

Title: exceRpt: A Comprehensive Analytic Platform for Extracellular RNA Analysis

Rozowsky J^{1,2*}, Kitchen R^{1,2*}, Park JJ^{1,2*}, Galeev TR^{1,2}, Diao J², Warrell J^{1,2}, Subramanian SL³, Thistlethwaite W³, Milosavljevic A³, Gerstein M^{1,2,4}

1. Program in Computational Biology and Bioinformatics, Yale University, New Haven, CT
2. Department of Molecular Biophysics and Biochemistry, Yale University, New Haven, CT
3. Bioinformatics Research Laboratory, Molecular and Human Genetics Department, Baylor College of Medicine, Houston, TX
4. Department of Computer Science, Yale University, New Haven, CT

Summary:

Extracellular RNAs (exRNA) are gaining considerable interest for their potential use as biomarkers and their role in intercellular communication. Small RNA-sequencing has been widely adopted to shed light on the vast diversity of exRNA in biofluids, but the field faces unique technical challenges. In particular, exRNA samples are more vulnerable to contamination and artifacts from different isolation techniques, are present in lower concentrations than cellular RNA, and some are of exogenous origin. To address these challenges we present the extracellular RNA processing toolkit (exceRpt), the standardized small RNA-seq analysis pipeline of the NIH Extracellular RNA Communication Consortium (ERCC) optimized for exRNA analysis. exceRpt is structured as a cascade of filters and quantifications prioritized based on the confidence of a given set of annotated RNA. The pipeline generates quality control reports and abundance estimates for RNA biotypes, and is also capable of characterizing alignments to exogenous genomes, which can be used to generate phylogenetic trees. exceRpt has uniformly processed all (currently ~2500) exRNA-seq datasets in the public exRNA atlas, and is available at genboree.org and github.gersteinlab.org/exceRpt.

Highlights and eTOC Blurb:

- exceRpt processes and analyzes exRNA profiling data
- Generates quality control metrics, RNA biotype abundance estimates, and processing reports
- User-friendly, browser-based graphical interface available
- Processes all RNA-seq datasets in the exRNA Atlas

Introduction:

Recent discoveries of extracellular RNA (exRNA) in the blood and other body fluids have added a new dimension to the paradigm of intercellular signaling (Patton et al., 2015; Skog et al., 2008). The relative stability of exRNA within extracellular vesicles (EVs) or bound to proteins or lipids (Yanez-Mo et al., 2015), coupled to the availability of sensitive and specific tools such as RNA-seq, underpins the emergence of exRNA profiling as an exciting new opportunity for biomarker discovery. exRNA-based liquid biopsy is particularly attractive as a non-invasive mode for monitoring disease due to the significantly increased accessibility of biofluids over tissues, thereby allowing more frequent and longitudinal sampling (Byron et al., 2016). With better characterization of the differences between profiles secreted by diseased and healthy tissues, the diagnostic and prognostic utility of exRNA-based profiling is increasingly becoming a reality (Akat et al., 2014; McKiernan et al., 2016; Yuan et al., 2016).

Despite rapid technological and methodological progress in isolating EVs and exRNA, the analysis and interpretation of exRNA data is uniquely challenging. Biochemical methods for extraction, purification, and sequencing of exRNAs are much more vulnerable to contamination and artifacts than cellular RNA preparations, in large

part due to relative low abundance (Danielson et al., 2017). Quality control prior to sequencing of samples derived from EV or exosome preparations is difficult due to the lack of reliable ‘housekeeping’ markers, such as the ratio of 18S and 28S ribosomal RNAs (Tataruch-Weinert et al., 2016). The variable presence of rRNA in mixtures of low- and high-density EVs (Lasser et al., 2017); deterministic cleavage of structured smallRNA (tRNAs and piRNAs) and longer RNA molecules; and imperfect annotation of miRNAs, piRNAs, and tRNAs all pose challenges for quantification and functional interpretation of exRNA. Furthermore, it has been suggested that exogenous exRNAs may be also present at detectable levels in some biofluids (Freedman et al., 2016; Yeri et al., 2017), and careful analysis is required to differentiate these sequences from endogenous RNA molecules. For these reasons, existing computational tools capable of analyzing smallRNA-seq data are not as well suited to the new field of exRNA analysis.

To address these analytical challenges, we present here the extracellular RNA processing tool (exceRpt). exceRpt is the primary smallRNA analysis pipeline of the NIH Extracellular RNA Communication Consortium (ERCC). By providing an optimized and standardized bioinformatics platform, exceRpt reduces technical bias and allows for cross-study analyses to potentiate meaningful insights into exRNA biology.

Results:

Filtering and Quantification Cascade:

The exceRpt pipeline is composed of a cascade of serial computational filters and alignments, whose order is designed to reflect an appropriate level of confidence in the various endogenous and exogenous sequences that may exist in the extracellular fraction (Figure 1A). To combat potential contamination in an RNA-seq library, reads aligned to known contaminants are removed before mapping to the host genome and to the exogenous sequences. The pipeline is highly modular (constructed as a makefile file containing shell, java, and R modules), allowing the user to define the order of which smallRNA annotations are used during read-mapping; it includes support for random-barcoded libraries and spike-in sequences for calibration. The general workflow comprises steps for preprocessing, endogenous alignment, and exogenous alignment (Figure 1A).

Preprocessing:

First, exceRpt automatically identifies and removes 3’ adapter sequences. If specified, it will also remove and store information from random-barcoded insert sequences, which are increasingly being used in smallRNA sequencing in an attempt to identify and compensate for ligation and/or amplification artifacts that have the potential to affect downstream quantification (Fu et al., 2014). If exogenous spike-in sequences were added to the library, exceRpt aligns adapter- and barcode-clipped reads against either standard NIST/ERCC sequences or sets of user-specified sequences. RNA-seq reads of consistently low-quality reads are removed, as are reads aligned to sequences in the NCBI’s UniVec/Vecscreen database in order to filter out common laboratory contaminants. Finally, reads aligned to all primary endogenous ribosomal RNAs (5S, 5.8S, 18S, 28S, and 45S), many of which are highly variable in abundance in EV preparations.

Endogenous Alignment:

RNA-seq reads are aligned to the host genome and transcriptome of human or mouse, and transcript abundances are calculated (RNAs are quantified using both raw read counts and normalized reads per million (RPM)). Because of the variety of RNA preparations available (totalRNA, smallRNA, miRNA), exceRpt allows the user to

prioritize the order in which annotations (miRBase, tRNAscan, piRNA, GENCODE, circRNA) are used for quantification based on our confidence in the presence of a given annotation in a given sample. For example, reads from a miRNA-seq prep can be assigned to miRBase miRNA annotations before piRNA annotations. Likewise, reads from long- or total-RNA preparations can be assigned to longer GENCODE transcripts before (or instead of) the other smallRNA libraries. This feature is particularly relevant for lower-confidence annotations; piRNAs, for example, are generally given lower priority than tRNAs to ensure correct read assignments.

Exogenous Alignment:

exceRpt has been designed to enable confident assessment of non-human sequences in biofluids after careful, explicit removal of as many known or likely contaminants as possible. Before interrogating non-contaminant and non-endogenous reads for potential exogenous sequences, exceRpt includes a second-pass alignment against the host genome and known repetitive sequences using significantly relaxed mapping criteria. This step serves to remove additional sequences that most likely derive from the host genome and makes exceRpt more conservative in the identification of exogenous sequences. Reads are then aligned to curated libraries of annotated exogenous miRNAs in miRBase and exogenous rRNA sequences in the Ribosomal Database Project (RDP), followed by alignment to the full genomes of all sequenced bacteria, viruses, plants, fungi, protists, metazoa, and selected vertebrates that are potentially part of the host diet. Existing approaches for exogenous sequence alignment remove degenerate sequences (i.e. those that co-occur across multiple species), which results in a loss of potentially valuable data as reads frequently align across multiple species/strains. By characterizing exogenous genome alignments generated by exceRpt in terms of the NCBI taxonomy tree and assigning reads to the most specific possible node in the phylogenetic tree (many reads can only be assigned to nodes higher up in the phylogenetic tree due to not uniquely mapping to a specific genome of a sub-species), users may obtain valuable information regarding the contribution of the flora to various exRNA samples and can generate phylogenies for cross-sample comparison. For example, in Figures 2B and 2C we present the phylogenetic trees of the reads that we assign to bacterial ribosomal and genome sequences for a specific saliva sample. In both trees, we find an abundance of reads assigned to the node corresponding to the genus *Streptococcus*. We have a high degree of confidence in these results given that the sets of short reads used for constructing these two trees are disjoint.

Pipeline Output and Analysis:

The pipeline generates bulk statistics for abundance of the various RNA biotypes in addition to sample-level quality control (QC) and processing reports. Due to the heterogeneity of the various exRNA isolation techniques, we developed QC metrics in collaboration with the ERCC for identification of clear experimental outliers (Figure 2D). These metrics are based on transcriptome-mapped read count and the proportion of reads that map to annotated RNA transcripts, since enrichment of RNA can be used to distinguish exRNA samples from DNA-enriched cellular contaminants (see STAR Methods). Descriptions of the post-processing output files and diagnostic plots generated by exceRpt are listed in Table 1. As a performance evaluation, we found that the endogenous miRNA abundance estimates produced by exceRpt are in close agreement with existing tools. Comparing exceRpt-filtered read counts for miRBase miRNAs, we obtain an average Pearson correlation of 99.98% to the counts produced by miRDeep2 (Figure S1). As another performance evaluation, running the same sample

through the pipeline with individual steps excluded shows the effect of the filters and alignments on downstream quantifications (Figure 1B). This analysis shows that the pre-filtering of low-quality and low-complexity reads and reads that align to UniVec or rRNA sequences account for a sizeable fraction of the total number sequenced and, without explicit removal, can align to the human genome leading to potential confounding and added quantification variability. UniVec-based contaminant filtering has the largest effect on the fraction of reads aligning to exogenous genomes, and its omission substantially increases the number of reads that appear to be, but are very likely not, exogenous in origin.

These bulk statistics can be used to differentiate biofluids (or tissues, if exceRpt is run on cellular samples) based on their RNA distribution. For example, results from samples selected from the exRNA Atlas (Figure 2A) show that, relative to other biofluids, saliva samples tend to have more reads that are unmapped or that map to exogenous genomes, which is consistent with saliva's high potential for bacterial contamination and exposure to the external environment. Moreover, abundance quantifications for specific RNA biotypes can show which miRNAs (or other RNA biotype) are most highly represented in a particular sample (Figure S2). This information is critical for understanding the composition of exRNA profiles and for interrogating their biological significance.

Discussion:

The exceRpt pipeline was built to address the need for a standardized bioinformatics processing platform in extracellular RNA research, and is structured as a principled, biologically driven series of alignment, filtering, and quantification steps where unmapped reads are used as inputs to the next step. The prioritization of steps is biased in favor of conservative estimates for RNA quantifications, with higher confidence RNA reference sets (by degree of expectation or annotation quality) having higher priority. To date, exceRpt has uniformly processed and applied QC standards to all of the datasets in the ERCC exRNA Atlas (<http://exrna-atlas.org/>) (Subramanian et al., 2015). The exceRpt pipeline enables a variety of user-specified customization, including RNA reference prioritization, random-barcoding, spike-in support, and detailed quantification reports. The pipeline is available as source (available at github.gersteinlab.org/exceRpt) or wrapped in a user-friendly, browser-based interface available at genboree.org.

Tables:**Table 1:** Description of Output Files

File Name	Description of File
QC Data	
exceRpt_DiagnosticPlots.pdf	All diagnostic plots automatically generated by the tool
exceRpt_readMappingSummary.txt	Read-alignment summary including total counts for each library
exceRpt_ReadLengths.txt	Read-lengths (after 3' adapters/barcodes are removed)
Raw Transcriptome Quantifications	
exceRpt_miRNA_ReadCounts.txt	miRNA read-counts quantifications
exceRpt_tRNA_ReadCounts.txt	tRNA read-counts quantifications
exceRpt_piRNA_ReadCounts.txt	piRNA read-counts quantifications
exceRpt_gencode_ReadCounts.txt	gencode read-counts quantifications
exceRpt_circularRNA_ReadCounts.txt	circularRNA read-count quantifications
Normalized Transcriptome Quantifications	
exceRpt_miRNA_ReadsPerMillion.txt	miRNA RPM quantifications
exceRpt_tRNA_ReadsPerMillion.txt	tRNA RPM quantifications
exceRpt_piRNA_ReadsPerMillion.txt	piRNA RPM quantifications
exceRpt_gencode_ReadsPerMillion.txt	gencode RPM quantifications
exceRpt_circularRNA_ReadsPerMillion.txt	circularRNA RPM quantifications
R Objects	
exceRpt_smallRNAQuants_ReadCounts.RData	All raw data (binary R object)
exceRpt_smallRNAQuants_ReadsPerMillion.RData	All normalized data (binary R object)

Figure Legends:**Figure 1**

(A): exceRpt schema: Samples in FASTA, FASTQ or SRA file formats are used as inputs to exceRpt. Adapter and random barcode sequences are removed, followed by a read-quality filter, optional spike-in quantification and removal, and UniVec contaminant removal. High quality filtered reads then enter the endogenous quantification engine, with RNA library prioritization defined by the user. After a second-pass endogenous genome and repetitive elements filter, reads are mapped to the exogenous miRNA, rRNA, and genomic libraries.

(B): Leave-one-out analysis: Running the pipeline multiple times with individual steps removed shows the effect of those steps on subsequent alignments. The sample used for this analysis was SRR822433, a plasma exRNA plasma sample. Low-quality and low-complexity reads and reads that align to UniVec or rRNA sequences account for a sizeable fraction of the total number sequenced. Removing the UniVec alignment step significantly increases the number of reads that, likely incorrectly, map to the exogenous genomes.

Figure 2

(A): Read distributions: exceRpt outputs endogenous alignment quantifications which can be used to compare RNA biotype distributions in exRNA samples. Here, saliva has a higher proportion of exogenous sequences than other samples, and urine has a higher proportion of tRNA sequences. Quantifications can also be performed for cellular

datasets, such as ENCODE samples, where the majority of reads align to long coding and non-coding RNAs in Gencode.

(B+C): Exogenous alignment phylogeny with genome reads: Exogenous sequence quantifications based on exogenous genome reads and rRNA reads can be represented using phylogenetic trees. The tree in (B) was constructed using 1.74M genome reads from a saliva small exRNA-seq sample, and the tree in (C) constructed from 1,127K ribosomal reads from the same saliva. Saliva biofluids are distinguished from other biofluids by their exposure to a robust and complex bacterial community in the oral cavity (Hasan et al., 2014), which causes a greater contribution of reads of bacterial origin (and not human genome) to the sample. In both the phylogenetic trees constructed using bacterial genome mapped (B) and ribosomal (C) reads, we find an abundance of reads assigned to the node corresponding to the genus *Streptococcus*.

(D): Quality Control metrics: ERCC QC metrics are based on number of transcriptome reads and ratio of RNA annotated reads to the genome reads. The horizontal and vertical lines define QC threshold minima, and most exRNA Atlas samples meet the standards and fall in the upper right quadrant.

Figure S1: miRNA quantification comparison with miRDeep2

miRDeep2 is a commonly used software package for identification of miRNAs in deep sequencing data. Read quantifications for miRNAs based on results from processing an exRNA plasma sample with miRDeep2 and exceRpt are highly correlated, with an average Pearson correlation of 99.98%. This suggests that miRNA quantification results from exceRpt are highly consistent with existing tools.

Figure S2: miRNA abundance for exRNA samples

exceRpt produces miRNA quantifications for exRNA samples, which can be used to see which miRNAs are enriched within which samples. Here, enriched miRNAs in samples from four different biofluids are shown.

Figure S3: Exogenous sequence characterization algorithm

For exogenous sequence alignment, reads are assigned to most specific node possible.

Figure S4:

(A): Genboree interface for exceRpt

Signing up for an account on genboree.org is the quickest and easiest way to start processing exRNA samples using exceRpt. The Genboree Workbench offers a graphical, browser-based interface for processing exRNA samples and storing exceRpt results. Shown is the working interface for Genboree, including navigation functionalities and locations for inputs and outputs.

(B): Docker implementation for exceRpt

exceRpt is also available as a Docker image at github.com/gersteinlab/exceRpt/. Shown are example commands for running a sample to the exceRpt pipeline using Docker. Detailed instructions are on the GitHub webpage.

Figure S5: Read distributions with unmapped reads

By including unmapped reads into the visualization of the read distributions for exRNA and ENCODE samples (Figure 2A), we see that saliva also has a higher proportion of unmapped reads compared to the other biofluids.

STAR Methods:**KEY RESOURCES TABLE:**

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Deposited Data		
exRNA Atlas	ERCC	https://exrna-atlas.org/
Human reference genome build GRCh38 (UCSC hg38)	Genome Reference Consortium	https://www.ncbi.nlm.nih.gov/grc/human
Human reference genome build GRCh37 (UCSC hg19)	Genome Reference Consortium	https://www.ncbi.nlm.nih.gov/grc/human
Mouse reference genome build GRCm38 (UCSC mm10)	Genome Reference Consortium	http://www.ncbi.nlm.nih.gov/grc/mouse
miRBase version 21	(Griffiths-Jones, 2004)	http://www.mirbase.org/
GtRNADB	(Chan and Lowe, 2009)	http://gtrnadb.ucsc.edu/
piRNABank	(Sai Lakshmi and Agrawal, 2008)	http://pirnabank.ibab.ac.in/
GenCode version 24 (hg38)	(Harrow et al., 2012)	http://www.encodegenes.org/
GenCode version 18 (hg19)	(Harrow et al., 2012)	http://www.encodegenes.org/
GenCode version M9 (mm10)	(Mudge and Harrow, 2015)	http://www.encodegenes.org/
circBase	(Glazar et al., 2014)	http://www.circbase.org/
UniVec	NCBI	ftp://ftp.ncbi.nlm.nih.gov/pub/UniVec/
Ribosomal Database Project	(Cole et al., 2014)	http://rdp.cme.msu.edu/
Software and Algorithms		
exceRpt version 4.6.2	This paper	http://genboree.org/theCommons/projects/exrna-tools-may2014/wiki/Small%20RNA-seq%20Pipeline
Java	Oracle Corporation	https://www.java.com/
R version 3.2	The R Project	https://www.r-project.org/
FASTX version 0.0.14	Hannon Lab	http://hannonlab.cshl.edu/fastx_toolkit/
STAR version 2.4.2a	(Dobin et al., 2013)	https://github.com/alexdobin/STAR/releases
Bowtie 2 version 2.2.6	(Langmead and Salzberg, 2012)	http://bowtie-bio.sourceforge.net/bowtie2/index.shtml
Samtools version 1.3.1	(Li et al., 2009)	http://www.htslib.org/
FastQC v0.11.2	Babraham Bioinformatics	http://www.bioinformatics.babraham.ac.uk/projects/fastqc/
SRA-Toolkit version 2.3	NCBI	https://trace.ncbi.nlm.nih.gov/Traces/sra/sra.cgi?view=software

PREPROCESSING

Input files of sequenced reads can be in FASTA, FASTQ, or SRA formats. exceRpt begins the preprocessing step by automatically identifying and removing 3' adapter sequences. Randomly barcoded 5' and/or 3' adapter sequences are increasingly being used in smallRNA sequencing in an attempt to identify and compensate for ligation and/or amplification artifacts. exceRpt is capable of removing and quantifying these biases at both the insert level, which reveals ligation/amplification bias, and the transcript level, which provides an opportunity to compensate for the bias by counting unique N-mer barcodes rather than counting the number of inserts. Random-barcode/UMI information from the raw reads is preserved throughout the alignment and quantification steps and may be used as an alternative to read-counting in the quantitation of smallRNAs. The pipeline then aligns against an input set of known spike-in sequences (if used in the library construction), followed by a filter to remove low-quality reads and reads with large homopolymer repeats using the FASTX toolkit (QFILTER_MIN_READ_FRAC = 80, QFILTER_MIN_QUAL = 20). As the final preprocessing step, exceRpt aligns reads to annotated sequences in the UniVec database (designed for filtration of common laboratory contaminants) and then to endogenous ribosomal RNAs, both of which are highly variable in abundance in EV preparations. The mapping to calibrators (spike-ins) step uses Bowtie2. All other preprocessing mapping steps use STAR.

ENDOGENOUS QUANTIFICATION

Reads that were not filtered out in the preprocessing steps of the pipeline are aligned to the endogenous genome and transcriptome of either human or mouse, using STAR. STAR uses the following parameters for the endogenous alignments by default, MIN_READ_LENGTH = 18, MAX_MISMATCH = 1, MISMATCH_OVER_L_MAX = 0.3, MATCH_N_MIN_OVER_L_READ = 0.9. Transcript abundances are calculated (RNAs are quantified using both raw read counts and normalized reads per million (RPM)). The user can prioritize the order in which annotations (miRBase, tRNAscan, piRNA, GENCODE, circRNA) are used for quantification based on confidence in the presence of a given RNA biotype in a given sample. By default the order for quantification is miRNAs, tRNAs, piRNAs, GENCODE transcripts (snRNAs, snoRNAs, miscRNAs, protein coding genes and lncRNAs) followed by circular RNAs based on our confidence of the likelihood of these annotations to be present in a small exRNA-Seq sample. The exceRpt pipeline can be used with the human host endogenous genome (either hg19 or GRCh38) or the mouse host endogenous genome (mm10) with corresponding annotations.

EXOGENOUS QUANTIFICATION

Before analyzing the remaining reads for potential exogenous sequences, exceRpt performs a second pass alignment using STAR, first against the host genome (allowing for novel gapped alignments) with more relaxed mapping criteria, and then against a database of known repetitive sequences. Reads are then aligned to curated libraries of annotated exogenous miRNAs in miRBase and exogenous rRNA sequences in the Ribosomal Database Project (RDP), followed by alignment to the full genomes of all sequenced bacteria, viruses, plants, fungi, protist, metazoa, and the following 12 vertebrate genomes: chicken, cod, cow, dog, duck, frog, horse, rabbit, pig, sheep, tilapia, and turkey. Multiple STAR indexes were constructed provided in the exceRpt database for the bacterial genomes, the plant genomes, the metazoan genomes, fungi, protest and virus genomes and the vertebrate genomes so that exogenous genome alignment, which is the most time-consuming step of the exceRpt pipeline, can be parallelized. exceRpt then uses STAR to align the remaining reads to these genomes. By default, exceRpt allows for no mismatches during this step (in order to be as conservative as possible in identifying possible exogenous sequences).

Since many exogenous genomes have a high degree of sequence similarity based on evolution, we find that many reads that align to an exogenous genome align to multiple genomes. We assign reads that align to exogenous genomes to the position in the phylogenetic taxonomy tree based on the node is most parsimonious with the different genomes that the read aligns (Figure S3). Reads that align to a unique genome are aligned to the corresponding leaf node in the phylogenetic tree while reads that align to multiple genomes are assigned to nodes higher up in the tree. exceRpt also independently performs this assignment for the reads that align to exogenous rRNA sequences in the context of the phylogenetic taxonomy tree constructed using rRNA sequences. One can compare the structures of the exogenous genomic and rRNA phylogenetic taxonomy trees for consistency, as the trees are constructed independently using the disjoint sets of exogenous genome and rRNA reads.

QUALITY METRICS

To evaluate the quality of the samples and to identify outliers, we developed QC data standards in collaboration with the ERCC which exceRpt evaluates uniformly on all input samples: (1) datasets are required to have at least 100,000 reads that overlap with

any annotated RNA transcript in the host genome, and (2) over 50% of the reads that map to the host genome also align to any RNA annotation. The first criterion ensures that enough reads are generated for quantification (the minimal read depth required for the minimal normalized expression of an annotated RNA should be greater than 1 RPM) and the second criterion ensures that the reads mostly align to RNA, as opposed to DNA contamination from cellular sources. We find that 95% of the ~2500 exRNA-Seq datasets that have been uniformly processed in the exRNA Atlas with exceRpt meet both criteria (Figure S3), with most datasets well above both thresholds.

QUANTIFICATION AND STATISTICAL ANALYSIS:

All quantification analyses were performed in R. Annotated RNAs are quantified using both raw read counts and are also normalized by sample across all RNA biotypes to reads per million mapped reads (RPM)). Diagnostic plots and output statistics are also automatically generated by exceRpt.

DATA AND SOFTWARE AVAILABILITY:

The graphical, browser-based, user-friendly interface for uploading and processing of exRNA-seq datasets with exceRpt is available at the Genboree Workbench: <http://genboree.org/theCommons/projects/exrna-tools-may2014/wiki/Small%20RNA-seq%20Pipeline>. The exceRpt source code may be downloaded and installed manually for the greatest degree of flexibility ([github.gersteinlab.org/exceRpt](https://github.com/gersteinlab/exceRpt)). For this option, the dependencies and databases used by the pipeline must be downloaded and installed, and the makefile must be properly configured for use (instructions for installation and use are available from the GitHub page). Alternatively, the exceRpt Docker image with all required dependencies may be used for installation on the user's own machine or cluster (github.com/gersteinlab/exceRpt). exceRpt also includes a script (`mergePipelineRuns.R`) to combine outputs from multiple samples for downstream comparative analysis.

ADDITIONAL RESOURCES:

The ERCC exRNA Atlas can be found here: <https://exrna-atlas.org/>

The ERCC quality control standards can be found here:

<https://exrna.org/resources/data/data-quality-control-standards/>

Supplemental Information:

Document S1. Figures S1–S5.

Author Contributions:

JR, RK, TRG, JW, WT and SSL developed the exceRpt pipeline software. JR, JJP, RK, and JD performed the analysis. JR and JJP wrote the manuscript. AM and MG supervised the project. All authors read and approved the contents of the paper.

Acknowledgements:

We acknowledge support from the NIH Common Fund for the Extracellular RNA Communication Consortium (ERCC) and from the AL Williams Professorship funds. JJP was supported by the NIH Medical Scientist Training Program Training Grant (T32GM007205).

References:

- Akat, K.M., Moore-McGriff, D., Morozov, P., Brown, M., Gogakos, T., Correa Da Rosa, J., Mihailovic, A., Sauer, M., Ji, R., Ramarathnam, A., *et al.* (2014). Comparative RNA-sequencing analysis of myocardial and circulating small RNAs in human heart failure and their utility as biomarkers. *Proc Natl Acad Sci U S A* *111*, 11151-11156.
- Byron, S.A., Van Keuren-Jensen, K.R., Engelthaler, D.M., Carpten, J.D., and Craig, D.W. (2016). Translating RNA sequencing into clinical diagnostics: opportunities and challenges. *Nat Rev Genet* *17*, 257-271.
- Chan, P.P., and Lowe, T.M. (2009). GtRNADB: a database of transfer RNA genes detected in genomic sequence. *Nucleic Acids Res* *37*, D93-97.
- Cole, J.R., Wang, Q., Fish, J.A., Chai, B., McGarrell, D.M., Sun, Y., Brown, C.T., Porras-Alfaro, A., Kuske, C.R., and Tiedje, J.M. (2014). Ribosomal Database Project: data and tools for high throughput rRNA analysis. *Nucleic Acids Res* *42*, D633-642.
- Danielson, K.M., Rubio, R., Abderazzaq, F., Das, S., and Wang, Y.E. (2017). High Throughput Sequencing of Extracellular RNA from Human Plasma. *PLoS One* *12*, e0164644.
- Dobin, A., Davis, C.A., Schlesinger, F., Drenkow, J., Zaleski, C., Jha, S., Batut, P., Chaisson, M., and Gingeras, T.R. (2013). STAR: ultrafast universal RNA-seq aligner. *Bioinformatics* *29*, 15-21.
- Freedman, J.E., Gerstein, M., Mick, E., Rozowsky, J., Levy, D., Kitchen, R., Das, S., Shah, R., Danielson, K., Beaulieu, L., *et al.* (2016). Diverse human extracellular RNAs are widely detected in human plasma. *Nat Commun* *7*, 11106.
- Fu, G.K., Xu, W., Wilhelmy, J., Mindrinos, M.N., Davis, R.W., Xiao, W., and Fodor, S.P. (2014). Molecular indexing enables quantitative targeted RNA sequencing and reveals poor efficiencies in standard library preparations. *Proc Natl Acad Sci U S A* *111*, 1891-1896.
- Glazar, P., Papavasileiou, P., and Rajewsky, N. (2014). circBase: a database for circular RNAs. *RNA* *20*, 1666-1670.
- Griffiths-Jones, S. (2004). The microRNA Registry. *Nucleic Acids Res* *32*, D109-111.
- Harrow, J., Frankish, A., Gonzalez, J.M., Tapanari, E., Diekhans, M., Kokocinski, F., Aken, B.L., Barrell, D., Zadissa, A., Searle, S., *et al.* (2012). GENCODE: the reference human genome annotation for The ENCODE Project. *Genome Res* *22*, 1760-1774.
- Hasan, N.A., Young, B.A., Minard-Smith, A.T., Saeed, K., Li, H., Heizer, E.M., McMillan, N.J., Isom, R., Abdullah, A.S., Bornman, D.M., *et al.* (2014). Microbial community profiling of human saliva using shotgun metagenomic sequencing. *PLoS One* *9*, e97699.
- Langmead, B., and Salzberg, S.L. (2012). Fast gapped-read alignment with Bowtie 2. *Nat Methods* *9*, 357-359.
- Lasser, C., Shelke, G.V., Yeri, A., Kim, D.K., Crescitelli, R., Raimondo, S., Sjostrand, M., Gho, Y.S., Van Keuren Jensen, K., and Lotvall, J. (2017). Two distinct extracellular RNA signatures released by a single cell type identified by microarray and next-generation sequencing. *RNA Biol* *14*, 58-72.
- Li, H., Handsaker, B., Wysoker, A., Fennell, T., Ruan, J., Homer, N., Marth, G., Abecasis, G., Durbin, R., and Genome Project Data Processing, S. (2009). The Sequence Alignment/Map format and SAMtools. *Bioinformatics* *25*, 2078-2079.

- McKiernan, J., Donovan, M.J., O'Neill, V., Bentink, S., Noerholm, M., Belzer, S., Skog, J., Kattan, M.W., Partin, A., Andriole, G., *et al.* (2016). A Novel Urine Exosome Gene Expression Assay to Predict High-grade Prostate Cancer at Initial Biopsy. *JAMA Oncol* 2, 882-889.
- Mudge, J.M., and Harrow, J. (2015). Creating reference gene annotation for the mouse C57BL6/J genome assembly. *Mamm Genome* 26, 366-378.
- Patton, J.G., Franklin, J.L., Weaver, A.M., Vickers, K., Zhang, B., Coffey, R.J., Ansel, K.M., Blemloch, R., Goga, A., Huang, B., *et al.* (2015). Biogenesis, delivery, and function of extracellular RNA. *J Extracell Vesicles* 4, 27494.
- Sai Lakshmi, S., and Agrawal, S. (2008). piRNABank: a web resource on classified and clustered Piwi-interacting RNAs. *Nucleic Acids Res* 36, D173-177.
- Skog, J., Wurdinger, T., van Rijn, S., Meijer, D.H., Gainche, L., Sena-Esteves, M., Curry, W.T., Jr., Carter, B.S., Krichevsky, A.M., and Breakefield, X.O. (2008). Glioblastoma microvesicles transport RNA and proteins that promote tumour growth and provide diagnostic biomarkers. *Nat Cell Biol* 10, 1470-1476.
- Subramanian, S.L., Kitchen, R.R., Alexander, R., Carter, B.S., Cheung, K.H., Laurent, L.C., Pico, A., Roberts, L.R., Roth, M.E., Rozowsky, J.S., *et al.* (2015). Integration of extracellular RNA profiling data using metadata, biomedical ontologies and Linked Data technologies. *J Extracell Vesicles* 4, 27497.
- Tataruch-Weinert, D., Musante, L., Kretz, O., and Holthofer, H. (2016). Urinary extracellular vesicles for RNA extraction: optimization of a protocol devoid of prokaryote contamination. *J Extracell Vesicles* 5, 30281.
- Yanez-Mo, M., Siljander, P.R., Andreu, Z., Zavec, A.B., Borrás, F.E., Buzas, E.I., Buzas, K., Casal, E., Cappello, F., Carvalho, J., *et al.* (2015). Biological properties of extracellular vesicles and their physiological functions. *J Extracell Vesicles* 4, 27066.
- Yeri, A., Courtright, A., Reiman, R., Carlson, E., Beecroft, T., Janss, A., Siniard, A., Richholt, R., Balak, C., Rozowsky, J., *et al.* (2017). Total Extracellular Small RNA Profiles from Plasma, Saliva, and Urine of Healthy Subjects. *Sci Rep* 7, 44061.
- Yuan, T., Huang, X., Woodcock, M., Du, M., Dittmar, R., Wang, Y., Tsai, S., Kohli, M., Boardman, L., Patel, T., *et al.* (2016). Plasma extracellular RNA profiles in healthy and cancer patients. *Sci Rep* 6, 19413.