Rapid and sensitive single-cell RNA sequencing with SHERRY2

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Abstract

Background: Prevalent single-cell transcriptomic profiling (scRNA-seq) methods are mainly based on the synthesis and enrichment of full-length double-stranded complementary DNA. These approaches are challenging to generate accurate quantification of transcripts when their abundance is low or their full-length amplifications are difficult.

Results: Based on our previous finding that Tn5 transposase can directly cut-and-tag DNA/RNA hetero-duplexes, we present SHERRY2, a specifically optimized protocol for scRNA-seq without second-strand cDNA synthesis. SHERRY2 is free of pre-amplification and eliminates the sequence-dependent bias. In comparison with other widely used scRNA-seq methods, SHERRY2 exhibits significantly higher sensitivity and accuracy even for single nuclei. Besides, SHERRY2 is simple and robust and can be easily scaled up to high-throughput experiments. When testing single lymphocytes and neuron nuclei, SHERRY2 not only obtained accurate countings of transcription factors and long non-coding RNAs, but also provided bias-free results that enriched genes in specific cellular components or functions, which outperformed other protocols. With a few thousand cells sequenced by SHERRY2, we confirmed the expression and dynamics of Myc in different cell types of germinal centers, which were previously only revealed by gene-specific amplification methods.

Conclusions: SHERRY2 is able to provide high sensitivity, high accuracy, and high throughput for those applications that require a high number of genes identified in each cell. It can reveal the subtle transcriptomic difference between cells and facilitate important biological discoveries.

Keywords: Single cell, RNA-seq, Tn5 transposase

Background

Many experimental methods for transcriptome profiling by next-generation sequencing (RNA-seq) have been developed to cover various scales of input samples, ranging from bulk samples to single cells or even subcellular components. High-quality single-cell RNA-seq (scRNA-seq) data can be used to reveal the kinetic details of gene expression and transitions between cell states or types. Prevalent scRNA-seq methods mainly rely on template switching and pre-amplification of complementary DNA (cDNA). However, large-scale scRNA-seq techniques, commonly operated in microdroplets or wells, have relatively low sensitivity. Single tube-based scRNA-seq approaches can typically...
produce higher coverage for low-abundance genes, but they still suffer from quantification bias due to insufficient reverse transcription and GC imbalance during amplification. Besides, their complex experimental methods are generally unsuitable for large-scale studies.

We have reported a highly reproducible and rapid library preparation method for RNA-seq, SHERRY, which can be applied to a minute amount of RNA samples [12]. The development of SHERRY was based on the recent discovery that Tn5 transposase can bind and cut RNA/DNA hetero-duplexes directly. With slight modifications, SHERRY could also be applied to various clinical metatranscriptome applications, such as the identification of SARS-CoV-2 and other pathogens [13].

Although SHERRY was applied to process single cells and achieved less biased quantification of gene expression in comparison with other scRNA-seq methods, the results still exhibited clear coverage bias toward the 3′-ends of transcripts, relatively low sensitivity, and low tolerance to endogenous DNA. In this work, we present an optimized method, SHERRY2, which addresses the limitations of SHERRY and is fully compatible with single cells and single nuclei with low RNA content. In comparison with prevalent RNA-seq methods, SHERRY2 showed higher sensitivity, better concordance with reference data, greater reproducibility between replicates, and superior scalability, allowing the method to be used to process a few thousand single cells per batch and thus reducing the time required to conduct experiments.

Results

SHERRY2 provides high sensitivity and even coverage across gene bodies for scRNA-seq

For scRNA-seq, RNA degradation and incompleteness of reverse transcription (RT) are two major factors that reduce gene detection sensitivity and coverage evenness. Although adding random RT primers facilitates the coverage of long transcripts, it requires the removal of ribosomal RNA, which is incompatible with scRNA-seq [13]. Spiking template-switching oligonucleotides also provides more uniform coverage, but this strategy has limited detection sensitivity and specificity [12].

We altered various experimental parameters of the original SHERRY protocol for both bulk (Additional file 1: Fig. S1-S2, Additional files 2 and 3) and single-cell inputs (Additional file 1: Fig. S3A). To protect RNA from degradation, we lowered the concentration of free Mg$^{2+}$, either by reducing the amount of total Mg$^{2+}$ or adding more dNTP to chelate Mg$^{2+}$ ions [14], and observed significant improvement in the coverage evenness of RNA-seq. To facilitate cDNA synthesis, we screened different reverse transcriptases and found that SuperScript IV (SSIV), working at a relatively high temperature with a low Mg$^{2+}$ concentration, could better overcome the secondary structure of RNA and hence simultaneously enhanced the sensitivity and uniformity of RNA-seq.

When RNA-seq was conducted using pictogram-level RNA inputs, sufficient amount of Tn5 transposase was important for high sensitivity, and Bst 3.0 DNA polymerase filled the gap left by Tn5 tagmentation more effectively than other enzymes. The protocol was insensitive to many experimental conditions, including the usage of single-strand DNA-binding proteins [15], the Tn5 inactivation, the concentration of extension polymerase, and the usage of hot-start polymerase.

We named this optimized protocol SHERRY2. Using RNA extracted from HEK293T cells as input, we compared the performance of SHERRY2 and the original SHERRY protocol. At the 10-ng level, both protocols identified more than 11,000 genes at saturation. At the 100-pg level, SHERRY2 performed better than SHERRY and detected 5.0% more genes at 0.6 million reads (Additional file 1: Fig. S2A). In addition, SHERRY2 greatly diminished 3′-end coverage bias (Additional file 1: Fig. S2B) and increased the unique mapping rate for 10-ng and 100-pg inputs (Additional file 1: Fig. S2C). We also constructed a bias-free RNA-seq library using 200-ng total RNA input via the conventional fragmentation-adorption method with the NEBNext E7770 kit (NEBNext). For 100-pg input, the gene overlap between NEBNext and SHERRY2 was greater than that between NEBNext and SHERRY (81.7% vs 78.4%) (Additional file 1: Fig. S2D), and the gene expression results of NEBNext and SHERRY2 were also more closely correlated ($R = 0.70$ vs $R = 0.65$) (Additional file 1: Fig. S2E).

The SHERRY2 protocol for scRNA-seq contains only four steps: reverse transcription, Tn5 tagmentation, gap-filling through extension, and PCR amplification. The entire SHERRY2 protocol can be completed within 3 h, 1 h less than the original SHERRY protocol, and still held its competence in costs (Additional file 1: Fig. S3B). Other high-sensitivity scRNA-seq methods such as SmartSeq2 may require much more time and more steps to be completed [3] (Fig. 1A). The one-tube workflow of SHERRY2 is readily scalable to high-throughput applications. SHERRY2 was able to detect 10,024 genes (FPKM > 1) on average within a single HEK293T cell at 1 million reads. When subsampling to 0.2 million reads, SHERRY2 still detected 8504 genes on average, which was 1622 (23.6%) more than SHERRY and 886 (11.6%) more than SmartSeq2 (Fig. 1B). In addition, the reproducibility of SHERRY2 was significantly higher than that of SHERRY or SmartSeq2 (Fig. 1C) due to its simplified workflow and stable performance. Moreover, the evenness of gene body coverage for SHERRY2 was much higher than that of the original SHERRY protocol (0.84 vs 0.72) and was
comparable to that of SmartSeq2 (0.84) (Fig. 1D). The exonic rate of SHERRY2 was also improved in comparison with that of SHERRY, likely due to the higher RT efficiency of the newly developed method (Fig. 1E).

Last but not least, scRNA-seq with SHERRY2 exhibited superior accuracy, as demonstrated by the significantly higher correlation between the SHERRY2 gene expression results and NEBNext libraries in comparison with that of SmartSeq2 ($R = 0.71$ vs $R = 0.67$) (Fig. 1F), since NEBNext fragmented mRNA before cDNA synthesis and amplified cDNA with very limited cycles which theoretically resulted in negligible bias at the transcriptome level. Especially, SHERRY2 showed high tolerance to GC content and was insensitive to the length of transcripts (Additional file 1: Fig. S4). Unlike SmartSeq2, for which the gene overlap and expression correlation with bulk RNA-seq showed clear declines when GC content was greater than 40%, SHERRY2 maintained these parameters at high and constant levels (82.6% overlap and $R = 0.76$) until the GC content reached 60%. Transcript length did not influence the accuracy of SHERRY2 or SmartSeq2, although SmartSeq2 exhibited a small degree of intolerance for transcripts longer than 800 bases.

**scRNA-seq for low RNA content cells**

For low RNA content cells, such as immune cells [16], we found that removal of intergenic DNA contaminations by DNase treatment was especially crucial for SHERRY2 scRNA-seq. In such cells, the open DNA regions of disassembled chromatin might be favored over DNA/DNA hybrids during Tn5 tagmentation. When DNase was omitted from the SHERRY2 protocol, more than 50% of

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**Fig. 1** The workflow and general performance of SHERRY2 on single-cell RNA-seq. A The workflow of SHERRY2 for scRNA-seq. Poly(A) tailed RNA is firstly released from single cells and reverse transcribed. The resulting RNA/cDNA hetero-duplex is then tagmented by Tn5 transposome, followed by gap repair and indexed PCR. Optionally, chromatin can be digested during lysis. The entire protocol is performed in one tube and takes 3 h. B Gene number (FPKM > 1) with SmartSeq2, SHERRY2, and SHERRY when subsampling reads to 0.1, 0.2, 0.4, 0.6, 0.8, and 1 million reads. C Pairwise correlation of gene expression within replicates for the three scRNA-seq protocols. The correlation $R$-value was calculated by a linear fitting model with normalized counts of overlapped genes. D Gene body coverage detected by the three scRNA-seq protocols. The gray region represents the standard deviation of the normalized depth among replicates. E Components of reads that were mapped to different regions of the genome using the three scRNA-seq protocols. The error bars show the standard deviation. F Gene expression correlation between single HEK293T cells and 200-ng RNA extracted from HEK293T cells. Single-cell data were acquired by the three scRNA-seq protocols. Bulk RNA results were acquired by the standard NEBNext protocol. The correlation $R$-value was calculated by a linear fitting model with normalized gene counts. The samples in B–F are single HEK293T cells. The $p$-values in B, C, and F were calculated by the Mann-Whitney U test.
reads sequenced from single mouse lymphocytes (Additional file 1: Fig. S5A) were mapped to intergenic regions, and only around 10% of reads were exonic reads (Fig. 2A).

Different DNases performed differently in SHERRY2 scRNA-seq. We tested five DNases (Additional file 1: Fig. S6A) and found three (NEB, Ambion, and TURBO DNase I) that worked and inactivated at higher temperatures increasing the intergenic rate unexpectedly, and this effect was probably due to RNA degradation at high temperatures with excess Mg2+ in the reaction buffer. In contrast, AG DNase I and gDW mix, which worked at room temperature, yielded ideal results.

We confirmed that all the five DNases could digest more than 99.5% of DNA (30 ng) by simply utilizing divalent ions of their respective storage buffer (Additional file 1: Fig. S6B, Additional file 4). Without adding extra divalent ions, the intergenic rates of single germline genes, and it contains diverse cell types with complex dynamics. Histologically, the GC can be separated into two micro-compartments, the dark zone and the light zone [17, 18]. By surface phenotyping, cells in the two compartments can be distinguished through CXCR4, CD83, and CD86 markers [19–21], with light zone cells being CXCR4hiCD83loCD86hi while dark zone cells CXCR4loCD83hiCD86lo. GC cell cycle between the dark zone and light zone states. Dark zone cells are highly proliferative and undergo somatic hypermutation, which generates a range of affinities against antigens. In the light zone, these B cells compete with each other for survival factors and help signals, which are mainly derived from follicular helper T cells. Those B cells that have acquired higher-affinity B cell receptors are selected to differentiate into plasma cells (PC) or memory B cells (MBC) or cycle back to the dark zone [18, 22–24]. Recently, a gray zone, consisting of CXCR4hiCD83hi cells with distinct gene expression patterns, was discovered and found to be involved in GC recycling [25]. The complex spatiotemporal dynamics of the GC and their underlying mechanisms are incompletely understood. To this end, sensitive scRNA-seq methods that can be used to detect gene expression with less bias are highly desirable.

We profiled 1248 sorted CXCR4hiCD86hi GC light zone cells with SHERRY2, and 1231 (98.6%) high-quality cells were retained for downstream analysis (Additional file 1: Fig. S5B). The gene expression levels of Cd19, Ccnd3, Fas, Cd86, and Cxcr4 were consistent with flow cytometry gating (Additional file 1: Fig. S7A), and no batch effect was observed (Additional file 1: Fig. S7B).

Unsupervised clustering identified seven clusters (Fig. 3A), two of which belonged to the gray zone, which was defined by co-expression of Cxcr4 and Cd83, as well as the ongoing cell division (enriched Ccnb1) [25] (Fig. 3B). We observed the expected downregulation of Bcl6 and S1pr2, the signature genes of GC B cells [26, 27], in memory B cell precursors (MPs) and plasma cell precursors (PPs). Specifically, Ccr6 was exclusively enriched in MPs [28], while Irf4 was upregulated in PPs, which was known to be mediated by the NF-kB pathway downstream of Cd40 stimulation [24]. It is worth noting that our results exhibited such Cd40 signaling effects as well (Additional file 1: Fig. S7C). Besides, Icam1 and Slam1 which were reported to be activated by Cd40 [29] were also observed (Additional file 1: Fig. S7D, Additional file 5). The relatively low expression levels of Prdm1 (not shown) and Gpr183 in PPs were consistent with the early stage of plasma cell development. In total, 1071 genes significantly up- or downregulated in specific clusters were identified.

The high sensitivity of SHERRY2 enabled the detection of Myc in 588 (47.8%) single GC light zone B cells. Using fluorescent protein reporting, Myc was found to mark light zone cells destined for dark zone re-entry [30], although Myc expression per se had been difficult to identify in specific cell types by low-sensitivity scRNA-seq approaches [31]. Consistent with previous findings [25, 29], the Myc expression was significantly higher in PPs (Fig. 3B, Additional file 1: Fig. S7E) and active in the gray zone cells (Fig. 3C). Light zone-1 also had a relatively
Fig. 2  scRNA-seq of low RNA-content samples with SHERRY2  A Proportions of genome regions covered by reads from SHERRY2 without DNase treatment, SHERRY2 with AG DNase I addition, SHERRY2 with AG DNase I and DNA carrier addition, and SmartSeq2  B Gene number (FPKM > 1) detected by SHERRY2 with AG DNase I addition, SHERRY2 with AG DNase I and DNA carrier addition, and SmartSeq2 when subsampling to 20, 50, 100, 200, 400, and 600 thousand reads. Only samples with an intergenic rate lower than 25% were counted. Samples in A and B were single lymphocyte cells from murine eyeball blood  C Library quality of SHERRY2 tested with different DNases, including gene number (FPKM > 1) at 0.25 million reads, coverage uniformity across gene body, and percentage of reads that were mapped to intergenic regions. The labels below the figure indicate the amounts and names of the DNases, as well as the EDTA concentration that was added during DNase inactivation. SmartSeq2 was also performed as a reference  D Components of reads covering different genome regions detected by SHERRY2 without DNase treatment, SHERRY2 with optimized AG DNases, and SmartSeq2  E Gene body coverage detected by SHERRY2 (with AG DNase I) and SmartSeq2. The gray region shows the standard deviation of the normalized depth among replicates  F Gene number (FPKM > 1) detected by SHERRY2 (with AG DNase I and DNA carrier) and SmartSeq2 when subsampling to 20, 50, 100, 200, 400, and 600 thousand reads  G Gene Ontology analysis of genes that were only detected by SHERRY2 (left) or SmartSeq2 (right). The top 20 most commonly occurred GO terms were shown. Samples in C-G were single B cells isolated from murine GC light zones. The p-values in B and F were calculated by the Mann-Whitney U test. The error bars in A and D show the standard deviation
higher portion of Myc+ cells, which are probably those destined for cyclic re-entry to the dark zone [30]. MPs also contained some cells that expressed Myc.

RNA velocity analysis (Fig. 3D) suggested that light zone-1 contained cells selected for dark zone re-entry, which were migrating to the gray zone and had Myc expression characterized by burst kinetics (Additional file 1: Fig. S7F). In addition, cells that appeared to have just entered the light zone were also identified. A few velocity vectors that moved to MPs were mixed in PPs, and these vectors were in the same direction as the downregulation of Myc. According to the velocity analysis, the aforementioned Myc-expressing MPs seemed to have a tendency to cycle back to the GC, suggesting that some MPs with Myc upregulation have the potential to re-participate in GC reactions.

We then assembled the BCR sequence for each cell to screen the usage of the IgH variable sequences, which were assigned in 1101 (89.4%) cells. As expected [32], IGHV1-72 dominated the NP-reactive GC response, and the coupled light chain was mainly IgL rather than IgK (Additional file 1: Fig. S8A, S8B). In addition, we identified CDR1 and CDR2 regions in 269 (24.4%) and 493 (44.8%) cells in which the Igh variable sequences were assigned, respectively (Additional file 1: Fig. S8C).

SHERRY2 revealed the differences in the usage frequencies of exons across cell types. The usage of a particular exon (chr11: 51,601,750–51,601,890) within the Hnrnpab transcript (Fig. 3E), which is widely expressed and encodes a protein that mainly functions in processing pre-mRNAs, was significantly biased among GC clusters. As shown in Fig. 3F, light zone-1 cells favored the inclusion of this exon.
Superior performance of SHERRY2 applied in snRNA-seq

Single-nucleus RNA-seq (snRNA-seq) has gained popularity since fresh and intact single cells are challenging to obtain in many applications. Hence, we tested the performance of SHERRY2 on snRNA-seq using single nuclei isolated from HEK293T cells. SHERRY2 detected 10,137 genes (RPM > 1) on average at 1 million reads, which was 4330 (74.6%) more than SmartSeq2, demonstrating that SHERRY2 had superior sensitivity for single nuclei (Fig. 4A). SHERRY2 still exhibited superior accuracy as it was significantly more correlated with NEBNext quantification results in comparison with SmartSeq2 ($R = 0.41$ vs $R = 0.39$) (Fig. 4B).

The high accuracy and sensitivity of SHERRY2 allowed better distinction between HEK293T cells and their nuclei, which had minimal differences. We performed principal component analysis (PCA) using RNA-seq data from NEBNext, SHERRY2, and SmartSeq2 (Fig. 4C). Single cells and nuclei prepared by SHERRY2 were much closer in distance to the bulk RNA results in comparison with those prepared with SmartSeq2. In addition, the expression pattern of the differential genes identified by SHERRY2 was more similar to that of NEBNext in comparison with SmartSeq2 (Additional file 1: Fig. S9).

Furthermore, we compared the performance of these two methods with hippocampal neurons since snRNA-seq is a popular method for studies of brain tissue due

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**Fig. 4** Sensitivity and accuracy of SHERRY2. A Gene number (RPM > 1) of single HEK293T nuclei detected by SHERRY2 and SmartSeq2 when subsampling reads to 0.1, 0.2, 0.4, 0.6, 0.8, and 1 million reads. B Gene expression correlation between single HEK293T nuclei and 200-ng RNA extracted from the HEK293T nuclei. Single-nucleus data were acquired by SHERRY2 and SmartSeq2. Bulk RNA results were acquired by the standard NEBNext protocol. The correlation $R$-value was calculated by a linear fitting model with normalized gene counts. C Clustering of HEK293T cellular and nuclear RNA-seq data from SHERRY2, SmartSeq2, and NEBNext using principal component analysis. The analysis utilized differentially expressed genes (adjusted $p$-value < 1e−4 and fold change > 2) between cells and nuclei detected by NEBNext. D Gene number (RPM > 1) of single neuron nuclei detected by SHERRY2 and SmartSeq2 when subsampling reads to 0.1, 0.2, 0.4, 0.6, 0.8, and 1 million reads. The nuclei were isolated from the mouse hippocampi that were freshly prepared or previously frozen at $-80 \, ^\circ C$. E Clustering of the single hippocampal neuron nuclei visualized by UMAP plot. The snRNA-seq library was prepared by SHERRY2. The analysis utilized genes expressed (counts > 0) in more than 4 nuclei. F Marker gene expression of different cell types on UMAP plot from E. The gradient colors correspond to the normalized counts of a specific gene ranging from 0 to 1. The $p$-values in A, B, and D were calculated by the Mann-Whitney $U$ test.
to the technical challenge of isolating intact single neurons. We constructed snRNA-seq libraries of the frozen and freshly prepared hippocampus with SHERRY2 and SmartSeq2. For both samples, SHERRY2 detected significantly more genes than SmartSeq2 (6600 vs 5331 at 1 million reads for frozen samples, 6686 vs 5769 at 1 million reads for fresh samples) (Fig. 4D), and still, Smart-seq2 tended to detect genes functionized in the mitochondrion (Additional file 1: Fig. S10A). Next, we sequenced a small number of fresh hippocampal neurons (176 nuclei) (Additional file 1: Fig. S10B) with SHERRY2 and classified their cell types correctly. The nuclei were non-supervisedly clustered into 4 distinct groups (Fig. 4E), after which they were re-clustered using marker genes identified by sNuc-Seq [33] (Additional file 1: Fig. S10C). The two clustering results were highly consistent. Neurons within the dentate gyrus (DG) and CA1, which occupy large areas of the hippocampus, could be assigned to cluster 0 and cluster 1, respectively, according to the high expression of Dock10, Slc4a4, and high expression of Pex5l and Hs61t3 (Fig. 4F). However, CA3 pyramidal cells were not shown in our results, probably due to the small number of samples. Cluster 3 that was featured with enriched Arx and Lhx6 could be annotated as GABAergic cells, which migrated from medial ganglionic eminence (MGE). Except for the aforementioned markers, the expression patterns of these three clusters acquired from sNuc-seq and SHERRY2 were very similar (Additional file 1: Fig. S10D). Cluster 2 was found to consist of cells with relatively high expression of Dpp10 and Tshz2, inferring that it might be contamination of cortex neurons. Moreover, our results revealed a long non-coding RNA (lncRNA) cluster [34] containing Meg3, Rian (Meg8), and Mirg (Meg9), which showed higher density in CA1 pyramidal cells and GABAergic cells while relatively sparse in DG granule cells (Additional file 1: Fig. S10E).

Discussion

SHERRY2 is a major improvement of our previously developed SHERRY [12], a Tn5 transposase-based RNA-seq method that eliminates the second-strand complementary DNA synthesis. Although the original SHERRY protocol has shown satisfactory simplicity to construct RNA-seq libraries using low amount of starting material, the coverage bias at 3’-ends of transcripts and fragmentation-prone DNA contaminant make it challenging to work with single cells. In MINERVA [13], a derivative of SHERRY that is specifically designed to work for the metatranscriptome of COVID-19 clinical samples, we have explored the various conditions to reduce DNA coverage. In SHERRY2, we further optimized the DNA reduction process and lead to a new protocol that can work for single cells and single nuclei, providing uniform coverage of whole transcripts and resist DNA contents.

There are three major advantages that SHERRY2 holds. First, SHERRY2 exhibits superior sensitivity and accuracy compared with SmartSeq2, a prevalent scRNA-seq method. What is more, from sequencing data of single GC B cells and single neuron nuclei, we found that SmartSeq2 biasedly detected genes involved in mitochondrial components. Though more genes were obtained by SHERRY2, there was no specific functional enrichment of these genes (Fig. 2G, Additional file 1: Fig. S10A). Thus, SHERRY2 would have more chances to facilitate biological discoveries that relied on subtle changes. Recently, SmartSeq3 [35], the upgraded protocol of SmartSeq2, has been reported to increase gene detection sensitivity. We have also compared the scRNA-seq data of HEK293T cells produced by SHERRY2 and SmartSeq3. SHERRY2 is able to detect over 10,000 genes at around 1 million reads, while SmartSeq3 cannot acquire the same number of genes even at 3-fold of sequencing depth (Additional file 1: Fig. S11A). Second, SHERRY2 retains great simplicity and expeditiousness, with the entire workflow taking around 3 h and with all reactions performed in one tube. The swift experimental pipeline ensures less RNA degradation, eliminates operational errors, and saves costs of supplies and labor. Third, SHERRY2 is highly robust and scalable. Procedural simplification not only reduces error cascade through step-wise operations, but also increases the tolerance of pipetting by offering easily handled volumes, leading to a significantly higher repeatability when in comparison with SmartSeq3 (Additional file 1: Fig. S11B). Besides, SHERRY2 contains richer information about exon junctions and coding regions across full-length transcripts, probably because SmartSeq3 is specifically optimized to quantify the 5’-end of transcripts (Additional file 1: Fig. S11C, S11D). SHERRY2 can be further developed to uncover more information from single cells. The simplicity and tolerance of protocol make it an ideal component to be incorporated into multi-omics studies. Moreover, SHERRY2 actually contains the strand-specific information of the transcript since it builds libraries from RNA/DNA duplex directly. Therefore, SHERRY2 can be potentially modified to differentiate the transcriptional strand of DNA. In addition, barcoded Tn5 tagmentation [36, 37] may also be applied to SHERRY2 to realize assembling full-length RNA molecules. Interestingly, when examining reads generated by SHERRY2 and SmartSeq2, we find that the cleavage sites of Tn5 tend to exhibit different sequence biases on substrate DNA and RNA/DNA duplex, which might give hints to understand the Tn5 mechanism (Additional file 1: Fig. S12).
There are a few remaining hitches of the current SHERRY2 protocol that need to be fixed in the future. The slightly unsatisfactory mapping rate may be compensated by slightly more sequencing reads. Without cDNA enrichment, the exogenous DNA from the environment or reagents still can be introduced after the lysis step and easily tagged by Tn5, thus impairing the performance of RNA-seq of low RNA content single cells or nuclei. Besides, it is still challenging to capture the complete 5′-end regions of transcripts for the limited processivity of reverse transcriptase. For example, CDR1 and CDR2 sequences in the Igh variable regions cannot be acquired for all GC cells (Additional file 1: Fig. S8C).

Conclusions
We present SHERRY2, an RNA-seq method designed for single cells and single nuclei. SHERRY2 is based on the direct tagmentation function of Tn5 transposase for RNA/DNA hetero-duplexes and overthrows prevalent single-cell RNA-seq chemistries which typically require pre-amplification of full-length transcripts, thus greatly improving the sensitivity of gene detection and eliminating the sequence-dependent bias. As a result, SHERRY2 can reveal the expression dynamics of transcription factors and IncRNAs, both of which typically harbor essential biological functions while at low abundance. Meanwhile, SHERRY2 maintains the simplicity of operation, with the whole process completed in one pot within 3 h, and hence elevates the throughput to a few thousand single cells/nuclei per experimental batch. As the simplest protocol of large-depth scRNA-seq, SHERRY2 has been validated in various challenging samples and can be seamlessly integrated into a wide range of applications.

Methods
Cell culture
HEK293T cell line was purchased from ATCC and incubated at 37 °C with 5% CO2 in Dulbecco’s modified Eagle medium (DMEM) (Gibco, 11965092), which was supplemented with 10% fetal bovine serum (FBS) (Gibco, 1600044) and 1% penicillin-streptomycin (Gibco, 15140122). Cells were dissociated by 0.05% Trypsin-EDTA (Gibco, 25300062) at 37 °C for 4 min and washed by DPBS (Gibco, 14190136).

For DNA or RNA extractions, we took ~ 10^6 suspended cells and followed the guideline of the PureLink Genomic DNA Mini Kit (Invitrogen, K182002) or RNeasy Mini Kit (Qiagen, 74104). The extracted RNA was further dealt with 20 U DNase I (NEB, M0303) for the removal of DNA and re-purified by the RNA Clean & Concentrator-5 Kit (Zymo Research, R1015).

For single-nuclei preparation, we followed the guideline of the Nuclei EZ Prep Kit (Sigma, NUC-101) and resuspended the nuclei into DPBS. Both single cells and single nuclei were sorted by FACSaria SORP flow cytometer (BD Biosciences).

Mice
For samples of germinal center B cells, C57BL/6 mice were originally from the Jackson Laboratory. Six- to 12-week-old, age- and sex-matched mice were used for the experiments.

For samples of the hippocampus nuclei and lymphocytes, aged 2-month-old male C57BL/6 mice were used and obtained from Charles River Laboratories. All mice were maintained under specific pathogen-free conditions and used in accordance with the governmental, Tsinghua University, and Capital Medical University guidelines for animal welfare.

GC light zone B cell preparation and sorting
To generate T cell-dependent GC responses in B6 mice, 100 μg NP-KLH (Biosearch Technologies, N-5060-5) plus 1 μg LPS (Sigma, L6143) emulsified in 100 μl 50% alum (Thermo, 77161) was utilized for intraperitoneal immunization.

The spleens isolated from 4 mice 13 days post-immunization were placed on a 70-μm cell strainer (Falcon, 08-771-2), which was previously wetted with MACS buffer (1% FBS and 5mM EDTA in PBS), and minced by flat end of the plunger of 2-ml syringes (Becton Dickinson, 301940). The splenocytes were passed through the strainer with MACS buffer into a 50-ml tube. The mixed red blood cells were then lysed by ACK lysis buffer (Thermo, A1049201). The cell suspension was further incubated with biotinylated 4-hydroxy-3-iodo-5-nitrophenylacetyl (NIP)15-BSA (Biosearch Technologies, N-1027-5) for 1.5 h and enriched by anti-biotin cell isolation kit (Miltenyi Biotec, 130-090-485) to get NP-reactive cells.

The enriched cells were blocked with 20 μg/ml 2.4G2 antibody (BioXCell, BE0307) and subsequently stained with APC-Cy7 (anti-B220, BD Biosciences, 552094), PE-Cy7 (anti-CD95, BD Biosciences, 557653), eF450 (anti-GL7, eBioscience, 48-5902-82), PE (anti-CXCR4, BioLegend, 146505). Also, 7-AAD (Biotium, 40037) was stained to exclude dead cells. All staining reactions were incubated for 1.5 h and enriched by anti-biotin cell isolation kit (Miltenyi Biotec, 130-090-485) to get NP-reactive cells.

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**Lymphocyte cell preparation and sorting**

The retro-orbital blood was taken from the eyeball of ether-anesthetized mice and dipped into a K2EDTA tube (BD Vacutainer, 367525). PBS was added to dilute the blood at ~ 50%. One milliliter of diluted blood was transferred into a clean 15-ml tube and incubated with 9 ml 1× red blood cell lysing solution (BD Pharm Lyse, 555899) at room temperature for 15 min avoiding light. The resulting red blood cells were washed twice with PBS containing 1% BSA at 200 g for 5 min, followed by staining with SYTOX green (Thermo, ST020) to identify intact cells. Single lymphocytes were sorted with FACS Aria SORP flow cytometer according to the gates shown in Additional file 1: Fig. S5A.

**Hippocampal nuclei preparation and sorting**

The isolated hippocampus tissue was transferred into a dounce homogenizer (Sigma, D9383) containing 2 ml of EZ Lysis Buffer (Sigma, NUC-101). The tissue was carefully dounced for 22 times with pestle A followed by 22 times with pestle B then transferred to a 15-ml tube. Next, 1 ml of EZ lysis buffer was added into the Dounce homogenizer to resuspend the residual nuclei then transferred to the same 15-ml tube. The samples were centrifuged at 300 g for 5 min. The supernatant was removed, and the pellet was resuspended in 100 μl of ice-cold PBS (Gibco, 10010023) with 1% BSA (NEB, B9000S) and 20 μM random primers (Thermo, SO181), and 1 mM dNTP containing 1 μl DNA was mixed with 22 μl reaction buffer containing 1× phi29 reaction buffer (NEB, M0269S), 20 μM random primers (Thermo, SO181), and 1 mM dNTP (NEB, N04465, and N04595), then they were incubated at 98 °C for 3 min and immediately cooled down at 4 °C for 20 min. Two microliters of phi29 DNA polymerase was added, and the amplification was carried out at 30 °C for 5 h, terminated at 65 °C for 10 min. The products were purified by Zymo columns.

**Generation of RNA-seq library**

We constructed NEBNext libraries with 200- and 10-ng RNA by following the guideline of the NEBNext Ultra II RNA Library Prep Kit for Illumina Kit (NEB, E7770). SmartSeq2 libraries with single cells were prepared following the protocol that was reported by Picelli et al. [3]. 10X libraries of 10,000 single hippocampal nuclei were constructed by Chromium Single Cell 3′ Reagent Kits (v3.1).

For scRNA-seq library of SHERRY2, single cells were sorted into 96-well plates containing 2 μl lysis buffer which consisted of 0.5% Triton X-100 (Sigma, T9284), 2 U SUPERaseln RNase Inhibitor (Thermo, AM2694), and 0.2 U AG DNase I (Thermo, 18068015). The plates were immediately spun down and incubated at 20 °C for 10 min for DNA digestion. The plates could be stored at ~80 °C or proceeded with the next step. Two microliters of inactivation buffer containing 5 μM OligodT3 (Sigma, Sangon), 5 mM dNTPs, and 1 mM EDTA (Thermo, AM9260G) was then added, and the reaction was incubated at 65 °C for 10 min and 72 °C for 3 min to facilitate RNA denaturation at the same time. Next, RT was performed by adding 6 μl RT mix (70U SuperScript IV (Thermo, 18090050), 1.7× SSIV buffer, 8.3 mM DTT, 10 U RRI, and 1.7 M Betaine (Sigma, B0300)); incubated at 50 °C for 50 min; and then inactivated the reverse transcriptase at 80 °C for 10 min. The resulting RNA/DNA hybrids mixed with 10-pg DNA carriers were taggedmented by 0.05 μl TTE Mix V50 (Vazyme, TD501) at 55 °C for 30 min, by adding 10 μl reaction mix containing 2× TTE buffer (20 mM Tris-HCl (Vazyme, TD501)) and 0.2 U AG DNase I; 6 U Bst 3.0 DNA polymerase (NEB, M0374M) within 1× Q5 high-fidelity master mix was utilized to repair the gap left by V50 at 72 °C for 15 min, followed by 80 °C for 5 min to terminate the reaction. Finally, 3 μl indexed primers mix (Vazyme, TD203) and 3 μl Q5 mix were added to perform PCR amplification. PCR was cycled as follows: 98 °C 30s for initial denaturation; 21 cycles of 20 s at 98 °C, 20 s at 60°C, and 2 min at 72 °C; and 72 °C for 5 min for the final extension. The indexed products were

**DNA carrier preparation**

One hundred nanograms of pTXB1 plasmids was firstly linearized by 10 U XbaI (NEB, R0145S) at 37 °C for 1 h and purified by Zymo columns. Then, we took 0.5-ng linearized plasmids for multiple displacement amplification (MDA), with all dTTPs replaced by dUTPs. Specifically, the 1 μl DNA was mixed with 22 μl reaction buffer containing 1× phi29 reaction buffer (NEB, M0269S), 20 μM random primers (Thermo, SO181), and 1 mM dNTP (NEB, N04465, and N04595), then they were incubated at 98 °C for 3 min and immediately cooled down at 4 °C for 20 min. Two microliters of phi29 DNA polymerase was added, and the amplification was carried out at 30 °C for 5 h, terminated at 65 °C for 10 min. The products were purified by Zymo columns.
merged and purified at 0.75× with VAHTS DNA Clean Beads (Vazyme, N411). Libraries were quantified with Qubit 2.0, and their fragment length distributions were checked by the Fragment Analyzer Automated CE System. Libraries were sequenced by Illumina NextSeq 500 or NovaSeq S4.

RNA-seq data analysis

Data quality
Adaptors and poly(A/T) sequences were trimmed, and bases with quality less than 20 and reads shorter than 20 bases were removed from the raw sequencing data by Cutadapt (v1.15) [38]. Clean reads were mapped to indexed genome (human: Gencode.v31, mouse: Gencode. vM23) by STAR (2.7.1a) [39]. Only unique alignment was utilized for downstream analysis. The mitochondrial and ribosomal ratios were counted with samtools (v1.10) [40]. The ratios of the coding region, UTR, intron, and intergenic region were counted with Picard tools (v2.17.6). The exonic rate was defined as the sum of the coding region and UTR ratios. For cells, Cufflinks (v2.2.1) [41] with exon annotations of protein-coding genes were used to count the gene number (FPKM > 1). For the nuclei, genes (RPM > 1) were counted by featureCounts (v1.5.1) [42] with transcript annotations. Coverage across the gene body was calculated by RSeQC (v.2.6.4) [43]. The coverage uniformity was the integral area between the coverage curve and the x-axis normalized by 100.

Gene Ontology analysis
We used genes that were detected in cell A while missed by cell B as “study” and combined the “study” genes with genes detected by cell B as “background.” The Gene Ontology analysis was performed by GOATools (v1.2.3) [44] and repeated between every two cells from different methods. GO terms (excluding electronic annotations) with adjusted p-value less than 0.01 were counted. All cells were firstly downsampled to 500K or 1M total reads.

Clustering and marker genes
For scRNA-seq and snRNA-seq, clustering followed the basic tutorials of Scanpy (v1.8.1) [45]. The cell type annotations were through manually checking the expression of well-known marker genes. Marker genes identified by SHERRY2 should satisfy the following conditions: (1) adjusted p-values calculated by the Mann-Whitney U test were less than 1e−3, (2) fold changes were greater than 1.5 or less than 0.67, and (3) the average normalized counts of the upregulated gene in the cell type or downregulated gene in the rest of cell types were greater than 0.3. For NEBNext, DESeq2 (v1.22.2) [46] was utilized to identify the differentially expressed genes (adjusted p-value < 1e−4, fold change > 2).

RNA velocity
Splicing and unsplicing mRNA were quantified by VeloCyto (v0.17.17) [10] with unique alignment. The generated loom file was utilized by scVelo (v0.2.4) [47] to profile velocity dynamics based on the clustering results of Scanpy.

BCR assembly
BCR sequences of each cell were assembled by MIXCR (v3.0.13) [48] with clean reads. The assembled BCR were realigned by IgBlast (v1.17.1) [49] to determine the clone types.

Exon usage
The frequency of exon usage in each cell was calculated by BRIE (v2.0.5) [50]. For each exon, cells satisfying the following conditions were retained: (1) counts of a gene which included the exon were greater than 10, (2) exon regions sided by the specific exon should be covered by greater than 50% with uniquely aligned reads, and (3) at least one read should detect junctions involved in this exon splicing events. A pairwise comparison of exon usage frequency was made between cell types which contained greater than 10 cells using the Mann-Whitney U test. The exons with a p-value less than 0.05 was further checked in IGV viewer to check whether transcript coverage was consistent with usage frequency. The passed ones were considered as significantly biased among cell types.

SmartSeq3 data reanalysis
SmartSeq3 [35] sequencing data of 117 single HEK293T cells was downloaded from ArrayExpress. The UMI and tag sequences at the 5′-end were firstly removed. Merged 5′-end reads and internal reads were then analyzed using the pipeline described in the “Data quality” section.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12915-022-01416-x.

Additional file 1: Fig. S1. Conditions for bulk RNA. Fig. S2. Improved performance of SHERRY2 on bulk RNA. Fig. S3. Conditions for single cells. Fig. S4. Accuracy of SHERRY2 for single cells. Fig. S5. Flow cytometry gating of single lymphocytes and single GC B cells. Fig. S6. DNase activity and performance in SHERRY2 library construction. Fig. S7. GC information gained from SHERRY2. Fig. S8. BCR sequences of single GC B cells identified by SHERRY2. Fig. S9. Expression pattern of differential genes between cells and nuclei in SHERRY2. Fig. S10. Clustering and annotation of hippocampal nuclei. Fig. S11. Comparison of SHERRY2 and SmartSeq3. Fig. S12. Sequence bias in tagmentation of different substrates.
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Authors’ contributions

Y.H. and J.W. conceived the study. L.D., B.L., Y.L., S.Z., Y.P., and J.S. performed the experiments. L.D. performed the data analyses. C.Z. and J.S. provided the samples. L.D., B.L., J.W., H.Q., J.S., and Y.H. wrote the manuscript with input from all authors. Y.H., J.W., and H.Q. supervised all aspects of this study. All authors read and approved the final manuscript.

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Availability of data and materials

All data generated or analyzed during this study are included in this published article, its supplementary information files, and publicly available repositories. The sequence data reported in this study have been deposited in the NCBI Sequence Read Archive (assession no. PRINAB79104) [51]. Data values of Additional file 1: Figs. S1, S2, S6B, and S7D were provided in Additional file 1: Fig. S48.

Declarations

Ethics approval and consent to participate

All mice were maintained under specific pathogen-free conditions and used in accordance with governmental, Tsinghua University, and Capital Medical University guidelines for animal welfare.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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