-circadian genes [21] in mouse liver as positive and negative controls, respectively. See Supplementary Table 1 for a full list of these gold-standard genes. With these gold-standard genes, we calculated metrics including the precision and recall rates given a P-value or q-value significance threshold (Figure 2A). We further calculated the AUC values of the ROC curves, as joint measures of sensitivity and specificity (Figure 2B).

To assess the reproducibility of each method, we compared the results from the four dark-dark datasets by calculating the number of overlapping genes, as well as the Jaccard and Sorensen index as metrics for similarity (Figure 3). Venn diagrams are used to display the number of overlapping cycling genes identified across different datasets by each method. The Jaccard index measures the pairwise similarities of the significant genes detected between each pair of datasets. Let $A_i$ and $A_j$ be the set of significant genes from dataset $i$ and $j$. The Jaccard similarity index is defined as follows:

$$J(A_i, A_j) = \frac{|A_i \cap A_j|}{|A_i \cup A_j|}$$

The Sorensen index is used to characterize similarity across all datasets [56]:

$$S(A_i, A_j, A_k, \ldots) = \frac{T}{T-1} \left( \sum_{i<j} |A_i \cap A_j| - \sum_{i<j<k} |A_i \cap A_j \cap A_k| + \sum_{i<j<k<l} |A_i \cap A_j \cap A_k \cap A_l| - \ldots \right)$$

where $T$ is the number of sets compared. Larger number of overlapping genes and larger Jaccard/Sorensen index values indicate higher reproducibility of the methods.

**Simulation setup**

Each simulated dataset consists of 6000 circadian and 6000 non-circadian gene profiles. Stationary circadian profiles with a period of 24 h are used in each simulation group, as outlined below. Note that when running the methods, we set the period range from 20 to 28 h for all methods except for eJTK_CYCLE and JTK_CYCLE, which either has a fixed period of 24 h or adjusts the period on the fly. The amplitude of the waveforms is sampled from a uniform distribution between 1 and 6; the phase shift is sampled from a uniform distribution between 0 and 24 h; and the noise term is sampled from a standard normal distribution. Flat waveforms are used to generate non-circadian profiles in all simulation groups except for testing against non-stationary waveforms where linear lines are used.

We first aimed to investigate whether higher sampling frequency or longer sampling time-window is more beneficial for each method. In this simulation group, we generated two datasets with different sampling frequencies and sampling time-windows. With six time points, we generated one dataset at 4 h/1 day and another at 8 h/2 days; with eight time points, we generated one dataset at 3 h/1 day and another at 6 h/2 days; with 12 time points, we generated one dataset at 2 h/1 day and another at 4 h/2 days.

Next, we assessed whether the inclusion of replicates can offset the effect of low sampling frequency in methods’ ability of detecting oscillations. Replicates are defined as multiple measurements taken at the same time point. Specifically, we generated two datasets consisting of the same number of observations, with or without replicates: one at 4 h/1 day X1 and the other at 8 h/1 day X2. The sampling designs of the other two pairs of datasets are 3 h/1 day X1 versus 6 h/1 day X2 and 2 h/1 day X1 versus 4 h/1 day X2.

Since biological rhythms can take on various waveforms, we generated three types of waveforms via simulation: stationary, non-stationary and asymmetric curves. Supplementary Table 4 includes models that we adopted in silico to generate the corresponding waveforms. Specifically, the stationary waveforms include cosine, cosine 2 and cosine peak curves; the non-stationary waveforms include cosine damp, trend exponential and trend linear curves; the asymmetric subgroup consists of only the saw-tooth waveform. We assessed the performance of the methods in identifying each category of the circadian waveforms.

The next three groups of simulations aimed to determine which methods are more robust to different levels of SNRs, uneven samplings and missing values. Specifically, we generated four datasets with SNRs of 0.5, 1, 2 and 3. SNR is defined by taking the ratio of the empirical variance of cosine function and the variance of the noise, the latter of which is fixed at one. Uneven samplings are defined as designs whose time points are not equally spaced. To investigate the effect of uneven samplings on performance, we generated datasets with one, two or four uneven samplings. With six time points, datasets with four uneven samplings cannot be generated as it would only have two time points. For missing data, we generated three levels of missing data (1, 5 and 10%) at three fixed, randomly selected time points.

Lastly, we generated three datasets with sampling patterns of 1 h/2, 2 h/2 and 4 h/2 days to compute the execution times for each method. We seek to identify the differences in computational efficiency among the methods and to explore the effect of increasing sampling resolution on the execution time. Each dataset consists of a total of 6000 genes. All execution times are reported by running on a Macbook Pro (15-inch, 2019) with 2.3 GHz 8-Core Intel Core i9 and 16 GB memory.

**Data and software availability**

MetaCycle is an open-source R package available at https://github.com/gangwug/MetaCycle and is also used for individual analysis for LS, JTK_CYCLE and ARSER. RAIN is a Bioconductor R package available at https://bioconductor.org/packages/rain/. eJTK_CYCLE was downloaded from https://github.com/alanhuutchison/empirical-JTK_CYCLE-with-asymmetry. Bio_CYCLE was downloaded from http://circadiomics.igb.uci.edu/BIO_CYCLE. All empirical datasets were downloaded from the NCBI Gene Expression Omnibus (https://www.ncbi.nlm.nih.gov/geo/). The accession numbers for dark–dark datasets are GSE11923, GSE30411 and GSE54652, respectively. The accession numbers for light–dark datasets are GSE59486, GSE36872, GSE36871 and GSE109938, respectively. The proteomic dataset
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**Key Points**

- Various methods have been developed for circadian rhythm detection on a genome-wide scale using omics technologies, yet there has not been a comprehensive summary and evaluation of all existing methods to date.
- Using gold-standard circadian and non-circadian genes, we systematically evaluated the accuracy and reproducibility of seven existing algorithms for circadian rhythm detection on empirical datasets generated from various omics platforms.
- We carried out extensive simulation studies to test each algorithm’s robustness to key variables, including sampling patterns, replicates, waveforms, signal-to-noise ratios, uneven samplings and missing values.
- We examined the distributions of the nominal P-values under the null and raised issues with multiple testing corrections using the Benjamini–Hochberg procedure due to gene-gene correlation and testing being overly conservative.
- We provide method selection guidelines for circadian rhythm detection, which are applicable to different types of high-throughput omics data.

**Supplementary Data**

Supplementary data are available online at https://academic.oup.com/bib.

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**References**

1. Li J, Grant GR, Hogenesch JB, et al. Considerations for RNA-seq analysis of circadian rhythms. *Methods Enzymol* 2015;551:349–67.
2. Panda S, Antoch MP, Miller BH, et al. Coordinated transcription of key pathways in the mouse by the circadian clock. *Cell* 2002;109:307–20.
3. Hughes ME, Hogenesch JB, Kornacker K. JTK_CYCLE: an efficient nonparametric algorithm for detecting rhythmic components in genome-scale data sets. *J Biol Rhythms* 2010;25:372–80.
4. Asher G, Sassone-Corsi P. Time for food: the intimate interplay between nutrition, metabolism, and the circadian clock. *Cell* 2015;161:84–92.
5. Partch CL, Green CB, Takahashi JS. Molecular architecture of the mammalian circadian clock. *Trends Cell Biol* 2014;24:90–9.
6. Roenneberg T, Merrow M. The circadian clock and human health. *Curr Biol* 2016;26:R432–43.
7. Levi F, Schibler U. Circadian rhythms: mechanisms and therapeutic implications. *Annu Rev Pharmacol Toxicol* 2007;47:593–628.
8. Glynn EF, Chen J, Mushegian AR. Detecting periodic patterns in unevenly spaced gene expression time series using Lomb-Scargle periodograms. *Bioinformatics* 2006;22:310–6.
9. Wijnen H, Naef F, Young MW. Molecular and statistical tools for circadian transcript profiling. *Methods Enzymol* 2005;393:341–65.
10. Wu G, Zhu J, Yu J, et al. Evaluation of five methods for genome-wide circadian gene identification. *J Biol Rhythms* 2014;29:231–42.
11. Yang R, Su Z. Analyzing circadian expression data by harmonic regression based on autoregressive spectral estimation. *Bioinformatics* 2010;26:i168–74.
12. Thaben PF, Westermark PO. Detecting rhythms in time series with RAIN. *J Biol Rhythms* 2014;29:391–400.
13. Hutchison AL, Maienschein-Cline M, Chiang AH, et al. Improved statistical methods enable greater sensitivity in rhythm detection for genome-wide data. *PLoS Comput Biol* 2015;11:e1004094.
14. Wu G, Anafi RC, Hughes ME, et al. MetaCycle: an integrated R package to evaluate periodicity in large scale data. *Bioinformatics* 2016;32:3351–3.
15. Agostinelli F, Ceglia N, Shabbaba B, et al. What time is it? Deep learning approaches for circadian rhythms. *Bioinformatics* 2016;32:i8–17.
16. Deckard A, Anafi RC, Hogenesch JB, et al. Design and analysis of large-scale biological rhythm studies: a comparison of algorithms for detecting periodic signals in biological data. *Bioinformatics* 2013;29:3174–80.
17. Hughes ME, Abruzzi KC, Allada R, et al. Guidelines for genome-scale analysis of biological rhythms. *J Biol Rhythms* 2017;32:380–93.
18. Hughes ME, DiTacchio L, Hayes KR, et al. Harmonics of circadian gene transcription in mammals. *PLoS Genet* 2009;5:e1000442.
19. Hughes ME, Hong HK, Chong JL, et al. Brain-specific rescue of clock reveals system-driven transcriptional rhythms in peripheral tissue. *PLoS Genet* 2012;8:e1002835.
20. Zhang R, Lahens NF, Ballance HI, et al. A circadian gene expression atlas in mammals: implications for biology and medicine. *Proc Natl Acad Sci U S A* 2014;111:16219–24.
21. Wu G, Zhu J, He F, et al. Gene and genome parameters of mammalian liver circadian genes (LCGs). *PLoS One* 2012;7:e46961.
22. Menet JS, Rodriguez J, Abruzzi KC, et al. Nascent-Seq reveals novel features of mouse circadian transcriptional regulation. *Elife* 2012;1:e00011.
23. Fang B, Everett LJ, Jager J, et al. Circadian enhancers coordinate multiple phases of rhythmic gene transcription in vivo. *Cell* 2014;159:1140–52.
24. Yang Y, Adebalu O, Wu G, et al. Cisplatin-DNA adduct repair of transcribed genes is controlled by two circadian programs in mouse tissues. *Proc Natl Acad Sci USA* 2018;115:E4777–85.
25. Huang d W, Sherman BT, Lempicki RA. Systematic and integrative analysis of large gene lists using DAVID bioinformatic resources. *Nat Protoc* 2009;4:44–57.
26. Huang d W, Sherman BT, Lempicki RA. Bioinformatics enrichment tools: paths toward the comprehensive functional analysis of large gene lists. *Nucleic Acids Res* 2009;37:1–13.
27. Robles MS, Cox J, Mann M. In-vivo quantitative proteomics reveals a key contribution of post-transcriptional mechanisms to the circadian regulation of liver metabolism. PLoS Genet 2014;10:e1004047.
28. Storey JD, Tibshirani R. Statistical significance for genomewide studies. Proc Natl Acad Sci USA 2003;100:9440–5.
29. Li J, Ji L. Adjusting multiple testing in multilocus analyses using the eigenvalues of a correlation matrix. Heredity (Edinb) 2005;95:221–7.
30. Conneely KN, Boehnke M. So many correlated tests, so little time! Rapid adjustment of P values for multiple correlated tests. Am J Hum Genet 2007;81:1158–68.
31. Schwartzman A, Lin X. The effect of correlation in false discovery rate estimation. Biometrika 2011;98:199–214.
32. Hutchison AL, Dinner AR. Bootstrapping and empirical Bayes methods improve rhythm detection in sparsely sampled data. J Biol Rhythms 2018;33:339–49.
33. Ceglia N, Liu Y, Chen S, et al. CircadiOmincs: circadian omic web portal. Nucleic Acids Res 2018;46:W157–62.
34. Masri S, Patel VR, Eckel-Mahan KL, et al. Circadian acetylome reveals regulation of mitochondrial metabolic pathways. Proc Natl Acad Sci USA 2013;110:3339–44.
35. Mauvoisin D, Atger F, Dayon L, et al. Circadian and feeding rhythms orchestrate the diurnal liver Acetylome. Cell Rep 2017;20:1729–43.
36. Dallmann R, Viola AU, Tarokh L, et al. The human circadian metabolome. Proc Natl Acad Sci USA 2012;109:2625–9.
37. Feng D, Lazar MA. Clocks, metabolism, and the epigenome. Mol Cell 2012;47:158–67.
38. Jiang Y, Wang R, Urrutia E, et al. CODEX2: full-spectrum copy number variation detection by high-throughput DNA sequencing. Genome Biol 2018;19:202.
39. Singer JM, Fu DY, Hughey JJ. Symphony: simulating large-scale, rhythmic data. PeerJ 2019;7:e6985.
40. Li X, Shi L, Zhang K, et al. CircGRDB: a database for the genome-wide deciphering circadian genes and regulators. Nucleic Acids Res 2018;46:D64–70.