

# 1 Rapid and Signal Crowdedness-Robust In-Situ Sequencing

## 2 through Hybrid Block Coding

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## 30 Abstract

31 Spatial transcriptomics technology has revolutionized our understanding of cell types and tissue  
32 organization, opening new possibilities for researchers to explore transcript distributions at subcellular  
33 levels. However, existing methods have limitations in resolution, sensitivity, or speed. To overcome  
34 these challenges, we introduce SPRINTseq (Spatially Resolved and signal-diluted Next-generation  
35 Targeted sequencing), an innovative in situ sequencing strategy that combines hybrid block coding and  
36 molecular dilution strategies. Our method enables fast and sensitive high-resolution data acquisition,  
37 as demonstrated by recovering over 142 million transcripts using a 108 gene panel from 453,843 cells  
38 from four mouse brain coronal slices in less than two days. Using this advanced technology, we uncover  
39 the cellular and subcellular molecular architecture of Alzheimer's disease, providing additional

1 information into abnormal cellular behaviors and their subcellular mRNA distribution. This improved  
2 spatial transcriptomics technology holds great promise for exploring complex biological processes and  
3 disease mechanisms.

4

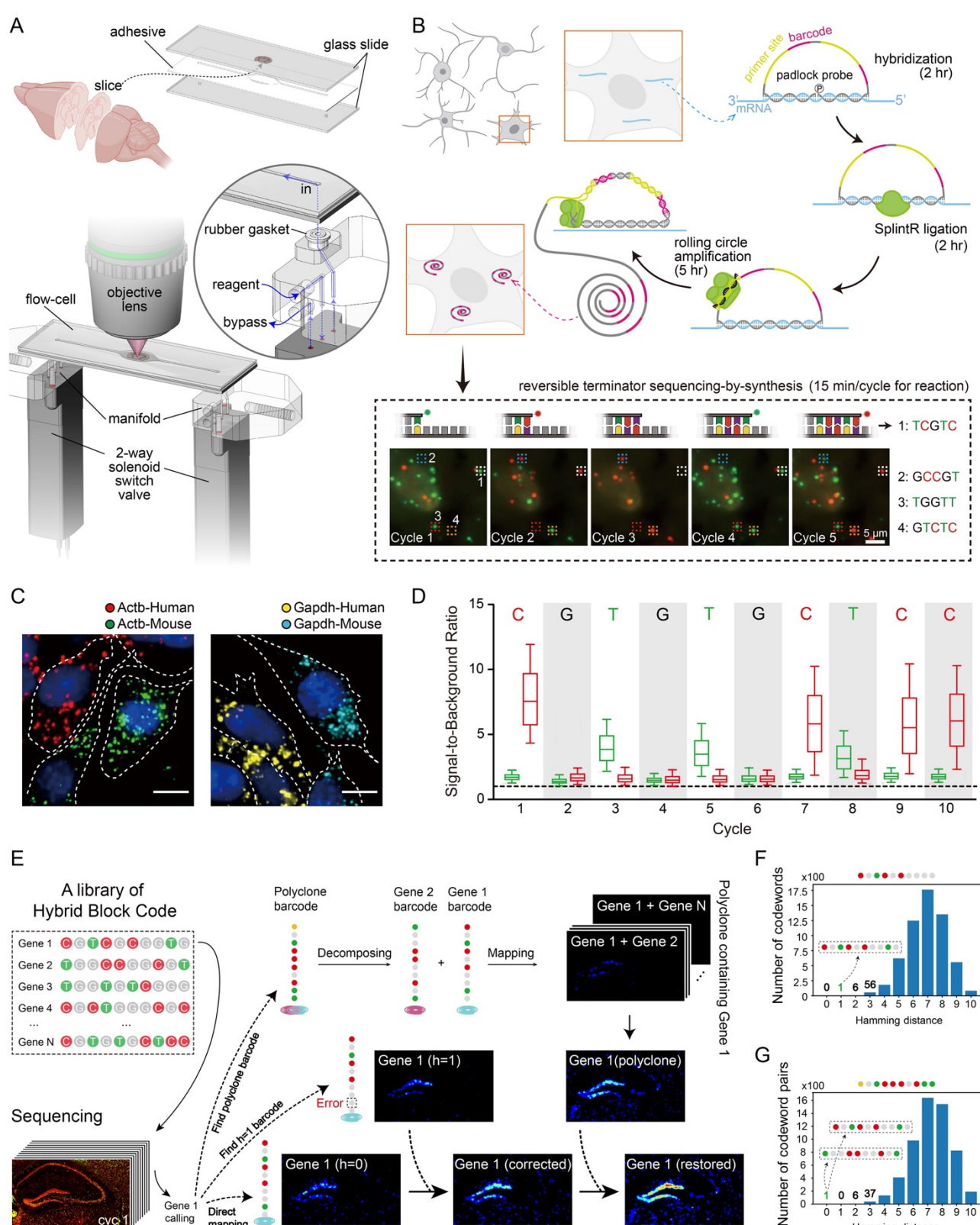
## 5 **Introduction**

6 The well-ordered structural organization of cells and the intrinsic heterogeneity among cells are  
7 essential characteristics of multi-cellular organisms. Various spatial transcriptomic technologies have  
8 been developed to help understand the nature of such spatial information and its functional properties.  
9 In-situ mRNA capture-based approaches offer a spatially-resolved expression landscape (1-5).  
10 Nevertheless, these methods have trouble providing finer information in space, such as subcellular  
11 RNA localization, whereas imaging-based spatial transcriptomic approaches offer near-optical limit  
12 resolution (6). An ideal spatial transcriptomic technology needs to be sensitive, accurate, scalable, and  
13 robust. Such currently available approaches are divided into two major categories, fluorescent in-situ  
14 hybridization (FISH) (7-11) and in-situ sequencing (ISS) (12-16). Those prevalent approaches face  
15 major challenges due to time-consuming workflows, complex reagents, and complicated imaging set-  
16 ups. FISH-based approaches are typically built upon single molecule detection schemes that rely on  
17 high-power, large numeric aperture objective lens and on specialized imaging techniques to overcome  
18 a low signal-to-background ratio (SBR) problem, and that usually leads to a sacrifice in the size of  
19 imaging field of view (FoV). The length of the mRNA of interest is also limited since many probe binding  
20 sites are commonly needed. While ISS-based methods usually apply in-situ amplification to increase  
21 the SBR ratio, molecular crowdedness during amplification or optical imaging are major challenges.  
22 Another common issue of currently available methods, the long experimental time, greatly hinders the  
23 scalability needed to handle a high-quantity of or large-size samples.

24

25 Here, we introduce a new ISS-based technology that uses sequencing-by-synthesis (SBS) chemistry  
26 to speed up the cyclic reactions needed for highly efficient information acquisition. We created a hybrid  
27 block code that is both signal-crowdedness and error robust, and that has a high decoding efficiency.  
28 Physical dilution of signal is adopted to further counteract ISS's intrinsic crowdedness issue. This  
29 method, SPatially Resolved and signal-diluted Next-generation Targeted sequencing (SPRINTseq), has  
30 greatly shortened the sequencing time it took (to within 9.5 hours) to profile a targeted transcriptome of  
31 a mouse brain coronal slice, and produced near optical diffraction-limit resolution. SPRINTseq produced  
32 a sub-micron precise, whole-slice-scale cellular atlas that contains subcellular location information for  
33 each transcript. Such an information-rich sub-cellular distribution of genes can be greatly affected by  
34 physiological conditions. Using a 108-gene panel, 4 slices of mouse coronal brain (from 2 normal  
35 mouse and 2 mouse with Alzheimer's disease) from sample to data could be profiled in 2 days, covering  
36 453,843 cells and 142,957,485 transcripts. We found that the degree of subcellular mRNA dispersion  
37 increased as glia cells activated in Alzheimer's disease. We also found that the mRNA distribution  
38 produced changes in orientation within the amyloid microenvironment. Additionally, the high

1 heterogeneity we found within inhibitory neurons may correlate with cell-type-specific responses during  
2 disease.  
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6  
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**Fig. 1. Workflow and performance of SPRINTseq: in-situ sequencing from sample to data within 20 hours. (A)** In-situ sequencing  
8 automation. Each mouse brain tissue slice was mounted on a slide and each slide was assembled into a flow cell and mounted between  
9 2 manifolds. The manifolds were controlled by 2-way solenoid switch valves that controlled the fluidic routes. The reagent flowed directly

1 into the flow cell while a bypass prevented reagent cross-contamination. **(B)** Schematic workflow: *in situ* barcoded clonal amplification and *in situ* sequencing. Probe hybridization, ligation, and rolling circle amplification were performed sequentially and took about 10 hours total. Each probe contained two identical “sequencing primer + barcode” blocks to increase the signal-to-background ratio. Barcodes were sequenced *in situ* using 2-color, reversible terminator sequencing-by-synthesis chemistry. Scale bar: 5  $\mu$ m. **(C)** SPRINTseq specificity demonstrated by inter-species gene detection on a human (HEK293T) and mouse cell (3T3) co-culture system. Scale bar: 10  $\mu$ m. **(D)** Characterization of 10-bp barcode sequencing. No obvious dephasing or signal decay was observed throughout 10 cycles. The boxes show the interquartile range, the lines in the boxes are the medians, and the bars show min/max values. The horizontal dashed line is SBR = 1. **(E)** Hybrid block codes achieve error-correctable and crowdedness-robust encoding. The barcodes are highly orthogonal between each other. A specific gene is mapped by looking for its exact barcode sequence and the one with correctable error (e.g., Hamming distance  $[h] = 1$ ). Also, all composed barcodes (composite codewords) containing that gene are corrected for mapping. By counteracting signal overlapping and sequencing error, a significant number of signals are rescued by this encoding strategy. **(F)** Error correction. When a sample readout sequence “CGTCGCGGGG” (one base error) is aligned with all barcodes (including an original barcode and a composed barcode) in the library, only one barcode (CGTCGCGGTG, Gene 1) with a Hamming distance equal to 1 is found, thus this sequence was mapped to Gene 1 and the error was corrected. **(G)** Polyclone barcode decomposition. When a sample readout sequence “AGTCCCGCTT” (a composed barcode containing two barcodes) is aligned with all the composed barcodes in the library, only one barcode pair (Gene 1: CGTCGCGGTG and Gene 2: TGGCCGGCGT) with a Hamming distance less than or equal to 1 exists. Thus, this polyclone (overlapping amplicons) can be decomposed and mapped to Gene 1 and Gene 2.

## 19 Result

### 21 Principle, operation, and performance of SPRINTseq.

22 A SPRINTseq experiment consists of two major parts, *in situ* barcoded clonal amplification and ISS. We  
23 constructed an integrated setup to automatically control fluidics, temperature, mechanical motion, and  
24 fluorescence imaging (Fig. 1A, Fig. S1 & S2). A slice of tissue was placed in a microfluidic flow-cell and  
25 various reagents were programmed to flow through it. A set of padlock probes we designed directly  
26 hybridized mRNAs with high specificity and then ligated to form a circular DNA template for rolling circle  
27 amplification (Fig. 1B). In about 10 hours, targeted genes were converted *in situ* into individual nanoball  
28 clones of corresponding barcodes. Each padlock probe contained two identical barcode blocks to  
29 double the signal of the sequencing reaction (Fig. S3), which came from fluorescent-labeled, reversible  
30 terminator nucleotide substrates. Each barcode block consisted of a gene-specific barcode and a  
31 universal sequencing primer binding sequence.

32 Because of its fast cyclic reaction time (Fig. S2; 15 min/cycle includes tissue blocking, substrate  
33 incorporation, and fluorophore cleavage steps), we used mature 2-color reversible terminator SBS  
34 chemistry for barcode readout (Fig. 1B). With an approximate  $2.2 \times 10^7$  pixel/s imaging speed, the  
35 sequencing process took only about 9.5 hours for a 10-base barcode reading on a coronal slice of  
36 mouse whole brain. The images were further processed for base calling and cell segmentation (Fig. S5  
37 & S6). Common imaging background noise caused by non-specific binding of the fluorescent nucleotide  
38 substrate was clearly reduced when free thiol groups in the tissue were blocked (Fig. S3A,B). Also, with  
39 multiple-barcode-block design (Fig. S3C-F), the SBR of SPRINTseq was significantly greater than those  
40 FISH-based methods (Fig. S7). Altogether, signal with sufficient SBR can be generated even on high  
41 auto-fluorescence tissue samples (Fig. S4). Amplified barcode clones were stably attached on the  
42 sample and drifted negligibly during the whole experimental process (Methods). Using only one probe  
43 per gene, targeting sensitivity was as high as 38% (Fig. S7D-G), mainly due to a low loss of amplified  
44 clones and no reverse transcription step in the protocol. In addition, during cyclic sequencing reactions,  
45 SPRINTseq exhibited high specificity (Fig. 1C) with very low signal decay or dephasing (Fig. 1D, Fig.

1 S8).

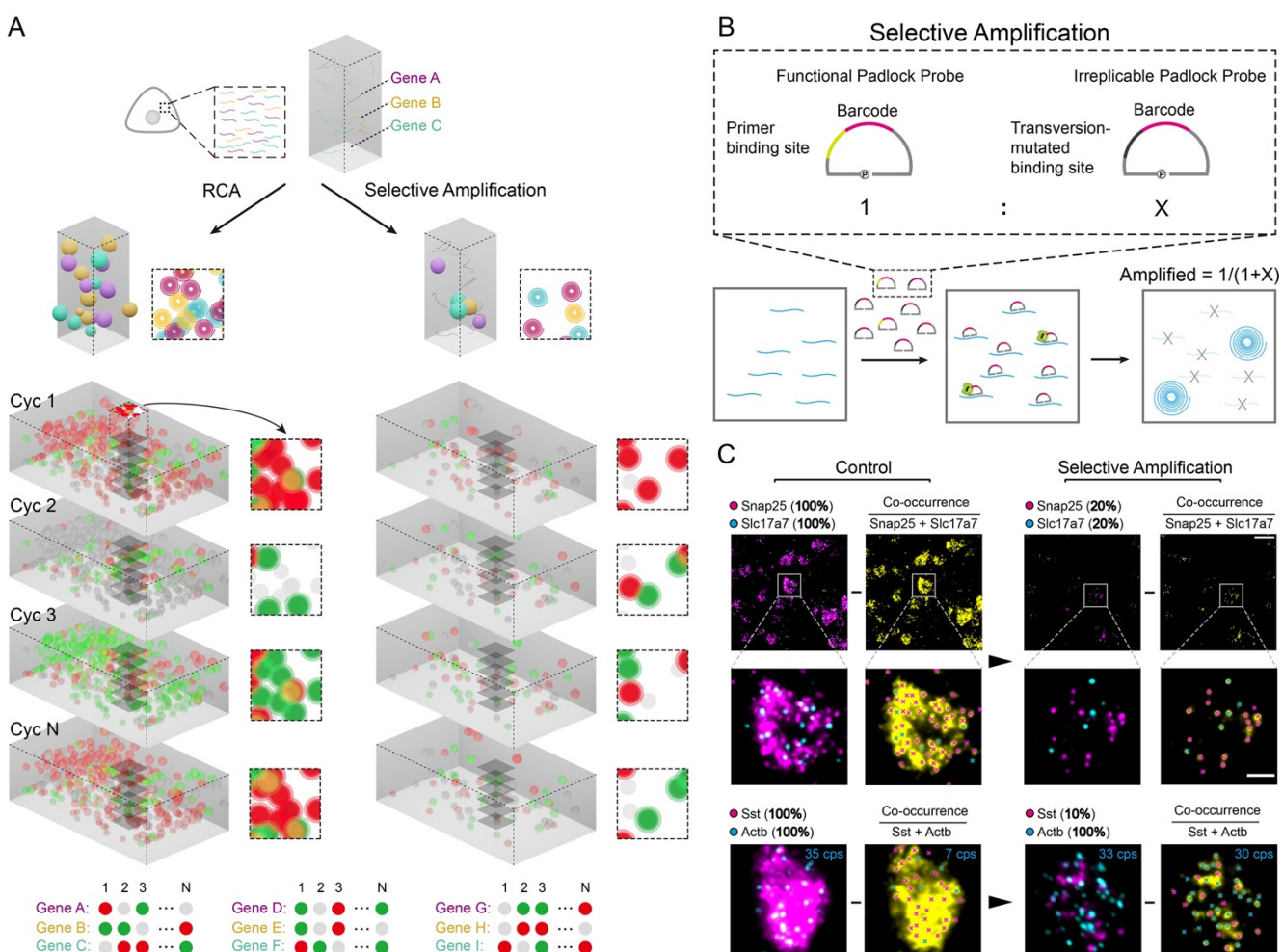
2  
3 We designed a highly efficient and orthogonal coding scheme, ‘hybrid block code’, that encodes  
4 barcodes in a way that tolerates signal crowdedness and corrects sequencing errors (Fig. 1E). We  
5 encoded  $2^k$  genes using n-bit ( $n > k$ ) codewords created from 0/1 string-based block code, and used  
6 the redundant bits to correct errors (Fig. S9). Moreover, we folded n-bit binary codewords to  $(n/2)$ -bit  
7 double-binary codewords by combining every two bits into a dual channel bit, thus shortening signal  
8 acquisition time. The dual-channel “1” signal at the same bit was avoided to reduce the overall signal  
9 density and to improve orthogonality between codewords. We then selected a subset of all codewords,  
10 among which almost any two could be uniquely split to enable the decomposition of overlapping signals.  
11 Hybrid block code naturally fits 2-color reversible terminator SBS. To convert the codeword into a DNA  
12 barcode sequence, we used the bases ‘C’ and ‘T’ to represent “1” in the two channels and base ‘G’ to  
13 represent “0”. Base ‘A’ was not used in the barcodes. After sequencing, all barcode reads were aligned  
14 against a library consisting of the original barcodes and all composed barcodes (e.g., CGG + TCG =  
15 ACG). When aligning a barcode that had one sequencing error against the library, only one barcode  
16 with a Hamming distance equal to 1 was obtained, thus achieving error correction (Fig. 1 F). Polyclone  
17 (overlapping amplicons) barcodes were aligned to all possible polyclone barcodes (all barcode pairs)  
18 in the library. Thus, a unique polyclone barcode with a Hamming distance equal to or less than 1 was  
19 found, and that allowed us to decompose the signal from overlapping amplicons (Fig. 1 G). Using  
20 barcodes called from an image sequence built with that design resulted in a significantly higher average  
21 Shannon entropy per image than those of existing methods (Table S3). Our refined experimental  
22 process and efficient coding scheme enables SPRINTseq to profile a coronal slice of mouse whole  
23 brain within 20 hours and with high sensitivity and accuracy.

24  
25 **Relieving crowdedness issues yields high quality ISS**

26 Signal crowdedness, a major challenge for ISS, occurs when numerous amplified signal spots fill the  
27 limited space within a cell and thus limit the total read count. In a swift scanning scheme, the locations  
28 of amplified clones after 2D-stacking projection are prone to signal overlap in crowded regions. In hybrid  
29 block code, though, signals are diluted by using the ‘bit-of-silence’ and ‘optical dilution’ is realized. Only  
30 a fraction of the barcodes was fluorescently labeled in each sequencing cycle (Fig. S9C). Such signal  
31 sparseness can be finetuned by the number and position of the ‘0’ signal in the barcodes, thus  
32 staggering the signals from highly expressed genes in different cycles.

33  
34 However, the optical dilution ratio has an upper limit within a certain number of sequencing cycles as  
35 the proportion of silent channels is limited by the Hamming weights of codewords. Inevitably, signal co-  
36 occurrence between highly-expressed genes will happen as more genes that need to be identified are  
37 included in the panel (Fig. 2A). In some cases, a signal from one super-highly expressed gene is so  
38 dense that the spots form ‘plaques’ and the signals from other genes are covered and their information

1 is unextractable. This crowdedness is caused by too many amplicons from highly abundant genes. Also,  
 2 some amplification events may be precluded in such a physically crowded environment, thus causing  
 3 an even greater bias in gene profiling. Optical dilution does not alleviate this issue.  
 4



5  
 6 **Fig. 2. SPRINTseq uses signal dilution to relieve signal crowdedness. (A)** Spatial crowdedness and physical dilution of signals. As  
 7 signals from rolling circle amplified (RCA) clones were projected onto a 2-D plane during focal-stacking, the signals inevitably overlapped  
 8 in crowded regions and became unrecognizable, thus causing information dropout. Through selective amplification, signals from partial  
 9 transcripts were physically eliminated, thus leaving a sparser environment. **(B)** Selective amplification. The proportion of amplification of  
 10 specific highly expressed genes can be controlled by mixing normal padlock probes and sequencing-primer binding-site transversion  
 11 probes (irreplicable), which achieve selective gene masking during amplification and yields a desired ratio. **(C)** Signal rescue through  
 12 selective amplification. Respective signals from Snap25/Sst (magenta) and Slc17a7/Actb (cyan) can be obtained through another  
 13 sequencing cycle. Because at least one base of the respective gene barcodes is designed to be the same (cycle and channel), a  
 14 crowdedness situation (co-occurrence, yellow) is simulated. Percentages represent the remaining fractions of gene signals after selective  
 15 amplification. The crosses represent the recognizable signals based on local maxima identification. In the example image, 20% (7/35) of  
 16 Actb reads can be extracted during co-occurrence with all Sst amplicons, while 91% (30/33) of the reads can be extracted during co-  
 17 occurrence after masking 90% of the Sst amplicons. Scale bars: 20 μm for the base images and 5 μm for the insets.

18  
 19 We used a selective amplification strategy to relieve physical crowdedness. For those highly expressed  
 20 transcripts, we doped the padlock probes with irreplicable ones to uniformly dilute their clonal  
 21 populations. Thus, partial transcripts of such genes are physically eliminated and other genes are able  
 22 to be better amplified and read (Fig. 2B). To demonstrate selective amplification, we simulated an  
 23 inevitable situation in which multiple genes are tested in a panel: two genes with dense signals appear

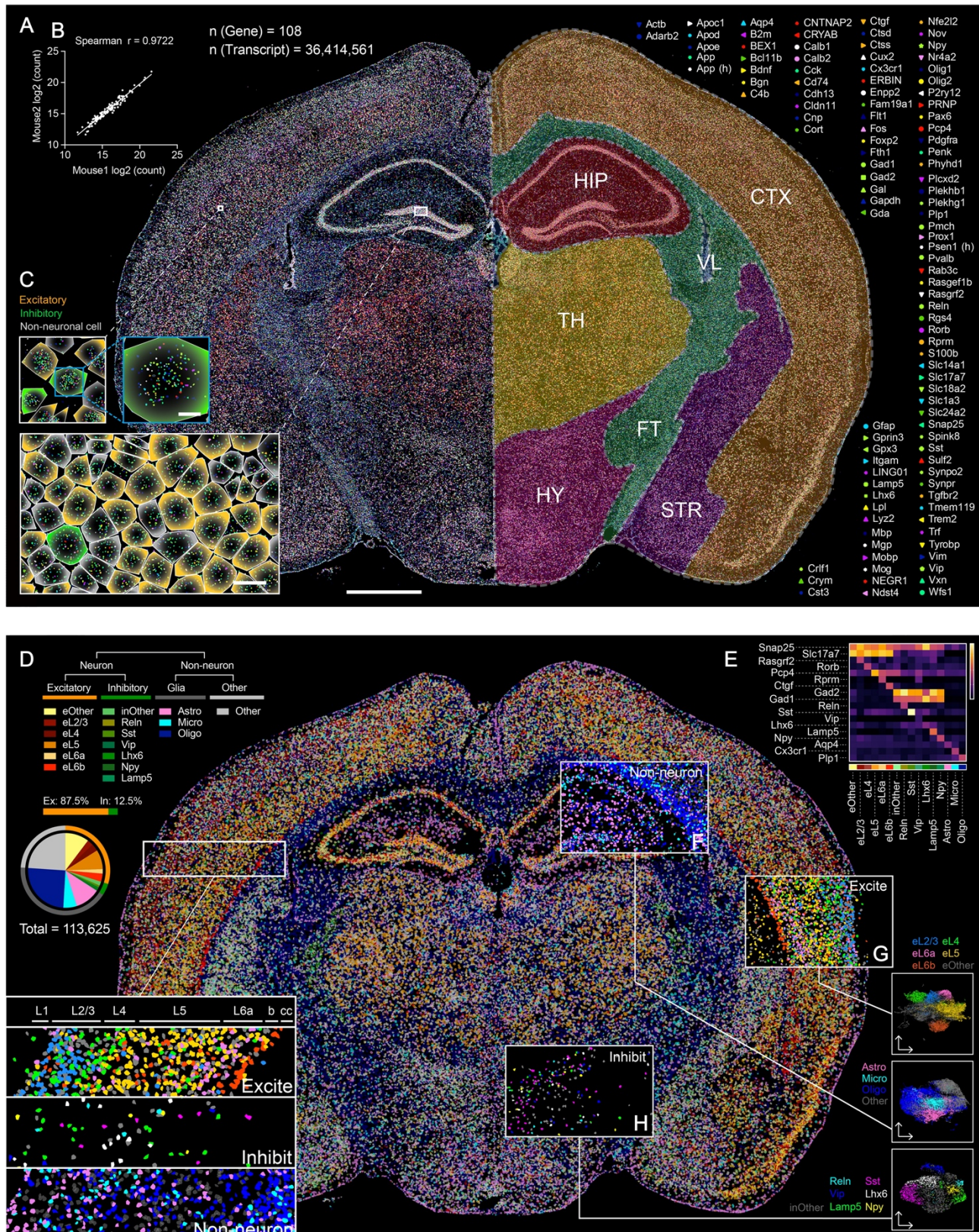
1 in the same reaction cycle and channel. Snap25 and Slc17a7 are highly expressed genes in excitatory  
2 neurons and when their signals appeared together, their clonal puncta joined together to form plaques  
3 in the images and the puncta were obscured (Fig. 2C). After masking 80% of both genes, each of their  
4 transcript clones were clearly differentiated. Though the masked signals' locations could not be  
5 recovered, the quantities of transcripts could be re-calibrated by multiplying the dilution ratios in gene-  
6 by-count expression matrix (Fig. S10). In another example, Sst, a GABAergic neuron subtype marker  
7 gene, dominated the expression in that cell type. The puncta of clonal signals from other genes such  
8 as Actb were difficult to discern. After diluting the Sst signal 10-fold, it was possible to digitally count  
9 mRNA clones, and many signals, including those of Actb and Sst, were rescued. Only 20% (7/35) of  
10 the Actb reads could be extracted when the Actb signal co-appeared with Sst's. After masking 90% of  
11 the Sst amplicon, more than 90% (30/33) of the Actb reads were extracted. Given prior knowledge, we  
12 can adjust the physical dilution ratios for genes with different abundances. This ratio needs to be high  
13 enough to identify clonal signals in highly expressed cells, but not so high as to cause false-zeroes in  
14 lowly expressed cells. Optical dilution and selective amplification are functionally complementary. The  
15 final dilution fold is the product of the physical dilution fold from selective amplification and the optical  
16 dilution fold.

17  
18 For regions that are still crowded after two-level dilution, the polyclonal signals that result mainly from  
19 the co-localized 2-D projections of two amplicons, can be decomposed into separate barcodes in hybrid  
20 block coding and their signals can be rescued (Fig. 1E,G). By combining optical dilution, physical  
21 dilution, and polyclone decomposition, SPRINTseq effectively relieves the signal crowdedness issue in  
22 ISS.

23  
24 **Single cell profiling with subcellular resolution of mouse brain coronal slices**  
25 We profiled mouse brain coronal slices (approximately  $10.2 \times 7.6 \text{ mm}^2$ ) at subcellular resolution using  
26 a 108-gene panel (Fig. 3A). The genes included both marker and disease-related genes selected for  
27 cell classification and status characterization (Extra Table 1) (17). To encode these genes, we designed  
28 a 10-base barcode set based on the hybrid block code (Fig. S5A-C). The minimum Hamming distance  
29 between any barcode pair in the 108-gene panel was 3, and the minimal Hamming distance between  
30 almost any ( $> 99.8\%$ ) barcode pairs that containing polyclone barcode was 3, which ensured error  
31 correction and polyclone decomposition. The silent bit (G) content of 52.4%, achieved a 23.8% optical  
32 dilution ratio while keeping a sufficient Hamming weight in codewords. Furthermore, selective  
33 amplification applied to seven highly-expressed genes (Actb, Mbp, Cst3, Penk, Snap25, Slc17a7, and  
34 Sst) to further reduced the clone number by 54.4%. As a result, about 10% of all signals were acquired  
35 per sequencing cycle, significantly diluting crowdedness. While 49.8% of all monoclonal reads were  
36 rescued from error correction, 31.6% of the total reads were rescued from polyclone decomposition.

37  
38 We identified 16,606,784 raw reads from the brain slice, and that number increased to 36,414,561 after

1 selective amplification was re-calibrated. Probe binding false positives and gene mapping were 0.05  
2 and 0.02 events per cell, respectively. The border of each general brain region was drawn using  
3 corresponding marker genes (Fig. 3A). Highly accurate and specific expression patterns were  
4 consistent with the in-situ hybridization results in the Allen Brain Atlas (Extra Supplement PDF). Brain  
5 replicates also showed high concordance at both whole and regional brain levels (Fig. 3B, Fig. S11).  
6



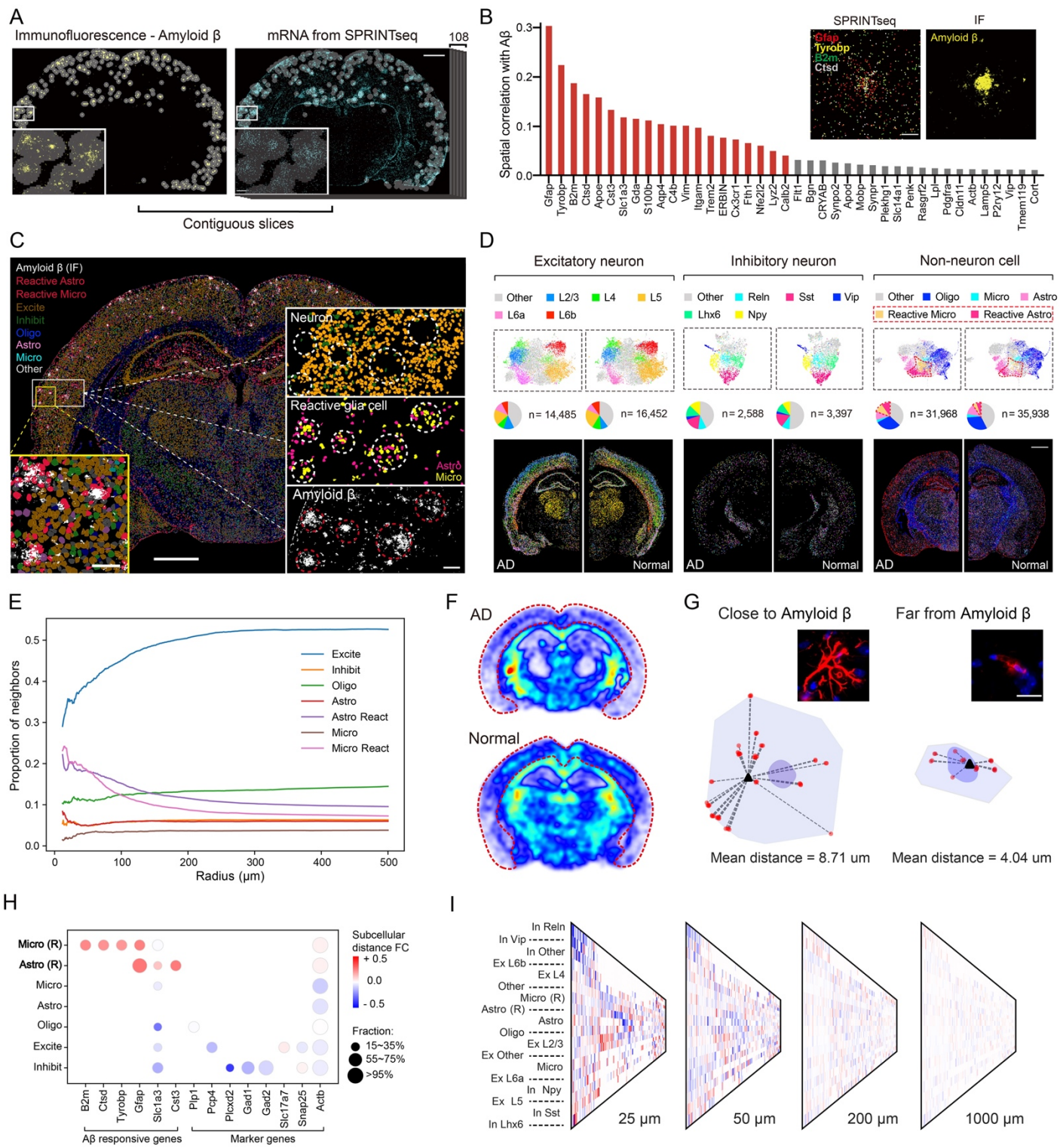
**Fig. 3. Single cell profiling with subcellular resolution of mouse brain coronal slices. (A)** The spatial landscape of 108 genes called by 10-cycle sequencing in mouse brain. Brain regions shown in the right half, the brain was defined by several regional marker genes (*Slc17a7*, *Gad2*, *Fth1*, *Enpp2*, *Pcp4*, *Pmch*), CTX: cortex, HIP: hippocampus, VL: lateral ventricle, TH: thalamus, FT: fiber tract, HY: hypothalamus, STR: striatum. Scale bar: 1 mm. **(B)** Spearman correlation of two mouse sequencing replicates, the brain slices were selected at the same position and the gene panel was the same. **(C)** Enlarged images showing gene locations (dots color/shape-coded to the genes) and cells from the insets in **(A)**. Cell border was showed by lines with different colors: orange for excitatory neuron, green for inhibitory neuron and gray for non-neuronal cell. Scale bar: 2  $\mu$ m for the single cell image and 10  $\mu$ m for the multiple cell images. **(D)**

1 Cell composition and spatial projection of a brain slice. Marker gene expression levels and Louvain shared nearest neighbor clustering  
2 identified 17 cell types (16 defined cell types and 1 other cell type shown in upper left corner), and their proportions are shown in the pie  
3 chart. Each color represents one unique cell type. The enlarged images at the bottom left show projections of all 17 cell types at the same  
4 locations in the cortex. The anatomical structure is labeled on top beginning with L1 (L: layer in the cortex, cc: corpus callosum) (E) Marker  
5 gene expression heatmap by each cell type. The cell type color code is the same as in (D). (F-H) Spatial projections of non-neuronal cells  
6 (astrocytes, microglia, and oligodendrocytes), excitatory neurons (eL2/3, eL4, eL5, eL6a, eL6b and others), and inhibitory neurons (Rhn  
7 Sst, Vip, Lhx6, Lamp5, Npy, and others), as well as their cell clustering visualizations through uniform manifold approximation plots.  
8

9 We determined cell nuclei centroids using DAPI staining, calculated cellular segmentation with nucleic  
10 images, and assigned sequenced RNA reads to their nearest nucleus centroid (Fig. 3C, Fig. S6). We  
11 obtained 146,137 cells in total and 113,625 (77.8%) passed quality-control. According to the major  
12 marker genes, cells were categorized into three major populations: excitatory neurons (Slc17a7+),  
13 inhibitory neurons (Gad1/Gad2+), and non-neuronal cells. Each population was then subdivided into  
14 detailed clusters through Louvain shared nearest neighbor clustering and the resulting 17 types were  
15 spatially projected to their original positions (Fig. 3 D & E, Fig. S12-13). Non-neuronal cells were divided  
16 into astrocyte, microglia, oligodendrocyte, and other cell types (Fig. 3F). Astrocytes and microglia are  
17 scattered across the whole brain, but astrocytes are more frequently distributed at tissue edges and  
18 the hippocampal region. Oligodendrocytes have a high density in the fiber tract region. Neuronal cells  
19 were classified according to their positions in the cortex (for excitatory neurons, Fig. 3G) and their  
20 subtypes (for inhibitory neurons, Fig. 3H). Excitatory neurons in the cortex can be clearly divided into  
21 six layers. They are also widely distributed in the hippocampus and thalamus. Inhibitory neurons are  
22 sparsely distributed in the cortex, striatum, hippocampus, hypothalamus, and reticular nucleus region  
23 of the thalamus.  
24

25 SPRINTseq offers an informative subcellular distribution and location of genes, as well as their  
26 correlations to other spatially distributed components. We assessed the degree of the mRNAs'  
27 subcellular dispersity using the average distances to their centroids, and also calculated the mRNA  
28 average distance to the nucleus centroid (Fig. S14). We then used those two parameters as coordinates  
29 for each gene in the panel and classified them all into one of three quadrants.  
30

31 The mRNAs of the genes in the first quadrant were widely dispersed within the cell and far from the  
32 nucleus. From that we inferred that they were diffused throughout the cytoplasm and expressed the  
33 protein needed inside the cell. Actb is one typical example of this type. The mRNAs of the genes in the  
34 third quadrant, such as Ctgf, were likely to be distributed on the endoplasmic reticulum near the nucleus  
35 and to express membrane proteins or secreted protein. Genes appearing in the fourth quadrant showed  
36 a certain polarity in their mRNA distributions; genes including Snap25 and Slc17a7 were in this category.  
37 The mRNA distribution polarity of signal-transduction-associated genes may be related to the polarity  
38 of neurons and other cells in the brain. No genes appeared in the second quadrant.  
39



**Fig. 4. Spatially-resolved cellular and subcellular changes of mouse brain with Alzheimer's disease (AD). (A)** Immunostained amyloid plaque (amyloid  $\beta$ ) and SPRINTseq analysis on contiguous slices. After alignment, the amyloid plaque positions were localized on the sequencing image. The 'Pan-plaque region' (regions circled in white) was defined around amyloid, and spatial correlation was calculated between all 108 genes and the amyloid plaque, respectively, in the pan-plaque regions. Bin size: 50 x 50  $\mu$ m. Scale bars: 80  $\mu$ m for enlarged images and 1 mm for brain slice images. **(B)** The most enriched genes, as shown by spatial correlation analysis. The top 20 genes in the graph are shown by red bars. The upper right images show the top 4 genes' (Gfap, Tyrobp, B2m, and Ctsd) actual aggregations (left, sequencing result) and at the same positions with amyloid plaque (right, immunofluorescence [IF], contiguous slice after alignment). Scale bar: 40  $\mu$ m. **(C)** Whole brain spatial projection for general cell types. Amyloid plaque IF in a contiguous slice is shown along with excitatory and inhibitory neurons and non-neuronal cells: oligodendrocyte (Oligo), astrocyte (Astro), microglia (Micro), and other. Reactive glia cells (responding to amyloid plaque) are labeled in red (lower left image). The 3 enlarged images on the right

1 show neuron and reactive glia cells and the amyloid plaque distributions at the same position. Scale bar: 1 mm for the whole brain image,  
2 100  $\mu$ m for lower left image and the multiple cell images on the right. (D) Normal and AD mouse brain direct cell composition comparison  
3 using combined classifications of the 16 cell types (see Fig. 3F-H). Scale bar: 1 mm (E) Relative cell density change as a function of  
4 distance to amyloid plaque. The measurements start at 11  $\mu$ m. (F) Oligodendrocyte density comparison of AD and normal mouse brain.  
5 The cortex region is encircled by a red dotted line. Oligodendrocyte density decreased globally in the AD mouse brain. (G) Diagram of  
6 subcellular dispersion. The photos show reactive astrocytes with Gfap protein in red and DAPI in blue. IF results confirm that subcellular  
7 dispersion of reactive astrocytes differed based on their distances from plaque. In the corresponding diagrams, the red dots represent  
8 Gfap mRNA, the black triangles are Gfap mRNA centroids, the pale purple shows cell body, and the dark purple shows cell nucleus.  
9 Scale bar: 10  $\mu$ m. (H) Heatmap of the degree of subcellular dispersion change. Y axis shows different general cell types as listed in (C).  
10 The values are the subcellular distance of cells close to amyloid plaque divided by that of cells far from amyloid plaque. (R): reactive, FC:  
11 fold change. (I) The orientation pattern as a function of distance. Y axis shows different cell types. The orientation of mRNAs was profiled  
12 in the cells whose distance to the nearest Amyloid  $\beta$  was 25  $\mu$ m, 50  $\mu$ m, 200  $\mu$ m and 1000  $\mu$ m, respectively.

## 13 Spatial profiling of the mouse brain with Alzheimer's disease

14 The 108-gene panel included many genes associated with Alzheimer's disease (AD), which is also  
15 pathologically associated with the distinct spatial distribution of amyloid plaques (Amyloid  $\beta$ ) that  
16 accumulate in the cortex and hippocampus, especially during aging (18). We used SPRINTseq to  
17 examine brain coronal slices from 10-month-old APP/PS1 and normal mice. First, the genes' overall  
18 expression levels were similar between AD and normal mice at the whole brain slice and regional levels  
19 (Fig. S15A). Next, we investigated gene-expression changes around amyloid plaques, whose typical  
20 size was over 100  $\mu$ m in diameter. We aligned the Amyloid  $\beta$  immunofluorescence images to our  
21 SPRINTseq images and defined the plaques and their surrounding areas as 'pan-plaque regions' (Fig.  
22 4A). Spatial correlation analysis between all genes and Amyloid  $\beta$  in those regions showed that many  
23 genes in our panel were potentially responsive to Amyloid  $\beta$  (Fig. 4B). The most relative genes included  
24 Gfap, Tyrobp, Ctsd, B2m, and Apoe, which express mainly in reactive astrocytes and microglia cells  
25 (19, 20). The aggregation of those genes was confirmed by raw sequencing images (Fig. 4B), and the  
26 ranking of spatially-correlated genes were independent of the bin size selected for analysis (Fig. S15B).  
27

28 At the cellular level, our SPRINTseq results showed that neurons were depleted from the Amyloid  $\beta$   
29 while the reactive glia cells, including microglia and astrocytes, were aggregated around it (Fig. 4C). To  
30 directly compare the cell compositions of AD and normal brains, we combined both data sets and  
31 classified them together before projection (Fig. 4D, Fig. S15C). The proportion of reactive glia cells in  
32 the AD mouse brain was more than twice that in the normal mouse brain. For other cell types, although  
33 the difference in quantity between AD and normal brains was marginal, the local densities showed  
34 different spatial dependencies to Amyloid  $\beta$  and those changes were not random (Fig. 4E, Fig. S15D).  
35 Clearly, the reactive glia cells were clustered favorably around Amyloid  $\beta$ , within 200  $\mu$ m, which reflected  
36 their roles of responding to microenvironmental changes and plaque clearance. The reduction of  
37 neuronal density near Amyloid  $\beta$  reflected neuron apoptosis in a plaque-rich microenvironment. Notably,  
38 the pathological effect of AD is not only associated with the local microenvironment but can be scaled  
39 to larger regions. For example, oligodendrocytes did not show significant density changes around  
40 Amyloid  $\beta$ , but their presence in the cortex region was generally less in the AD than in the normal mouse  
41 brain (Fig. 4F, Fig. S15E).  
42

1 SPRINTseq extended the analysis' spatial resolution to the subcellular level, enabling us to find many  
2 cells that showed distinct differences that were related to the cellular distance to Amyloid  $\beta$ . In reactive  
3 glia cells, the subcellular dispersion of mRNAs was increased in Amyloid  $\beta$  adjacent cells (distance  
4 threshold: 25  $\mu$ m) than in more distant cells (Fig. 4G & H). This was largely because the cytoplasmic  
5 size of reactive glia cells near Amyloid  $\beta$  increased because of increased and elongated surface bumps.  
6 Disease-associated microglia (DAM) have been reported to play a vital role in AD and the function is  
7 conserved in mice and human (21, 22). Interestingly, the Amyloid  $\beta$  adjacent reactive microglia we found  
8 may largely correspond to reported plaque-phagocytic microglia (XO4+) in gene expression pattern  
9 (Tyrobp+, Apoe+, B2m+ and low Cx3cr1) and behavior (spatially enriched around plaques) (23). The  
10 enlarged cell body might suggest their function in phagocytosis and other regulation process within the  
11 micro-environment, whereas XO4- microglia can't migrate towards plaques and thus are far away from  
12 plaques. The spatial information can be used as a new dimension for confirming plaque-phagocytic  
13 microglia.

14  
15 Additionally, in a cell that is directly adjacent to a plaque, the location of one of its gene's mRNA  
16 orientations with respect to the nearest Amyloid  $\beta$  can characterize the relative proximity of that gene.  
17 So, we calculated the included angle between the mRNA's orientation, the nucleus centroid, and  
18 Amyloid  $\beta$ 's direction. A smaller angle indicated that the mRNA tended to be closer to its nearest plaque  
19 (Fig. S16A). Changes in the distance to the nucleus can also be calculated and combined with the  
20 orientation calculation to better describe the mRNA's tendency to approach or retreat (Fig. S16B). Much  
21 heterogeneity was found within cells especially neurons, suggesting a different cell response pattern  
22 during disease (Fig. S16C). Such orientation was dependent on the distance between a cell and  
23 amyloid (Fig. 4I, Fig. S16D). This orientation feature pattern disappeared at longer distances, as the  
24 effect of Amyloid  $\beta$  on distant cells became weaker. Altogether, Amyloid  $\beta$  might generally affect its  
25 nearby cell morphology and potentially alter sub-compartment architecture in various cell types.

26  
27  
28 **Discussion**  
29  
30 SPRINTseq is an intrinsically high-speed spatial sequencing method that can finish a mouse brain  
31 coronal slice profile at subcellular resolution within 20 hours, including both sample preparation (10  
32 hours, and several slices can be prepared in parallel) and cyclic sequencing (9.5 hours). Due to a  
33 combination of highly effective barcode coding and robust SBS chemistry, SPRINTseq is significantly  
34 faster than most other approaches. Besides, we developed three orthogonal approaches to relieve the  
35 conventional ISS's signal crowdedness issue. Among those approaches, optical dilution holds great  
36 potential and deserves further examination. For example, with constant codeword Hamming weights,  
37 more silent bits can be gained by increasing the barcode length. Multiple padlocks with different

1 barcodes can also be applied to detect more transcripts of highly expressed genes, if necessary, which  
2 resembles selective amplification but is lossless in spatial information.

3  
4 One major advantage of this method is the single-molecule high spatial resolution that gives sub-cellular  
5 localization information. This is important for addressing questions of cellular interactions in the context  
6 of gradients or proximity to specific tissue features such as disease lesions. In mouse models of  
7 Alzheimer's disease, subpopulations of microglia with distinct transcriptomic phenotypes have been  
8 identified via single-cell RNA-sequencing (24). These seemingly AD-associated microglia bearing  
9 stronger inflammatory signatures and were hypothesized to engulf Amyloid  $\beta$  plaques (24). We  
10 described the distribution of different types of cells near Amyloid  $\beta$  in Alzheimer's disease mouse model,  
11 and we identified reactive glia cells with a distinct expression pattern that are responsive to Amyloid  $\beta$ ,  
12 similar to what was previously reported (21, 25). The spatial information, including their distance to  
13 Amyloid  $\beta$ , can further confirm their functional and phenotypic diversity. In addition, subcellular mRNA  
14 distribution was also informative, showing specific transcript distribution patterns within the cell  
15 including polar, random, and centripetal distributions. Extensive changes in these subcellular  
16 distributions in glia cells and neurons are found within the Amyloid  $\beta$  microenvironment, most likely  
17 caused by changes in the cytoplasm morphology and membrane position, consistent with previous  
18 reports (24). Cell functions including phagocytosis, stimulus sensing and response, and cell-to-cell  
19 interactions are closely linked with these structural alterations during disease development (23, 24, 26).

20  
21 Subcellular mRNA distribution has high heterogeneity across cell types and subtypes. All reactive glia  
22 cells were classified as such based on their gene expression, but importantly the mRNA dispersion  
23 degree is higher in cells that are closer to plaques within the same type (the larger cytoplasm size is a  
24 typical phenotypic feature of glia cell activation in addition to specific gene expression). This new facet  
25 of in-situ sequencing data has added one more dimension to the conventional cellular-level gene  
26 expression matrix. We assert that a combination of gene expression and subcellular mRNA distribution  
27 will improve the understanding of complex transcriptional molecular functions in tissues and lead to  
28 more accurate and more quantitative characterizations of cell types and states.

29  
30 At present, SPRINTseq is yet to be improved in some aspects. Firstly, cell segmentation in brain is a  
31 challenge in the field, because cells (including neurons and glia cells) are intricately distributed in the  
32 brain, and there is no proper dye that can perfectly define cell boundaries and also is compatible with  
33 in-situ sequencing chemistry. Currently we did not take long projection structure (such as axon) into  
34 account. One possible improvement is to use expression-guided machine learning approaches for a  
35 better cell segmentation. Besides, the scalability holds a great potential to be further improved by  
36 engineering optimization. For example, current total time is limited by imaging speed (~40 min to cover  
37 a mouse brain coronal slice) which is incomparable to sequencing reaction (~15 min for each cycle) in  
38 each cycle. Larger FoV lens with lower magnification could reduce the sequencing time spent by half

1 without damaging data quality due to the coding strategy. The current encoding scheme has about 140  
2 original codewords when fewer than 5% of all codeword pairs (including the original codeword and all  
3 composite codewords) have Hamming distances less than 3, a preferred condition for error correction.  
4 Information acquisition efficiency decreases when more codewords are used to encode a greater  
5 number of genes. To encode all 20,000 genes in the human genome, a simple solution is to splice two  
6 current 10-bp barcodes into a 20-bp barcode while maintaining the current dilution fold ( $140^2 \sim 20,000$ ).  
7 However, when more genes are read, a large dilution fold is needed to maintain crowdedness-robust  
8 signal quality, which likely requires a longer barcode.

9

10 **Acknowledgments**

11 The authors thank BIOPIC sequencing center for experimental assistant. Funding was provided by  
12 National Natural Science Foundation of China Grant 21927802 (Y.H., J.W., P.F.), 22050002 (Y.H.) and  
13 T2188102 (Y.H.), Beijing Municipal Science and Technology Commission Grant Z211100003321006  
14 (Y.H.), and Beijing Advanced Innovation Center for Genomics.

15

16 **Author contributions**

17 Conceptualization: Y.H. and J.W. Experiment: T.C., W.H., M.J., J.L., J.S., Z.C., Y.P., and G.W. Data  
18 analysis: T.C., W.H., M.J., Z.L., M.T., P.F., X.R., and Y.H. Writing: T.C., W.H., M.J., J.W., and Y.H.

19

20 **Competing interests**

21 All authors declare no competing interests.

22

23 **Supplementary Materials in the followed pages.**

24

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