Problem Solving Protocol

Sensitive, reliable and robust circRNA detection from RNA-seq with CirComPara2

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Abstract
Circular RNAs (circRNAs) are a large class of covalently closed RNA molecules originating by a process called back-splicing. CircRNAs are emerging as functional RNAs involved in the regulation of biological processes as well as in disease and cancer mechanisms. Current computational methods for circRNA identification from RNA-seq experiments are characterized by low discovery rates and performance dependent on the analysed data set. We developed CirComPara2 (https://github.com/enrico-gaffo/CirComPara2), a new automated computational pipeline for circRNA discovery and quantification, which consistently achieves high recall rates without losing precision by combining multiple circRNA detection methods. In our benchmark analysis, CirComPara2 outperformed state-of-the-art circRNA discovery tools and proved to be a reliable and robust method for comprehensive transcriptome characterization.

Key words: circRNAs; bioinformatics; computational pipeline; RNA-seq

Introduction
Recent research uncovered that eukaryotic transcriptomes comprise thousands of stable circular RNAs (circRNAs) originating by a process called back-splicing, where the transcript 3' and 5' ends are covalently joined [1]. Rather than being transcriptional by-products, circRNA molecules exert critical functions in cell biology through different mechanisms [2]. By interacting with microRNAs, circRNAs can regulate gene expression and govern important oncogenic axes [3]; moreover, similar to long non-coding RNAs, they can control diverse cellular processes by decaying RNA-binding proteins and scaffolding molecular complexes [4]. CircRNAs can also function as templates for translation to encode functional peptides [1, 5] and regulate the transcription of their parental gene [6]. Nowadays, circRNAs are considered key players that can impact all cancer hallmarks [7, 8]. The discovery of circRNA regulatory roles and their potential as biomarkers given by higher stability compared to linear RNAs [9] has actuated the integration of circRNA investigation in conventional transcriptomics, especially in cancer research and studies of pathological conditions [10, 11], including viral infections [12].

Studies of circRNAs rapidly increased in pace thanks to the development of bioinformatics tools that identify the sequences spanning circRNA back-splice junctions from total RNA-seq data. To date, several methods for circRNA identification have been developed [13]; most of them select the back-splice junction reads (BJR) by screening the output of read aligner tools that allow chimeric spliced read mappings to the reference
circRNA discovery, including circRNA_finder [29], CIRI2 [27], such. We applied nine widely used computational pipelines for negative predictions (FNs), i.e. true circRNAs not identified as evaluate the characteristics of circRNA detection method false-

We simulated RNA-seq expression data of 5680 circRNAs from abundant circRNAs

Results

DCC [30], Findcirc (FC) [28], Segemehl (SE) and CIRCexplorer2 [31].

Interestingly, Hansen [24] observed that circRNA detection methods largely agreed on true predictions. In contrast, circRNAs identified by single methods were enriched in false-positive guesses (FPs) and suggested selecting circRNAs commonly predicted by two or more methods to obtain dependable results.

We formerly implemented CirComPara [25], an automated computational pipeline combining four circRNA detection methods, including CIRCexplorer [26], CIRI2 [27], Findcirc [28] and Segemehl [17]. CirComPara controlled the FP number by considering only the circRNAs commonly detected by two or more methods.

In CirComPara2, we have considerably improved our tool by: (i) including five additional circRNA detection methods, (ii) updating the software of the already integrated tools, (iii) implementing a more accurate counting of the back-spliced reads, (iv) increasing the analysis pipeline flexibility and (v) including the procedure to calculate the linear expression related to the circRNAs.

In this work, we first show that nine widely used circRNA detection tools could miss circRNAs of interest. Then, we confirm that CirComPara2 correctly reports circRNAs overlooked by other methods and achieves significantly higher sensitivity with no loss of precision. Moreover, assessment on simulated data and 142 public RNA-seq samples demonstrated the consistent higher performance of CirComPara2 compared with state-of-the-art methods. Further, we show that the CirComPara2 expression estimates highly correlate with the true circRNA abundance of simulated data. Finally, we discuss the computational cost of the CirComPara2 approach.

CircRNA detection methods could miss abundant circRNAs

We simulated RNA-seq expression data of 5680 circRNAs from the whole human genome (‘simulated data set’; see Methods) to evaluate the characteristics of circRNA detection method false-negative predictions (FNs), i.e. true circRNAs not identified as such. We applied nine widely used computational pipelines for circRNA discovery, including circRNA_finder (CF) [29], CIRI2 [27], DCC [30], Findcirc (FC) [28], Segemehl (SE) and CIRCexplorer2 [31]. CIRCexplorer2 was coupled to each of BWA (C2BW), Segemehl (C2SE), STAR (C2ST) and TopHat-Fusion (C2TH) aligners, thus composing four different pipelines.

On average, 49% of FNs detected by each method showed higher expression than the overall circRNA median expression (Figure 1A), suggesting that nearly half of the missed circRNAs had a considerable expression level no matter which method was applied. Besides, the expression distribution of the FNs was similar to the correctly identified circRNAs, i.e. the true-positive findings (TPs), whereas the false-positive (FP) expression was generally low.

Multiple method combinations increase the detection sensitivity

We further examined the 1945 circRNAs undetected by one or more tools, referred to as the ‘FN set’ from now on, by counting how many circRNAs in the FN set each method could detect. Interestingly, only 4% circRNAs of the FN set were undetected by all methods (Figure 1B), whereas 96% could be identified at least by one among the nine tools. Specifically, 1% FNs were detected individually by Segemehl, C2BW and C2TH, and 95% were commonly identified by various combinations of two or more tools. Almost half the FNs (48%) were conjointly detected by eight out of nine methods, with Segemehl, DCC and C2SE providing the most inclusive predictions. Instead, Findcirc showed the least number of recovered FNs. However, no method entirely covered the predictions of Findcirc, indicating some specificity of its algorithm.

Overall, this analysis suggested that algorithms with possibly different and complementary features can compensate for each other’s weak points and improve the detection rate if applied together.

Workflow and features of CirComPara2

Following the observation reported in the previous paragraph, we enhanced CirComPara by including more circRNA detection methods to improve its sensitivity. Moreover, we introduced new features that made CirComPara2 more flexible, computationally efficient and resilient.

CirComPara2 implements a fully automated computational pipeline for circRNA detection, quantification, annotation and integration with linear gene expression data (Figure 2A). Several parameters are available to customize the analysis workflow and the integrated methods. The minimal input consists of the RNA-seq reads in FASTQ format and a reference genome in FASTA format. Optionally, the user can also provide the gene annotation in GTF format. The software will then build the genome indexes for each read aligner and perform the necessary file format conversions. Previously computed indexes can be reused as input to save computing time.

The CirComPara2 workflow comprises an optional pre-processing of the input raw reads by Trimomatic [32] to trim or discard low-quality reads. Statistics of the pre-processing steps are produced with the FastQC tool [33]. Next, the reads are aligned collinearly to the reference genome using HISAT2 [34] to (i) identify the reads that are later used for linear transcript analysis (Figure 2A) and (ii) extract the reads not collinearly aligned, which are used to detect back-splices. The linear gene and transcript expression analysis is performed with StringTie [35] and produces files that can be easily imported into packages for downstream expression analysis, such as tximport [36] and txmeta [37]. The circRNA analysis aligns the collinearly unmapped reads independently with five methods allowing chimeric alignments, namely Bowtie2 [38], BWA-MEM, Segemehl, STAR and TopHat-Fusion (Figure 2B). The chimeric aligner outputs are subsequently parsed by six circRNA detection tools, which compose the nine different circRNA detection sub-pipelines presented in the previous paragraph. Of note, the computationally expensive chimeric alignment step is performed only once per aligner and reused by multiple circRNA detection tools, boosting CirComPara2 efficiency. For instance, the same alignments from STAR are passed to CIRCexplorer2, circRNA_finder and DCC (Figure 2B). The outputs of the various tools are automatically handled, converting them into a standard
Commonly used circRNA detection methods may overlook some highly expressed circRNAs. A. The expression level of predicted circRNAs. BJ:R: back-splice junction read counts; C2BW: CIRCexplorer2 on BWA; C2SE: CIRCexplorer2 on Segemehl; C2ST: CIRCexplorer2 on STAR; C2TH: CIRCexplorer2 on TopHat-Fusion; CF: circRNA_finder; CIRI: CIRI2; FC: Findcirc; SE: Segemehl; FP: false-positive; FN: false-negative; TP: true-positive. B. Number of methods detecting circRNAs not detected by other methods and the number of the FNs detected. Colour refers to the number of methods conjointly detecting circRNAs missed by other tools. The vertical bars show the number of FN circRNAs detected by the methods indicated in the coloured dots below the bars. The bars denote disjoint circRNA sets. Grey dots indicate the methods failing to detect the circRNAs considered in the bar chart on the top. The horizontal bars on the right represent the overall number of FN circRNAs detected by the methods. The horizontal bar shows the percentage of detected FNs by grouping method combinations according to the number of combined methods.

CirComPara2 is available as stand-alone software (https://github.com/egaffo/circompara2) and Docker image (https://hub.docker.com/r/egaffo/circompara2), which facilitates installation, portability and reproducibility of the analysis.

Optimal method combination for circRNA detection
Concurrently to the improvement of CirComPara2 sensitivity, we wanted to control the number of the introduced FPs to preserve a high precision. As observed in a previous study [24], the combination of specific methods does not ameliorate the discovery of true circRNAs. Accordingly, our method combination assessment on the simulated data showed that combining circRNA_finder or Segemehl with other methods increased the FP number (Supplementary Figure S1 available online at http://biob.oxfordjournals.org/). For this reason, circRNA_finder and Segemehl were excluded in the CirComPara2 default method combination (Figure 2B); nevertheless, these two methods can be included if enabled by the user.

Further, Hansen [24] observed that circRNAs predicted by multiple methods were enriched in reliable findings and suggested using the shared output from two (or more) algorithms.
To determine an optimal default setting to use in CirComPara2, we evaluated the amount of the recovered FNs against the introduced FPs in relation to the number of methods sharing the predictions. As expected, considering the predictions from all methods resulted in the highest recall (0.99) and the lowest precision (0.90) among the combination strategies (Figure 2C). Further, excluding predictions from single methods, i.e. selecting only the circRNAs commonly detected by two or more tools, showed a slightly reduced recall (0.98) with a substantially increased precision (0.99). This large precision gain indicated that most of the FPs were predicted by single methods. Further increasing the number of conjoint methods (i.e. from three-or-more to all seven conjoint methods) led to a considerable decrease of recall (0.96–0.68) with only a modest gain in precision (0.99–1.00). In summary, we confirmed that the larger the number of methods sharing the output circRNAs, the more reliable the findings but also observed that the amount of circRNAs discarded increased accordingly.

To evenly weigh recall and precision in ranking the method combination strategies, we calculated the $F_1$-score for each combination. The best trade-off between recall and precision, indicated by the highest $F_1$-score (0.99), was obtained with the two-or-more method strategy (Figure 2C). Therefore, the default CirComPara2 parameters have been set to simultaneously use seven circRNA detection methods ($C2BW$, $C2SE$, $C2ST$, $C2TH$, $CIRI2$, $DCC$ and $FC$) and later discard the circRNAs not shared by at least two of these methods.

**CirComPara2 outperforms other methods in simulated data**

We next set CirComPara2 with the two-or-more method strategy and compared it with the single methods on the simulated data plus CIRI-full [41] (CIRFU) and the former implementation of CirComPara (CCP1) (Figure 3A). CirComPara2 obtained the highest $F_1$-score (0.99) by achieving the highest recall (0.98) while holding a precision comparable to or higher than the other algorithms (0.99 versus 0.92–1.00), confirming that CirComPara2 rectified the true circRNAs missed by the other methods. Notably, CirComPara2 identified 7% additional true circRNAs compared to its former implementation (Supplementary Figure S2 available online at http://bib.oxfordjournals.org/).

To assess the extent of the annotation-guided method contribution to CirComPara2 predictions, we performed the analysis also with a pruned gene annotation input to the algorithms (see Methods). As expected, the CIRExplorer2 pipelines showed a dramatic reduction (up to $-0.14$) of the recall and $F_1$-scores (Supplementary Figure S3 available online at http://bib.oxfordjournals.org/). Instead, CirComPara2 maintained the highest $F_1$-score (0.98), suggesting that it can be efficient when applied to RNA-seq data of organisms with incomplete or poor genome annotation by leveraging the embedded annotation-independent tools.

A typical circRNA expression analysis usually involves post-detection data cleaning to remove background noise signals or mapping errors and poorly expressed circRNAs of little interest [42]. Consequently, the circRNAs with small back-splice junction read counts (BJRs) are routinely filtered out. As shown in Figure 1A, FPs generally have small BJRs, suggesting that more reliable circRNAs can be retained by simply filtering according to expression abundance. We applied this procedure to our data, progressively filtering out circRNAs with less than 2 up to 10 BJRs. We stress that removing circRNAs with $\leq 10$ BJRs was an extreme filter since it was close to the overall median BJR (Figure 1A). As expected, by increasing the minimum-BJR
filter threshold, all methods scored higher precision but with a corresponding reduced recall, indicating that true circRNAs were discarded as well (Figure 3B; Supplementary Figure S5 available online at http://bib.oxfordjournals.org/). Nevertheless, CirComPara2 maintained the highest recall compared to other methods at equal precision (Figure 3B; Supplementary Tables S1 available online at http://bib.oxfordjournals.org/) and obtained the highest area under the precision-recall curve (Supplementary Table S2 available online at http://bib.oxfordjournals.org/), suggesting that the circRNAs recovered by CirComPara2 were valid findings with a considerable abundance that may represent relevant circRNAs in actual experiments.

**Benchmark real data sets**

The gold standard for evaluating circRNA detection methods on real RNA-seq data is to compare ribosomal RNA-depleted (ribo⁻) with circRNA-enriched sequencing libraries. The most used technique is the additional treatment of the ribo⁻ library with RNase R to exploit the exonuclease degradation resistance of circRNAs derived by their lack of a single-stranded 3’ end [22–24, 27, 43]. Accordingly, we collected a total of 142 public real RNA-seq data samples for which sample-matched ribo⁻ and RNase R-treated libraries were available (Table 1; Supplementary Table S3 available online at http://bib.oxfordjournals.org/). This validation data set comprised samples of human cell lines and various human, Rhesus macaque and mice tissues from six independent studies.

The matched RNase R-treated samples were used as a control to define the TP circRNAs (see Methods). Assuming that the RNase R treatment would deplete the linear more than the circular transcripts, we considered TPs the circRNAs having circular-to-linear expression proportion (CLP) higher or equal in the RNase R-treated compared to the ribo⁻ matched samples according to at least one method. In this way, we accounted for different sequencing library depths between the matched samples. Moreover, each circRNA host–gene linear expression was estimated commonly to all the detection tools and independently of the circRNA abundance estimated by each method, allowing to remove possible advantages given by specific method quantification approaches.

**CirComPara2 performs consistently better than single methods**

The performance of the methods on the real RNA-seq data agreed with the analysis of simulated data and previous benchmark study results [23, 24]. We did not include CIRI-full because it showed inferior performance than CIRI2 in the simulated data. Segemehl showed high recall (median 0.75; Figure 4A) but the lowest precision (median 0.92; Figure 4B), whereas C2BW, as expected from annotation-based algorithms, showed the most reliable predictions (median precision 0.97; Figure 4B). Similar to the evaluation in the simulated data set, we computed the \( F_1 \)-scores in the real data set. According to \( F_1 \)-score medians, CirComPara2 had the highest (0.91) and significantly different value \( q < 0.001; \) Figure 4C), and substantially outperformed the next best method, Segemehl (median \( F_1 \)-score 0.82), with a 0.09 \( F_1 \)-score difference (Supplementary Table S4 available online at http://bib.oxfordjournals.org/). The highest \( F_1 \)-score, achieved by CirComPara2, resulted from a significantly larger recall (median 0.86; \( q < 0.001; \) Figure 4A) and a negligible loss of precision compared to the other methods (0.01 median reduction to the most precise method; Figure 4B). Moreover, CirComPara2 had the narrowest interquartile range of \( F_1 \)-scores across the real data sets (0.11; Figure 4C; Supplementary Table S4 available online at http://bib.oxfordjournals.org/), proving that it is robust and almost unaffected by the experimental scenario. CirComPara2 showed considerable and significant improvements also compared to its former implementation \( F_1 \)-score \( q < 0.001; \) precision \( q = 1.0; \) recall \( q < 0.001; \) Supplementary Table S4 available online at http://bib.oxfordjournals.org/), identifying in average 47% more true circRNAs than its former
Table 1. Real data sets of rRNA- and rRNA-/RNase R-treated matched pairs of samples used to benchmark the circRNA detection methods

| SRA/GEO/BIGD ID | Tissue(s)/cell line(s) | Study reference |
|-----------------|-----------------------|-----------------|
| SRR3476956, SRR1636985, SRR1636986; SRR3476958, SRR1637089, SRR1637090. SRR3479244, SRR3479243 | HeLa, HEK293 | [27] |
| SRR3479244, SRR3479243 | HeLa | [44] |
| CRA001838 (10 samples) | HeLa | [45] |
| SRR444655, SRR444975, SRR445016, SRR444974 | Hs68 | [46] |
| GSE130905 (10 samples) | HeLa | [47] |
| CRA001838 (10 samples) | HeLa | [48] |
| SRR3479244, SRR3479243 | Hs68 | [49] |
| GSE113120 (24 samples) | 22Rv1, 42D, PC3, V16, LNCaP | [50] |
| PRJCA000751 (88 samples) | Homo sapiens (17 tissues) | [51] |
| | Mus musculus (14 tissues) | |
| | Macaca mulatta (13 tissues) | |

Tissues and cell lines are from humans unless specified. GEO: Gene Expression Omnibus [50], BIGD NGDC: National Genomics Data Center [51].

*Data set GSE130905 used an improved protocol for circRNA enrichment [45].

Quantification of circRNA expression

Most circRNA expression quantification tools, including CIRC-explorer2, circRNA_finder, CIRI2, DCC and Findcirc, consider BJR counts to estimate the circular transcript abundance. However, different sets of BJRs may contribute to the expression estimates, especially if the tools were based on different chimeric aligners (Figure 5A). The method combination strategy, which we demonstrated to have superior detection performance, poses the question of which tool expression estimate should be considered or how to combine multiple expression estimates.

The former version of CirComPara estimated the circRNA abundance as the library-size normalized BJR median across the methods detecting a circRNA. Instead, CirComPara2 extracts the BJR from each method and for each circRNA to count only the non-redundant BJR identifiers (Figure 5B). By this approach, the information brought by each method is preserved.

Recent tools such as CIRIquant, Sailfish-cir and CircAST proposed a circRNA sequence reconstruction followed by read re-alignment to improve circRNA expression estimation. These methods have been devised for quantification purposes and require external tools for circRNA detection.

To evaluate the quantification accuracy of CirComPara2, we compared the true BJR counts of the simulated circRNAs with the expression estimates. We also compared the estimates from the other tools, plus CIRIquant, Sailfish-cir and CircAST. As in Zhang et al. [46], we computed the Pearson correlation coefficient between the estimated and the simulated BJR of each circRNA (Figure 5C). CirComPara2 achieved the highest correlation (r = 0.76), followed by Segemehl (r = 0.75), C2SE (r = 0.73), CIRIquant (r = 0.73) and DCC (r = 0.72). Notably, CirComPara2 largely improved over its former implementation (r = 0.63) and showed overall the most accurate expression estimates without requiring additional re-alignment steps.

Computation time and memory requirements

When running the circRNA detection methods on the simulated data, we observed that the chimeric alignment steps were the most computational-resource demanding tasks of the pipelines. In contrast, parsing the chimeric alignment output required < 1% of the resources used in the alignment step in both computation time and peak memory. Only CIRI2 showed more memory usage to process the alignments (~30 GB compared to the ~6 GB of BWA-MEM) because it ran with eight parallel processes.

CirComPara2 inherited from and extended the CirComPara1 parsimonious computational design that does not repeat tasks in common between the integrated methods. For instance, the linear read alignment is common to all the circRNA subpipelines and is performed only once. Further, the chimeric alignments of BWA-MEM are used for both CIRI2 and CIRC-explorer2. Likewise, the STAR alignments are shared by three parsers, i.e. DCC, CF and C2ST. CirComPara2 runtime and memory requirements are comparable to its former implementation since the most computational-resource demanding tools (STAR and Segemehl) were already included in CirComPara1.
Figure 5. CircRNA expression estimates from CirComPara2. A, The overlap between the different sets of the back-splice junction reads (BJRs) identified by four chimeric aligners (BWA-MEM, Segemehl, STAR and TopHat-Fusion/TopHat2). B, The CirComPara2 BJR count approach: the upper panel shows the non-redundant BJR count of CirComPara2 (CCP2) compared to other methods (for simplicity, only three methods are displayed), which is based on unique BJR identifiers (BJR IDs); the lower panel highlights how CCP2 can preserve the information even when no BJR is in common between circRNA detection methods. C, The estimated circRNA expression in BJR compared to the true expression from the simulated data set. The Pearson correlation coefficient ($r$) values are reported in the boxes. CIRIquant, CircAST and Sailfish-cir were inputted with the circRNAs predicted by CirComPara2.

When processing the simulated data set (~3.7 million paired-end reads, 100 bp long, from the whole human genome), CirComPara2 saved ~20% computation time compared to the sequential independent running of the seven method full pipelines (SeqMet; Figure 6). Besides, CirComPara2 provides concurrent running of independent tasks, including the circRNA detection methods, which can reduce the computation time. Setting CirComPara2 to run multiple (up to seven) tasks in parallel
Figure 6. Computation time and memory requirements of circRNA detection methods. $F_1$-scores of 14 circRNA detection pipelines compared to their computing time in seconds (s) and peak random access memory (in GigaBytes) required to process $\sim$3.7 million paired-end (PE) reads of 100 bp length, mapped to the human genome (GRCh38) and using eight CPU threads for each tool. Plus, one hypothetical pipeline running sequentially the seven methods used in CirComPara2 (SeqMet) and CirComPara2 set with maximal parallelization settings (mpCCP2) allowing to run up to seven tasks in parallel, for peak usage of 56 CPU threads in total.

(mpCCP2) saved $\sim 74\%$ computing time compared to the SeqMet, and $\sim 68\%$ compared to CirComPara2 with no augmented parallelization (Figure 6). The mpCCP2 setting reduces the computation time up to the longest program execution (C2TH) but at the cost of higher peak memory and CPU cores requirements: 56 CPU cores and up to 120 GB of random access memory were used to run in parallel all the aligners on the human genome, and CIRI2 with eight parallel processes.

Finally, the CirComPara2 running time scaled linearly with the number of input sequencing reads (Supplementary Figure S5 available online at http://bib.oxfordjournals.org/). CirComPara2, set with the mpCCP2 parameters and applied to a multiple-sample batch, showed a runtime bounded only by the analysis of the sample having the largest input size (Supplementary Figure S5 available online at http://bib.oxfordjournals.org/). This result demonstrates that the CirComPara2 augmented parallelization harnessed the system’s computational power by consuming multiple short tasks concurrently to longer ones also from different samples, eventually saving computation time.

Discussion

In the early days after circRNA discovery, bioinformaticians put significant effort into developing circRNA detection methods with highly precise predictions. To this aim, the current circRNA detection algorithms apply filtering procedures to remove FPs, but that may also reject true circRNAs [21-24], increasing the number of FNs and the risk of overlooking circRNAs of interest, as suggested by our analysis. Moreover, the frequent practice of filtering out low-count circRNAs to improve precision may result in the loss of differentially expressed elements in comparative studies, as demonstrated by systematic studies on gene expression [52, 53]. Notably, unlike FPs, experimental validations cannot amend the bias derived by FN errors. These considerations prompted us to consider a method evaluation metric that equally weighted precision and recall, such as the $F_1$-score.

Interestingly, the evaluation by $F_1$-score in our simulated data analysis highlighted that some methods with diverging precision and recall, such as CIRI2 and Segemehl (Figure 3A), were equally ranked. Moreover, the real data set analysis showed an ample recall variation of each method across the samples, while precision scores were tighter both between and within methods. These results confirmed that choosing a single method for circRNA discovery is problematic since no single best performing method exists. We showed that an approach leveraging the advantages brought by each of the integrated methods allows CirComPara2 to perform better than the single methods consistently in different experimental contexts. Besides, we showed that CirComPara2 predictions are more inclusive and robust than single-method ones, even upon low-count filtering, indicating that the recovered circRNAs missed by other methods are not merely of low abundance.

To our knowledge, CirComPara has been the first bioinformatics tool to combine multiple circRNA detection methods in an automated software pipeline. Since its former implementation, the continuous upgrade of CirComPara evolved into a substantially improved new tool, CirComPara2, with an efficacious method combination strategy. Other computational pipelines that employed multiple circRNA detection tools, such as RAISE [54] and circRNAwrap [55], considered the circRNAs predicted by the union or the intersection of all methods, respectively, without validating which method combination was best. Our data show that those two approaches could be suboptimal as they suffer either low precision or sensitivity, whereas the best trade-off between precision and recall is achieved with predictions of methods taken pairwise.

Recently, large circRNA databases have been compiled using method combination strategies [56]. For instance, circAtlas2.0 [57] retained circRNAs identified by at least two methods among CIRI2, CIRCexplorer2, Findirc and DCC; the same combination but replacing DCC with circRNA_finder has been used in CircRic [58]. Applying these two approaches to our real data sets analysis, we observed that they performed better than most of the algorithms except CirComPara2 (Supplementary...
considered the realignment of the reads onto pseudo-reference
circRNA detection to estimate the circRNA expression. They
and CircAtlas2.0 applied a quantification step downstream of
additional realignment of the reads. CircRNAwrap, RAISE
dant counts of BJR identified by multiple tools without relying
expression estimates into unified values that eliminate redun-
documentation [59].

freeing bioinformaticians from the several difficulties posed by
pipeline ready-to-use and portable through a Docker container,
the automated and computationally efficient CirComPara2
tools available to the scientific community. Instead, we made
databases in future works.

better recall. Such a result suggests that CirComPara2 could
which achieved similar precision but slightly more than 0.2
Table S4 available online at http://bib.oxfordjournals.org/),
demonstrated the asset given by the CirComPara2 method com-
combination strategy for circRNA discovery, which provides robust
and inclusive predictions in diverse biological contexts. With
CirComPara2, we aim to provide a helpful bioinformatics tool
to obtain a more comprehensive picture of transcriptomes and
boost the understanding of circRNA features, biological and
pathogenetic roles.

Methods

Simulated data set

CircRNA reads were simulated with the CIRI_simulator from
the CIRI2 tool suite using the whole GRCh38 human genome and
Gene expression V29 gene annotation.

The parameters used in CIRI_simulator were: -C 20 -LC 0 -R 1
-LR 1 -L 101 -E 1 -CHR 1 0 -M 250 -M2 450 -PM 0 -S 70 -S2 0 -SE 0
-PSI 10.

A total of 10% (30% for the highly pruned annotation simul-
ation) of the gene annotation for the simulated circRNA parent
genes was removed from the annotation file and used as input
to circRNA detection methods and to simulate linear transcript
reads with Polyester (the reads_per_transcript parameter was set
to 300). The linear transcript read files were then concatenated
to the circRNA read files.

Code for generating the simulated data is available as a
software tool CCP_simulator at https://github.com/egaffo/CCP_simu-
lator. The parameter ANNOPARTS used in CCP_simulator for
the two filters on gene annotation (standard and high pruning)
was set to ‘85,4,5,1,0,0,0’ and ‘60,15,10,2,5,6,3,5,3’, respectively.

Real data sets

Overall, 71 samples with matched rRNA depletion and rRNA
depletion followed by RNase R treatment libraries from six stud-
ies (Table 1), for a total of 124 samples processed, were retrieved
from Gene Expression Omnibus (GEO) or the National Genomics
Data Center (NGDC) databases.

Reads from PRJCA000751 were trimmed to 150 bp in the pre-
processing phase, as reported in the original work.

Method predictions in genomic scaffolds or from the mito-
ochondrial genome were not considered. CircRNAs predicted with
a length shorter than the library read length or longer than the
longest gene expressed in the sample (computed as genes with
TPM ≥ 1 computed by Stringtie v2.1.4) were filtered out.

CircRNA detection methods’ parameters

The following gene and gene annotations from the Ensembl
database were used in the analyses: GRCh38 human genome
and v97 gene annotation, Mmul 10 Macaca mulatta genome and v101
gene annotation, and GRCm38 Mus musculus genome and v101
gene annotation.

CirComPara2 default parameters were set for the analy-
yses, which are as follows: adaptors from the Trimmomatic
v0.39 TruSeq3-PE-2.fa file; PREPROCESSOR = ‘trimmomatic’;
PREPROCESSOR_PARMS = ‘MAXINFO=40:0.5 LEADING:20 TRAIL-
ing:20 SLIDINGWINDOW:4:30 MINLEN:50 AVGQUAL:26’. For the
PRJCA000751 data sets, the CROP:150 option was appended
to the parameter. STAR_PARMS = ‘-runRNaseGend 123 —
outFilterOverhangMin 15 15 15 —alignSjoverhangMin 15 —
alignSJDBoverhangMin 15 —seedSearchStartLmax 30 —outFilter
ScoreMin 1 —outFilterMatchNmin 1 —outFilterMismatchNmax
2 —chimSegmentMin 15 —chimScoreMin 15 —chimScore
Separation 10 —chimJunctionOverhangMin 15’. CIRCRNA_
Assuming that RNase R should degrade linear transcripts and the control library and the biochemical variability of the exoribonuclease in RNase R treatment have been adjusted by considering the control sample and to RNase R-sensitive circRNAs.

The precision was defined as TP/(TP + FP). The recall was computed as TP/(TP + FN) and F$_1$-score as (2 × Precision × Recall)/(Precision + Recall), where TP and FN denote the true-positive and false-negative numbers.

Wilcoxon one-tailed paired tests were used to compare CirComPara2 greater recall and F$_1$-score or lower precision with each method. Bonferroni multiple test correction was applied to compute adjusted P-values (q-values). The q-values reported in the main text refer to the highest value among the pairwise comparisons for recall and F$_1$-score, whereas to the lowest value for precision comparisons.

Software versions

The following software versions of the circRNA detection methods and chimeric read aligners were used in this study: CIRCexplorer2 [31] v2.3.8, CircRNA_finder [29] v1.1, CIRI2 [27] v2.0.6, DCC v0.4.8 [30], Findcirc [28] v1.2, Segemehl [17] v0.3.4 (used also by CIRCexplorer2), BWA MEM [16] v0.7.15 (used by CIRI2 and CIRCexplorer2), STAR [15] v2.6.1e (used by circRNA_finder, DCC and CIRCexplorer2) and TopHat2 [14] v2.1.0 (used for the TopHat-Fusion algorithm by CIRCexplorer2). CirComPara2 v0.12.1 was used to run the analysis. Other method versions used were as follows: CirComPara v0.1.3, CIRI-full v2.0 (the embedded CIRI-vis had to be replaced with an updated version CIRI-vis v1.4), CIRIquant v1.1.2, CircAST v1.0.2 [71] and Sailfish-circ v0.11a.

### Key Points

- Current circRNA detection methods achieve either high precision or high recall, possibly overlooking circRNAs of interest.
- Extensive tests on simulated RNA-seq expression data determined the optimal method integration strategy for circRNA detection.
- CirComPara2 achieves high detection recall with no loss of precision by combining seven circRNA detection methods.
- At benchmarking on 142 real data sets, CirComPara2 consistently outperforms other methods regardless of the biological context and the genome annotation quality.

### Supplementary Data

Supplementary data are available online at Briefings in Bioinformatics.

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Data Availability

The data underlying this article are available in the Gene Expression Omnibus (https://www.ncbi.nlm.nih.gov/geo/) and the National Genomics Data Center (https://www.ncbi.nlm.nih.gov/geo/). The data can be accessed with SRR3476956, SRR1636985, SRR1636986, SRR3476958, SRR1637089, SRR1637090, SRR3479244, SRR3479243, GSE130905, SRR444655, SRR444975, SRR445016, SRR444974, GSE113120 and CRA001838 and FR/CA000751 accession numbers. Further details are referenced in Table 1 of this manuscript.

The code of CirComPara2 is available on GitHub at https://github.com/egaffo/circompara2.

Author Contributions

Conceptualization, E.G.; Data Curation, E.G.; Formal Analysis, E.G. and A.B.; Funding Acquisition, E.G. and S.B.; Methodology, E.G. and A.B.; Project Administration, S.B.; Resources, S.B.; Software, E.G. and A.B.; Supervision, E.G. and S.B.; Visualization, E.G.; Writing-Original Draft, E.G., A.B. and S.B.; Writing-Review & Editing, E.G., A.B., A.D.M. and S.B.

All authors read and approved the final manuscript.

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