

Immuno-SABER enables highly multiplexed and amplified protein imaging in tissues

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Spatial mapping of proteins in tissues is hindered by limitations in multiplexing, sensitivity and throughput. Here we report immunostaining with signal amplification by exchange reaction (Immuno-SABER), which achieves highly multiplexed signal amplification via DNA-barcoded antibodies and orthogonal DNA concatemers generated by primer exchange reaction (PER). SABER offers independently programmable signal amplification without *in situ* enzymatic reactions, and intrinsic scalability to rapidly amplify and visualize a large number of targets when combined with fast exchange cycles of fluorescent imager strands. We demonstrate 5- to 180-fold signal amplification in diverse samples (cultured cells, cryosections, formalin-fixed paraffin-embedded sections and whole-mount tissues), as well as simultaneous signal amplification for ten different proteins using standard equipment and workflows. We also combined SABER with expansion microscopy to enable rapid, multiplexed super-resolution tissue imaging. Immuno-SABER presents an effective and accessible platform for multiplexed and amplified imaging of proteins with high sensitivity and throughput.

In situ protein imaging using immunofluorescence maps specific targets within their native environment. Accurate representation of cell type and state requires visualization of multiple markers. However, conventional methods offer limited multiplexing of targets (typically no more than five) owing to spectral overlap of fluorophores. Several recent methods achieve higher multiplexing, but often with decreased sensitivity, throughput or accessibility. Multiplexed ion beam imaging (MIBI)^{1,2}, imaging mass cytometry (IMC)³ and multiplexed vibrational imaging⁴ use specialized instruments to point-scan small fields (for example, ~2–5 min per 50 µm × 50 µm) and hence are exceedingly slow for large centimeter-scale tissue sections. Fluorescence methods utilizing sequential antibody staining (for example, MxIF⁵, CycIF^{6,7} and 4i⁸) offer high accessibility and relatively fast image acquisition, but require multiple slow cycles of primary antibody incubation (typically hours per cycle) and can take weeks to image tens of targets^{5–7}. Alternatively, DNA-barcoding methods simultaneously apply ten or more antibodies, labeled with orthogonal DNA barcodes, followed by fast sequential barcode readout either through rapid binding and unbinding of fluorescent imager strands using DNA-Exchange (DEI)^{9–11} or by *in situ* polymerization of fluorescent dNTP analogs (CODEX)¹², reducing experimental time to a couple of days. However, to overcome the overlap of antibody host species, high multiplexing requires conjugating DNA strands to primary antibodies, which results in lower signal (due to the lack of amplification from secondary antibodies) and decreased sensitivity, especially for low-abundance targets in tissues with high autofluorescence and

scattering and low antigen access. Limited signal further lengthens the image acquisition times and reduces throughput. *In situ* signal amplification is thus needed to improve signal, throughput and sensitivity. The ideal amplification method should be scalable and compatible with rapid multiplexing, individually tunable for each target to accommodate the high dynamic range of the proteome¹³, and applicable for spatially overlapping dense targets. However, existing methods do not satisfy these requirements.

Tyramide signal amplification (TSA)¹⁴ amplifies the signal through covalent binding of diffusive tyramide molecules in the vicinity of the target. Owing to the lack of orthogonal chemistries, TSA amplifies one target per round, and necessitates slow rounds of (typically microwave-based) antibody removal for spectral multiplexing^{15,16}. In contrast, by using orthogonal sequences, DNA-based methods allow simultaneous signal amplification. In rolling circle amplification (RCA)¹⁷, a processive polymerase acts on a circular template to synthesize long concatenated repeats. RCA offers high levels of amplification and potential for multiplexing, but the *in situ* enzymatic reaction is hard to control or tune for individual targets¹⁸. TSA and RCA may also lead to blurring of signals and decreased resolution, respectively, owing to spreading of the tyramide molecules or the large size of the amplicons (reaching from 250 nm to over 1 µm in radius^{19,20}). Branched DNA assemblies^{21,22}, such as RNAscope²³, generate complex tree structures for stable binding of fluorescent DNA strands, whereas the hybridization chain reaction (HCR) utilizes triggered assembly of metastable fluorophore-conjugated hairpins^{24–26}. The structural complexity of existing

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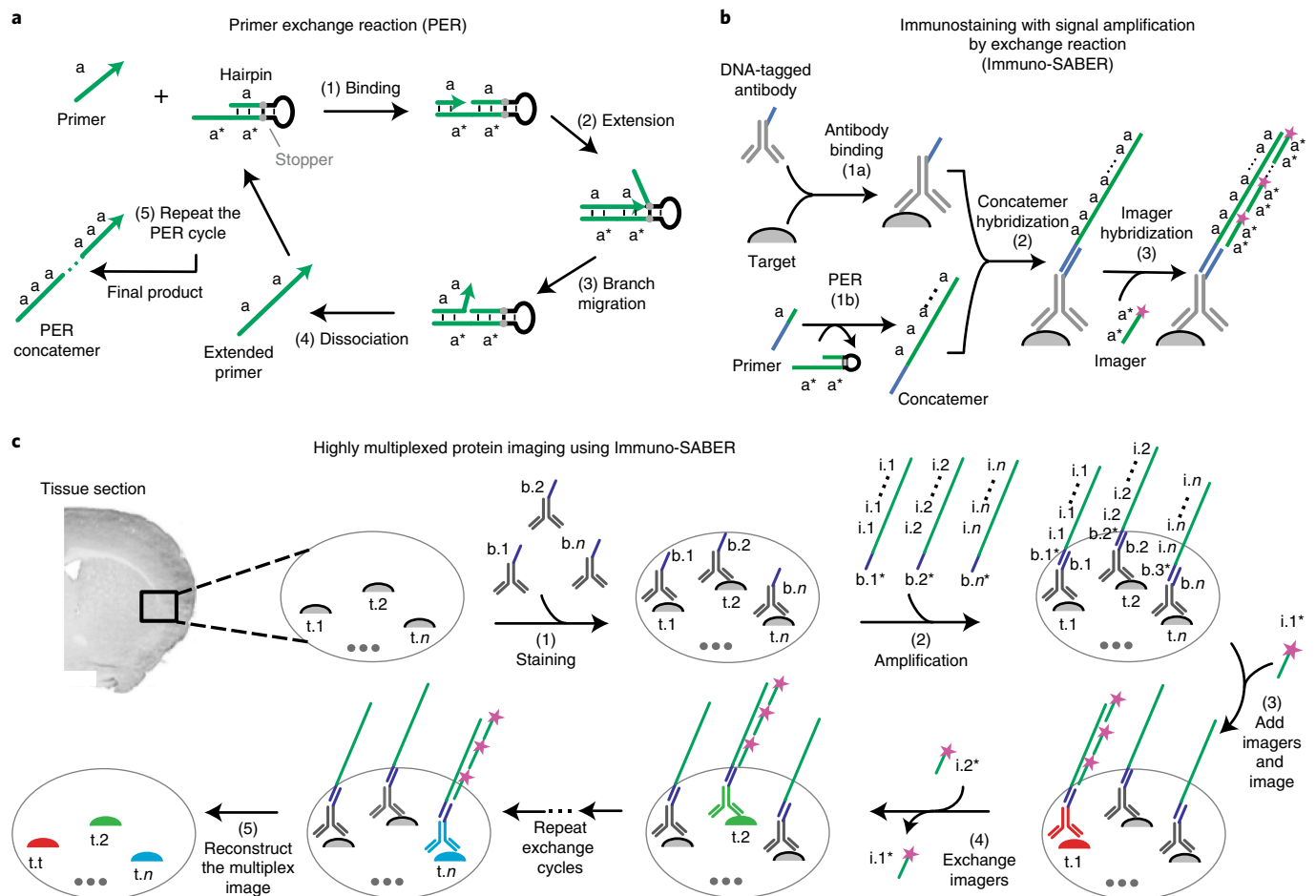


Fig. 1 | Immuno-SABER schematic. **a**, PER mechanism²⁹. (1) A 9-nucleotide primer sequence *a* binds to the single-stranded *a** sequence on a hairpin (the asterisk indicates reverse complementarity). (2) The primer is extended by a strand-displacing polymerase (for example, Bst) isothermally and autonomously. The hairpin features a stopper sequence that halts polymerization, which releases the polymerase. (3) The newly synthesized *a* sequence is displaced from the hairpin through branch migration. (4) The extended primer and the hairpin autonomously dissociate. (5) Repetition of this copy-and-release process produces a long concatemer of *a*. **b**, Immuno-SABER schematic. (1a) Antibodies conjugated with bridge strands are used to simultaneously stain multiple targets. (1b) Primer sequences (green) are independently extended to a controlled length using PER. (2) Concatemers hybridize to the bridge sequence (blue) on the antibody. (3) Fluorophore (purple star)-labeled 20-nucleotide DNA imager strands hybridize to the repeated binding sites on the concatemers. Each imager is designed to bind to a dimer of the unit primer sequence. **c**, Exchange-SABER schematic. (1) Different biological targets (*t*.1 to *t*.*n*) are labeled with antibodies conjugated to orthogonal bridge strands (*b*.1 to *b*.*n*). (2) Orthogonal pre-extended concatemers are hybridized (via bridge complements *b*.1* to *b*.*n**) to the bridge strands on the antibodies simultaneously. (3) Target *t*.1 is visualized by hybridization of imager *i*.1* to the *i*.1 sites on the concatemer bound to *b*.1 on the corresponding antibody. (4) Multiple targets can be imaged sequentially by hybridization and dehybridization of orthogonal imagers in multiple rapid exchange cycles. (5) The images are computationally aligned and pseudocolored to overlay different targets.

DNA-assembly-based platforms could present potential challenges for designing highly multiplexed orthogonal systems. In practice, simultaneous signal amplification for proteins beyond spectral multiplexing (3–5 targets) has not been demonstrated^{27,28}.

Here we report Immuno-SABER, a highly multiplexed and individually controllable signal amplification method free of *in situ* enzymatic reactions (Fig. 1). After staining a sample with multiple DNA-barcoded primary antibodies, SABER involves hybridization of these barcodes to orthogonal single-stranded DNA concatemers, which are generated in a preprogrammed manner via primer exchange reactions (PERs)²⁹ (Fig. 1a) and bind multiple fluorophore-bearing imager strands for multiplexed signal amplification³⁰ (Fig. 1b). Rapid spectrally unlimited multiplexing can be achieved via Exchange-SABER, which combines SABER amplification with rapid exchange cycles of fluorescent imager strands^{9–11} (Fig. 1c). We validated Immuno-SABER in diverse samples, including cultured cells, cryosections, formalin-fixed paraffin-embedded (FFPE)

sections and whole-mount tissues, and demonstrated 5- to 180-fold signal amplification that was independently tunable, covering the range conventionally achieved by secondary antibodies to TSA. The simple design of SABER amplifiers makes it scalable. We further demonstrate simultaneous signal amplification and imaging of ten protein targets in tissues. Finally, we combine SABER with expansion microscopy (ExM)³¹ to achieve rapid highly multiplexed super-resolution tissue imaging.

Results

Validation of *in situ* signal amplification by Immuno-SABER.

For *in situ* signal amplification, Immuno-SABER relies on controlled *in vitro* synthesis of amplifier concatemers by PER, followed by programmed *in situ* assembly. PER utilizes a catalytic hairpin for controlled extension of a short primer sequence in an iterative manner (as in Fig. 1a). Primer designs rely on a three-letter code (made of only A, T and C nucleotides), where G nucleotides are avoided in

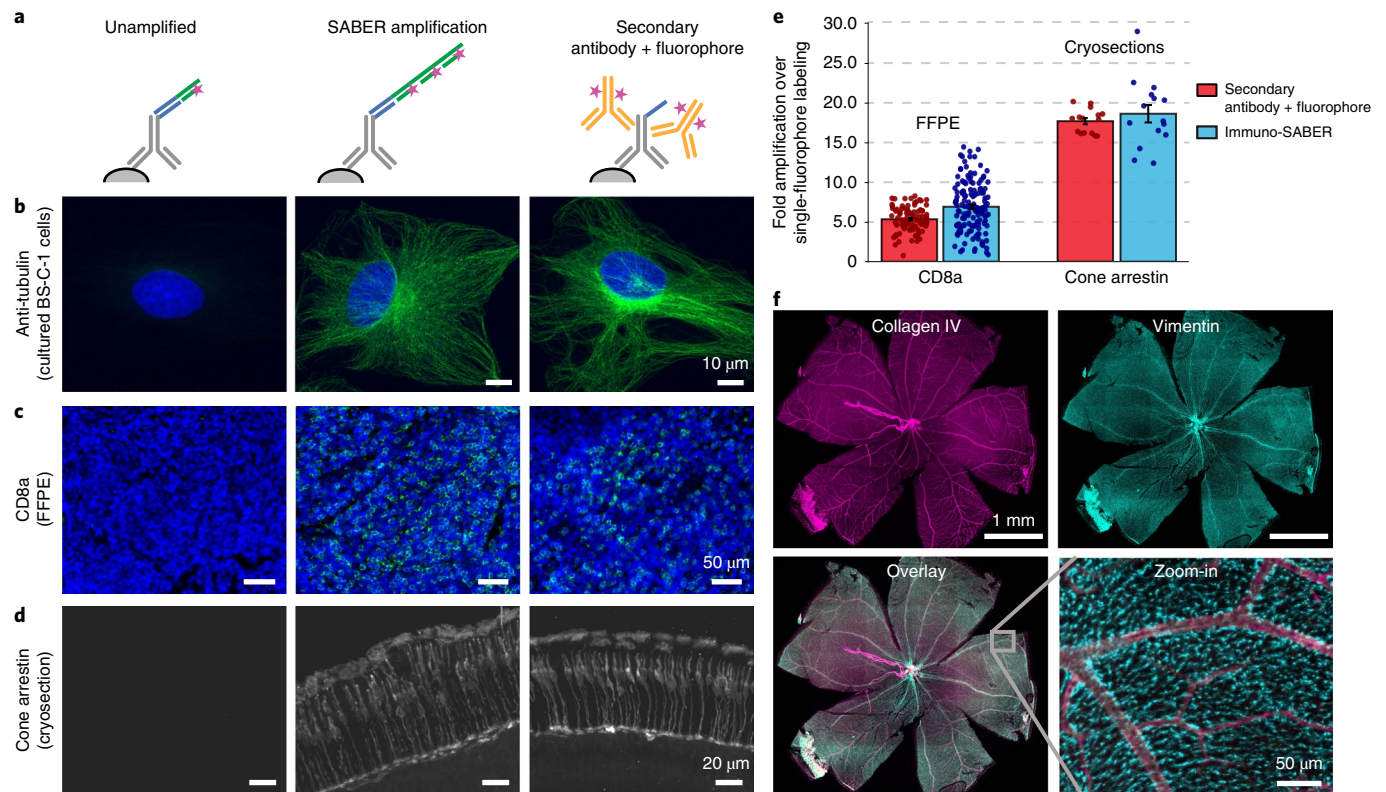


Fig. 2 | Validation and quantification of signal amplification by Immuno-SABER. **a**, Cultured BS-C-1 cells were immunostained for α -tubulin and the following three conditions were prepared for comparison: an unamplified condition, in which unextended primers with a single binding site for the imager (with Alexa Fluor 647) were hybridized to the bridge on the antibody; an amplified condition, in which the extended concatemer was hybridized for signal amplification (linear amplification); and a control condition, in which conventional antibody staining was performed with Alexa Fluor 647-conjugated secondary antibodies. **b**, Representative images of each condition from **a** (maximum projections from the confocal z stack taken with a 63 \times objective). **c**, Representative images for CD8a staining (labeled with an Atto488 imager) in human tonsil FFPE sections (single-plane large-area scans with a 20 \times objective cropped to show a region of the interfollicular zone that is rich in CD8 $^{+}$ cells). **d**, Cone arrestin staining in mouse retina cryosections (maximum projection of a confocal z stack taken with a 20 \times objective). See the Methods for experimental details. **e**, The level of signal amplification by Immuno-SABER was quantified by measuring the background-subtracted mean fluorescence for several ROIs in the tissues and is expressed as the fold signal amplification over the unamplified signal level. Conventional secondary antibody amplification was also quantified similarly and is shown as reference. For CD8a FFPE sections, $n=80$ (for unamplified), $n=100$ (for SABER) and $n=94$ (for conventional) rectangular ROIs (each covering 0.03–1.20 mm 2 tissue regions; consecutive tissue sections are used for the three conditions). For cone arrestin, $n=6$ images were taken from two retina samples. Error bars, s.e.m. **f**, Immuno-SABER was performed in whole-mount retina sections for collagen IV and vimentin. Maximum projections from confocal z stacks are displayed.

the primer sequence and in the reaction mixture^{29,30}. Hence, the C nucleotide after the template on the hairpin acts as a stopper for the polymerase. This simple step-by-step synthesis offers tight control of the reaction by external parameters (such as hairpin or dNTP concentration, reaction time and temperature) as well as high programmability and yields long DNA concatemers of desired lengths reaching >500 nucleotides³⁰. For a modular design, we conjugated antibodies with 42-nucleotide DNA sequences ('bridges'; from the orthogonal library we designed previously³⁰), which enable hybridization of concatemers through their 5' sections of sequence that are complementary to the bridges. To preserve antigen recognition, we optimized our protocol to conjugate 1–3 oligonucleotides per antibody (see Supplementary Note 1). We also developed an optional purification strategy based on toehold displacement-mediated DNA affinity pull-down (Supplementary Fig. 1).

For application of our in vitro extension and in situ assembly strategy on biological specimens, labeling specificity, resolution and amplification efficiency are the main considerations. First, to evaluate the specificity and preservation of morphology, we performed Immuno-SABER staining of microtubules in cultured cells as a test

case for a densely arranged structural protein target. We observed specific staining, clear tubular morphology and a similar staining pattern to conventional immunostaining with fluorophore-conjugated secondary antibodies (Fig. 2a,b). The resolution of the images, as quantified by the full-width-half-maximum (FWHM) of the fitted line plots across the thin microtubules, was unaltered as compared to the secondary antibody control (Supplementary Fig. 2a–c). These results demonstrated the suitability of Immuno-SABER for in situ labeling of proteins, with high resolution and high specificity even in dense arrangements.

For validation of the labeling strategy in tissue samples and quantification of the signal amplification, we used the following two types of tissue preparations: 5- μ m-thick FFPE human tonsil sections and 40- μ m-thick mouse retina cryosections. Both tissue types provide a good validation platform owing to the morphologically distinct and well-conserved organization of different cell types that can be identified by well-established biomarkers. The retina has a layered organization of different types of cells whereas tonsils feature multiple germinal centers with stereotypic organization of a large number of distinct cell types.

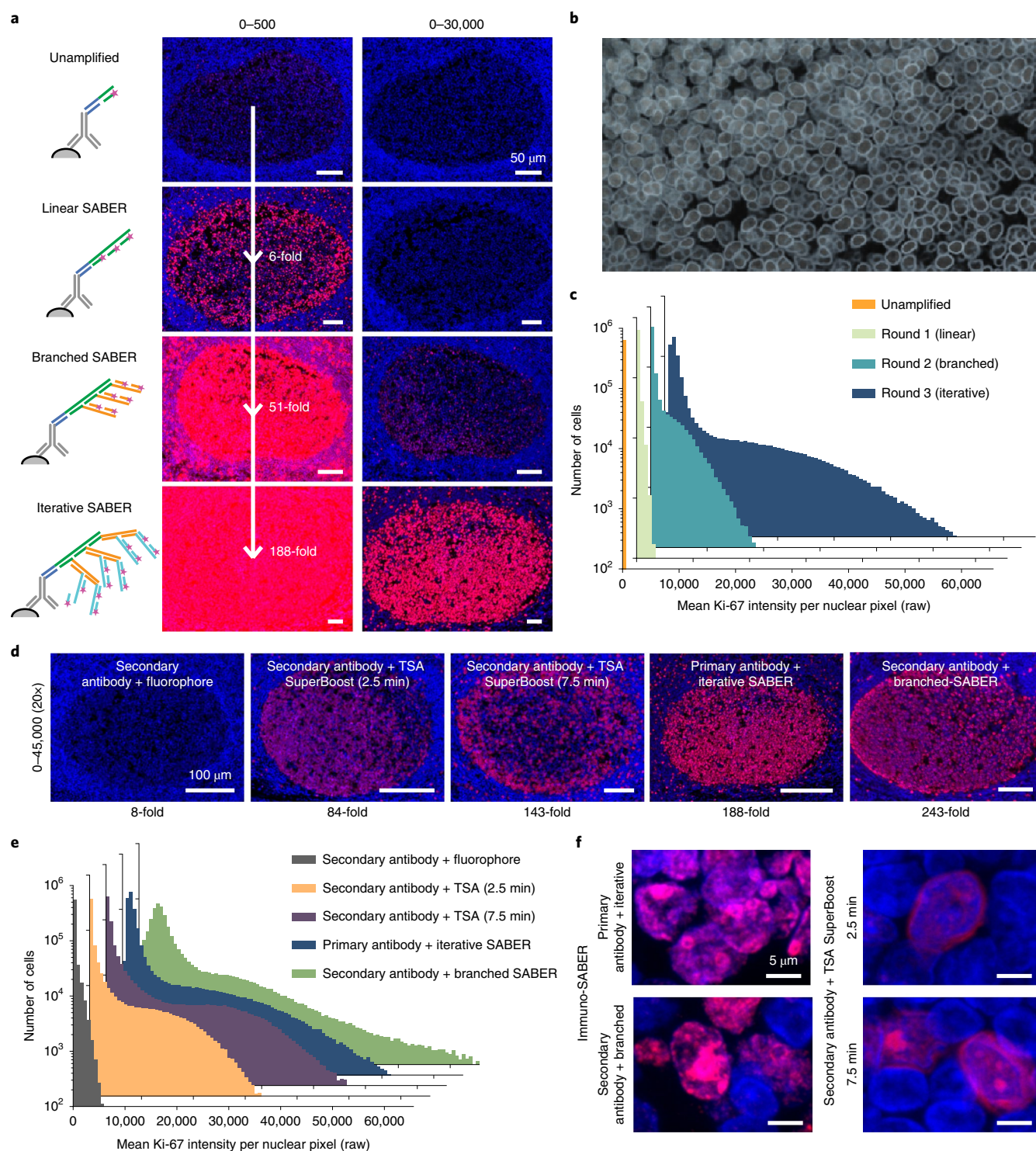


Fig. 3 | Immuno-SABER signal can be further amplified through branching. a, Nuclear Ki-67 (Alexa Fluor 647, red) imaging with DAPI (blue) in the Ki-67-rich germinal center of FFPE human tonsil sections shown after up to three rounds of amplification (iterative SABER). Sixteen-bit images were scaled to two different maximum pixel values (500 and 30,000) to allow visual comparison. Signals were quantified in each case versus the unamplified sample and the fold changes are provided. **b**, Machine-learning-based contouring of the nuclei for quantification of signal per cell. See Supplementary Fig. 5a and the Methods for more information. **c**, Mean Ki-67 signal intensity for each nucleus was obtained from automated segmentation and the histogram was plotted for the whole tissue section for each condition. The consecutive sections each contain 636,479–717,176 identified nuclei. **d**, Images show germinal centers in FFPE human tonsil sections with Ki-67 labeling (red) with conventional secondary antibody–fluorophore staining, TSA (with HRP-conjugated secondary antibodies) for 2.5 and 7.5 min, primary antibodies using iterative SABER amplification or secondary antibodies using branched SABER amplification. TSA was applied using poly(HRP)conjugated secondary antibodies from a commercial SuperBoost Kit with 2.5 or 7.5 min of incubation with tyramide–Alexa Fluor 647. The amplification levels are noted below the images. **e**, Histograms to visualize mean nuclear signal level were plotted for the conditions in **d**. The consecutive tissue sections each contain 586,183–717,176 cells. **f**, Samples were imaged with a confocal microscope at a higher resolution with 63 \times magnification to evaluate signal blurring. Images with different scaling are displayed in Supplementary Fig. 5f.

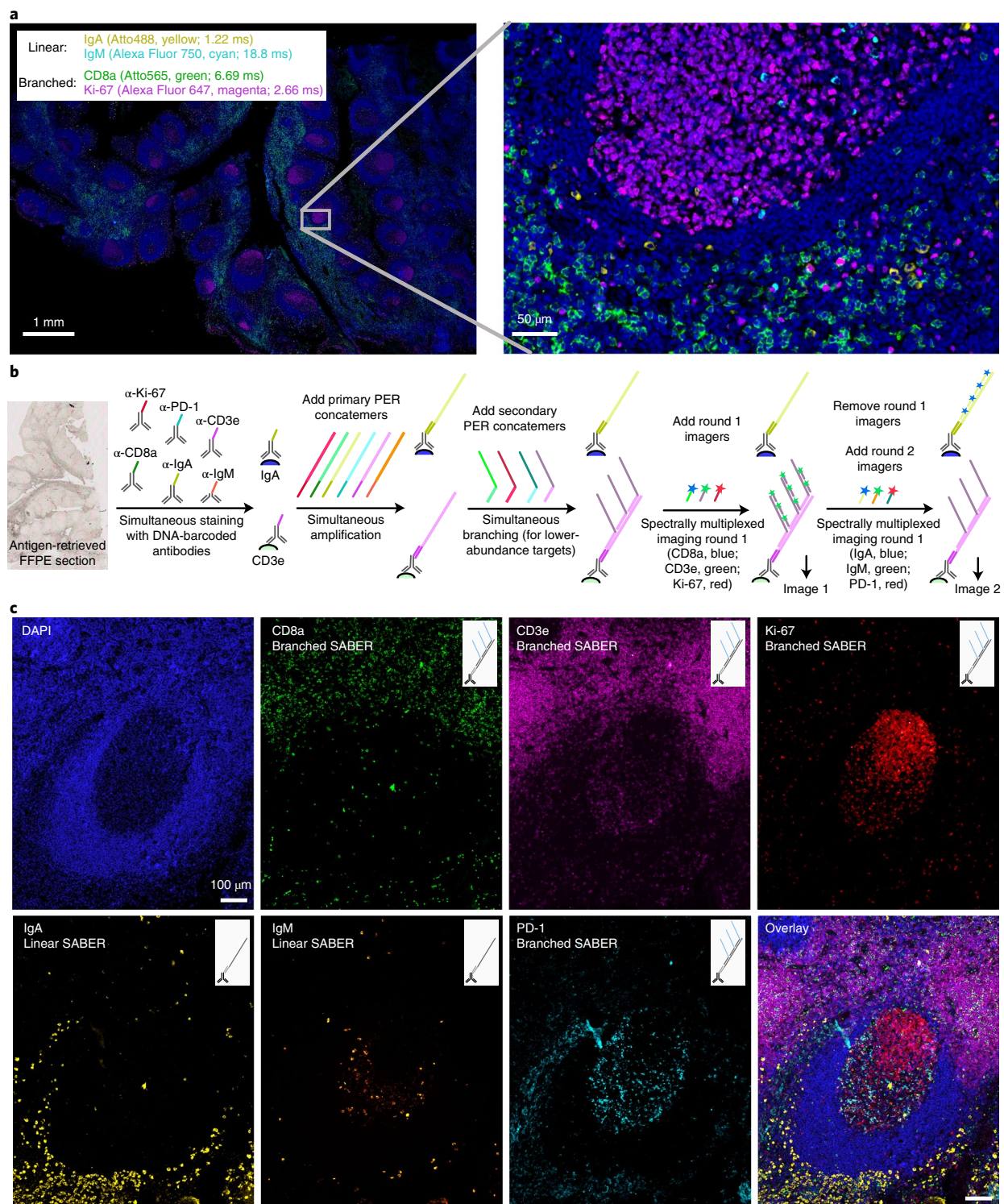


Fig. 4 | Immuno-SABER and Exchange-SABER in FFPE human tonsil samples. a, Left, centimeter-scale whole-slide imaging of human tonsil sections with five-color spectral multiplexing (DAPI and four other targets). Right, an expanded view of the region marked with the gray box showing four-target imaging with subcellular resolution with a $\times 20$ objective. Exposure times for each target (autoexposure setting) are indicated in a white box. For IgA and IgM (higher copy number), linear amplification yields high enough signal to achieve autoexposure times of 1–20 ms under optimized conditions. For Ki-67 and CD8a (lower copy number)^{47,48}, branched amplification (one round of branching) was applied to allow auto-exposure times of 2–10 ms. **b**, Higher multiplexing via Exchange-SABER. The schematic for the multiplexed imaging workflow where all antibodies were applied simultaneously, followed by simultaneous amplification and sequential rounds of imaging. IgA and CD3e labeling structures are shown as examples to illustrate the workflow for linear and branched SABER on the same sample. **c**, Images show an expanded view of a germinal center in human FFPE tonsil sections imaged in seven colors (DAPI and six other targets) with a single exchange round (round 1, top row; round 2, bottom row). Four of the targets, CD8a (Atto488), CD3e (Alexa Fluor 647), Ki-67 (Alexa Fluor 750) and PD-1 (Alexa Fluor 750), were visualized with simultaneous branched SABER amplification, whereas IgA (Atto488) and IgM (Atto565) were visualized with linear amplification.

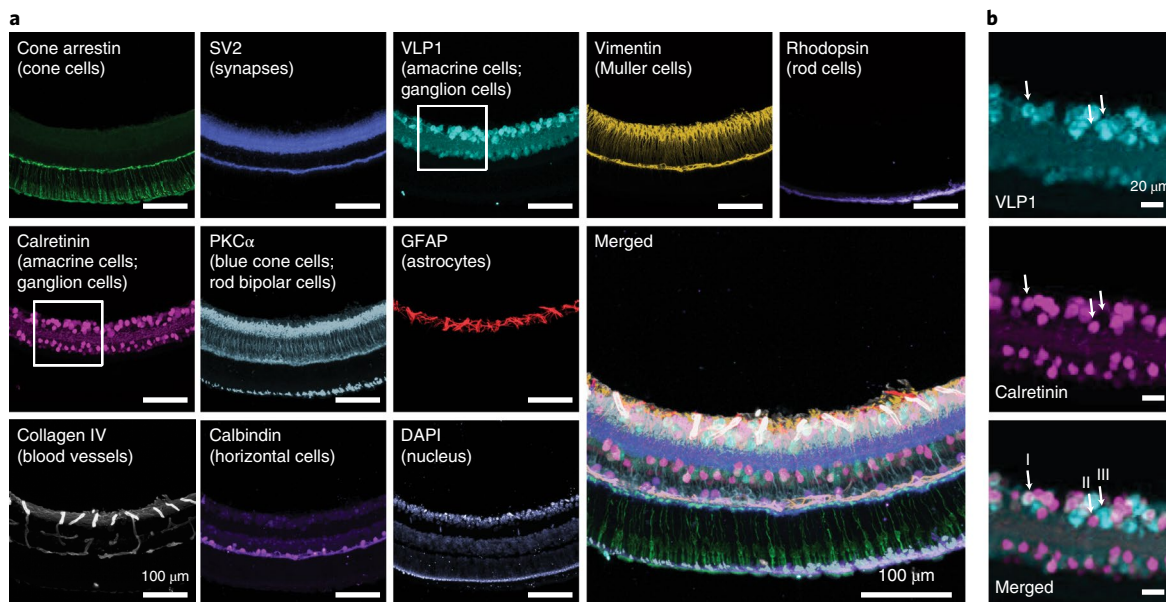


Fig. 5 | Exchange-SABER in mouse retina cryosections. a, Ten protein targets labeling various retinal cell types were visualized in 40- μ m-thick mouse retina cryosections. The markers targeted with Immuno-SABER were rhodopsin (rod photoreceptors), GFAP (astrocytes), vimentin (Muller cells⁹), collagen IV (blood vessels), three calcium-binding proteins^{49,50} (VLP1, calretinin (found in a subset of amacrine and ganglion cells) and calbindin (note that, although calbindin was suggested to be also found in a subset of amacrine and ganglion cells, the anti-calbindin antibody used here mostly labels horizontal cells⁵⁰)) and PKC α (blue cone cells and rod bipolar cells⁵⁰). The sections were first incubated with all of the DNA-conjugated antibodies simultaneously. All SABER concatemers were then added simultaneously to the sample, followed by washing and sequential incorporation of the imager strands and multiround imaging. A z stack of images was acquired for each target, and DAPI was imaged in every exchange cycle to monitor sample drift. The maximum projected images of each stack were computationally aligned using a subpixel registration algorithm with DAPI as the drift marker⁹, and pseudocolored for the overlay presentation. **b**, Expanded view of the area marked by the white rectangle in **a**. Three cell subtypes (marked with arrows: I, VLP1⁺ and calretinin⁺; II, VLP1⁻ and calretinin⁺; III, VLP1⁺ and calretinin⁻) can be differentiated on the basis of VLP1 and calretinin expression.

For the FFPE samples, we imaged the T cell membrane marker CD8a. CD8a⁺ cells are largely present in the marginal zone (outside of germinal centers)³². For mouse retina cryosections, we imaged cone arrestin, a specific marker of cone photoreceptor cells⁹. Both markers showed signal patterns consistent with the expected distribution of the markers, suggesting high specificity (Fig. 2c,d). For both targets and sample types, Immuno-SABER yielded similar or slightly higher fluorescence signal than conventional fluorophore-conjugated secondary antibody staining using the same fluorophore. We quantified the signal amplification level as 6.7-fold for CD8a and 19.4-fold for cone arrestin. For comparison, conventional secondary antibody staining yielded 5.3- and 17.5-fold signal amplification, respectively (Fig. 2e). Overall, our strategy generated an improvement of approximately 5- to 20-fold in the signal level as compared to the unamplified control, reaching amplification in the range of secondary antibodies. The degree of in situ signal amplification may depend on multiple factors, including the abundance and organization of targets, the antibodies (for example, clonality and conjugation efficiency) and the method of quantification (the unamplified signal level, thresholding and background subtraction), as well as the experimental conditions and properties of the SABER sequences (for example, concatemer length³⁰).

Despite reaching >500 nucleotides in length, SABER concatemers can effectively penetrate relatively thick samples. We validated the concatemer penetration capability in whole-mount preparations of mouse retina by successfully staining for the Muller cell marker vimentin and the blood vessel marker collagen IV, which were both predominantly detected in the 100- μ m region between the nerve fiber layer and the outer plexiform layer of the retina, as expected³³ (Fig. 2f and Supplementary Fig. 2d). This high penetration may potentially be attributed to SABER concatemers being largely linear

DNA structures that are designed using a three-letter code to minimize secondary structures.

Enhancement of signal through branching. To enable further signal enhancement, which would increase the sensitivity for the proteins of lower abundance or improve imaging throughput, we developed a sequential amplification strategy where independently extended secondary concatemers can be branched off the primary concatemer to create more binding sites for fluorescent imagers (branched SABER; Supplementary Fig. 3a). We performed similar tests to check the effect of an additional amplification round targeting the same proteins in cell culture, human FFPE tonsil tissues and mouse retina cryosections (Supplementary Fig. 3b,c). With one round of branching, we obtained an additional 2.8-fold signal amplification for CD8a in FFPE sections and a 8.4-fold signal amplification for cone arrestin in cryosections as compared to a single round (Supplementary Fig. 3d). Microtubule staining in cells indicates that the specificity of labeling and the high-resolution morphology of the target were still preserved with branched SABER (Supplementary Fig. 3e). FWHM estimations demonstrated that the sizes of the structures were still bound by the diffraction-limit and were not significantly altered (Supplementary Fig. 3f).

To evaluate the preservation of access to the target and variation in overall signal distribution, we performed baseline quantification and linear and branched amplification for a ubiquitous protein (lamin B) in the same sample of HeLa cells using DNA-conjugated secondary antibodies and varying lengths of concatemers and branches (Supplementary Fig. 4). We did not detect a significant difference in the baseline signal for concatemers up to 700 nucleotides in length (for linear amplification), suggesting that the length range we utilized in our experiments did not cause decreased

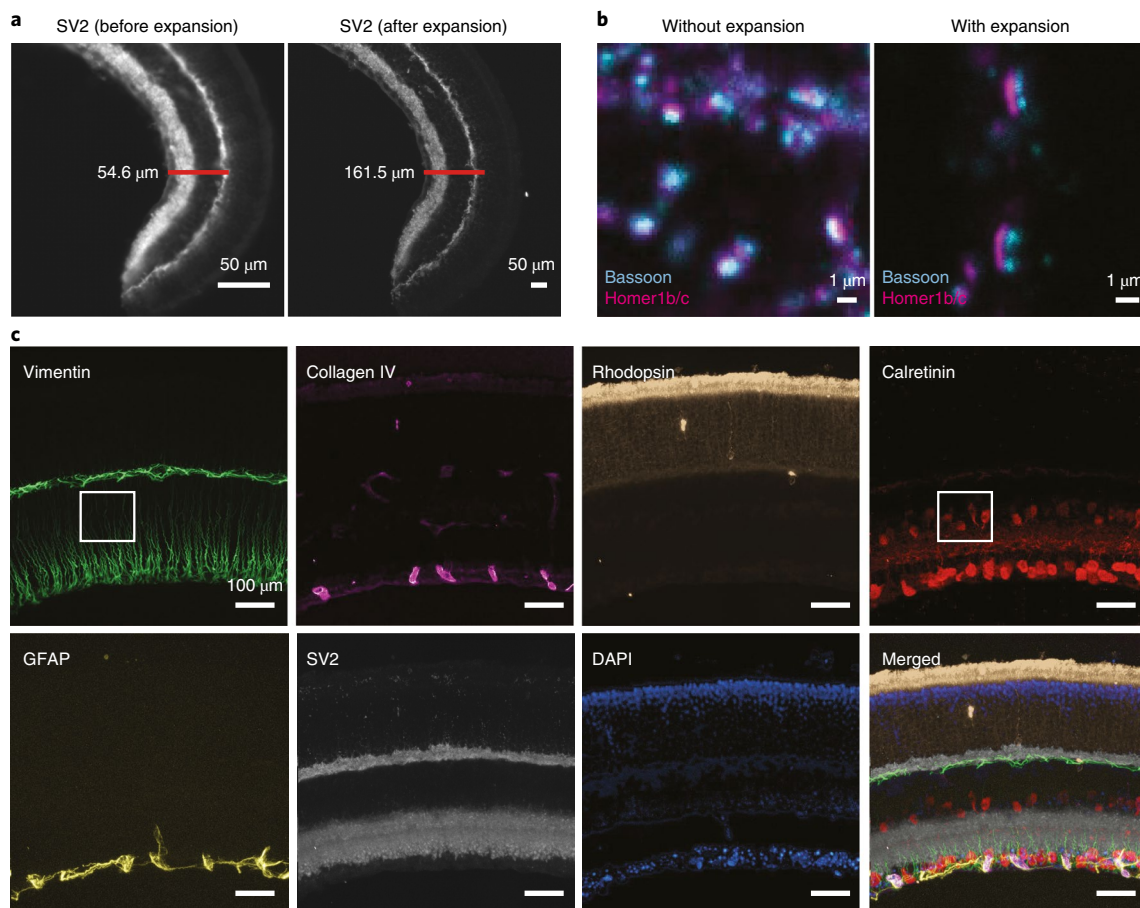


Fig. 6 | Multiplexed super-resolution imaging using Expansion-SABER. **a**, Forty-micrometer-thick mouse retina cryosections were stained for SV2 using DNA-conjugated anti-SV2 antibodies, followed by SABER concatemer hybridization. Images were acquired before hydrogel formation and after hydrogel formation and expansion (approximately threefold) using the original expansion protocol^{31,39}. **b**, Images of pre- and postsynaptic sites of neuronal synapses in fixed primary mouse hippocampal neuron culture samples with and without expansion (different fields of view are shown). The presynaptic sites were labeled with anti-Bassoon antibodies and the post-synaptic sites were labeled with anti-Homer1b/c antibodies. DNA-conjugated secondary antibodies were used to target anti-Bassoon and anti-Homer1b/c primary antibodies, and SABER concatemers were then applied. **c**, ExM imaging of six protein targets in the originally 40-μm-thick mouse retina section (expanded approximately threefold) with Exchange-SABER. Two exchange rounds with Atto488-, Atto565- and Alexa Fluor 647-conjugated imager strands were performed to visualize all six targets in the expanded samples. DAPI was imaged in both rounds to serve as a registration marker. The images are maximum projections of z stacks, drift corrected using DAPI channels and pseudocolored for presentation. An expanded view of the boxed region is available in Supplementary Fig. 9b.

access (Supplementary Fig. 4b,g). For reference, we also compared the baseline signal to the signal from secondary antibodies, which, on average, were conjugated to five fluorophores. We were able to see that in this case signal level was approximately fivefold higher than the single-fluorophore baseline, further demonstrating comparable antigen access (Supplementary Fig. 4c). For the increasing concatemer lengths, the expected amplification trend was obtained across samples with quantitatively consistent intensity values. Blue bars in Supplementary Fig. 4b,g can be compared for more insights; for example, for linear amplification, approximately two-fold higher signal intensities were obtained with the long primary concatemers (700 nucleotides) versus the short primary concatemers (350 nucleotides).

Immuno-SABER allows further improvement of sensitivity by performing multiple branching rounds (iterative SABER). We applied this strategy both in tonsil FFPE sections and mouse retina cryosections, and performed three sequential amplification rounds following primary antibody staining. We targeted the proliferating cell nuclear marker Ki-67 that is mainly found in the germinal center (that is, the dark zone) in tonsil sections and the synaptic marker

SV2 that is found mainly in synapses of both the outer and inner plexiform layers⁹. Figure 3a shows Ki-67-rich germinal centers stained at different amplification levels, displayed at two different contrast scales to allow visual comparison of the signal.

To follow the change in the signal level for individual nuclei in the whole-section scans, we developed a machine learning algorithm that automatically annotates DAPI-defined nuclear contours (Fig. 3b and Supplementary Fig. 5a) for nuclear segmentation. This allowed us to assign mean Ki-67 signal intensity to individual nuclei in sections of 550,000–750,000 cells each, and plot histograms for each amplification condition (Fig. 3c). The total nuclear signal for each condition gave a basic quantitative estimate of the amplification level with respect to the unamplified case, yielding 6-, 51- and 188-fold signal amplification, respectively for linear, branched and iterative SABER (Fig. 3a). Although the amplification process is exponential in nature, throughout the three rounds the coefficient of variation (s.d. over the mean) for the population remained at a similar level (round 1 = 1.76; round 2 = 1.65; round 3 = 1.55), suggesting that increased amplification level does not create a substantial variation in the distribution.

For these samples, concatemer hybridizations can also be performed for shorter durations (for example, with incubation times down to 75 min per round (instead of overnight for the longer primary concatemer, and 3 h for branching as in Fig. 3a) without significant difference in signal level (Supplementary Fig. 5b,c). Iterative SABER (three rounds) was also applied for SV2 in thicker mouse retina cryosections, generating ~80-fold signal amplification (Supplementary Fig. 5d).

For such high levels of amplification, catalyzed reporter-deposition-based TSA is considered a gold standard, with reports of 10- to 1,000-fold signal amplification^{14,34,35}. This can be further improved 2- to 10-fold via poly-horseradish peroxidase (HRP)-conjugated secondary antibodies. However, TSA is not suitable for simultaneous multiplexed amplification owing to the lack of orthogonal chemistries and difficulty in controlling the in situ reaction, and is also not ideal for high-resolution imaging owing to spreading of the fluorescent tyramide molecules, which causes blurring²⁰. To investigate how Immuno-SABER performs in comparison, we first utilized a commercial kit for conventional TSA with mono-HRP-conjugated secondary antibodies and Alexa Fluor 647-tyramide. In retina cryosections, we observed that iterative SABER (with conjugated primary antibodies) could amplify the signal to a higher level than TSA with secondary antibodies (Supplementary Fig. 5e). We also used the commercial SuperBoost kit that features poly-HRP-conjugated secondary antibodies for Ki-67 staining of FFPE tonsil sections, which yielded up to 143-fold signal amplification (Fig. 3d,e). Comparatively, both iterative SABER (188-fold; Fig. 3a) with primary antibodies and branched SABER (two rounds) using secondary antibodies (243-fold) yielded higher signal than TSA (Fig. 3d,e). Where species overlap is not of concern (single or low-plex imaging), linear or branched SABER with secondary antibodies can be performed for increased signal with better-preserved morphology.

At this high amplification level, Immuno-SABER still provided crisp staining and preserved the subcellular morphology, whereas TSA staining resulted in blurring of the signal and spilling over of the label from the target compartments (Ki-67 is a nuclear protein that is highly enriched in nucleoli), as signal coming from outside of DAPI-labeled nuclei could be observed in the high-magnification confocal images (Fig. 3f and Supplementary Fig. 5f,g). Increasing the tyramide incubation period to 10 min (2–10 min being the manufacturer-recommended optimization range) worsened the blurring (Supplementary Fig. 5f,g).

Simultaneous multiplexed amplification by Immuno-SABER.

Outside of a preference for single-stranded concatemers that is facilitated by the use of a three-letter code, PER has few inherent restrictions in sequence design, making design of orthogonal sequences straightforward. We previously developed a computational pipeline to design orthogonal PER primer-hairpin pairs with maximum extension efficiency and minimum crosstalk by utilizing in silico simulations using NUPACK³⁶ and designed 50 orthogonal sequences to enable multiplexed imaging^{29,30}. Here we tested 32 primer sequences and extended each into long DNA concatemers in a controlled fashion up to the target length range of 600–700 nucleotides (Supplementary Fig. 6a). As extension efficiency is sequence dependent, reaction conditions were optimized for each primer by modulating the hairpin concentration and reaction time to obtain concatemers of similar length (Supplementary Table 1). All of the tested primers extended into long concatemers in the desired length range, with 31 of 32 sequences (except sequence 51) yielding a predominant long concatemer band, albeit with some heterogeneity in the distribution of shorter reaction products (Supplementary Fig. 6a). Next, we evaluated the orthogonality of detection (crosstalk check) in situ through an imaging-based multiwell plate assay. We targeted α -tubulin with DNA-conjugated antibodies in cells, and

applied each concatemer in separate wells with its cognate imagers or with all other imagers. All 32 sequences yielded specific staining, and we observed minimal crosstalk (crosstalk signal/cognate signal = 4%) for only one non-cognate imager-primer pair (Supplementary Fig. 6b–e), suggesting that all other sequence combinations are suitable for multiplexed detection.

Next, we sought to validate the applicability of Immuno-SABER for independently programmed multiplexed signal amplification in tissues. We first tested spectral multiplexing on 5- μ m-thick human FFPE tonsil sections. FFPE samples are the standard preparations for clinical settings and for archival purposes. However, these preparations suffer from autofluorescence and low antigen access, making amplification of the target signal a necessity. Clinical immunohistochemistry procedures typically utilize amplification methods such as chromogenic reactions to improve sensitivity, limiting the multiplexing capability to only one target per section. Multiple thin sections have to be prepared to stain different markers, which is not ideal as biopsy-based sample collection is invasive and samples can be scarce. Imaging throughput is another big concern, as typically centimeter-scale tissues need to be imaged by whole-slide scanning.

After standard preparation protocols for deparaffinization and antigen retrieval, we simultaneously stained the samples for Ki-67, CD8a, IgA and IgM using antibodies conjugated to orthogonal bridge sequences. Among these targets, IgA and IgM are expressed at high levels, whereas Ki-67 and CD8a are more moderately expressed (as a proxy, in the tonsil tissue, the RNA expression level of IgJ, the joining chain domain for multimeric IgA and IgM, was found to be 1,360 tags per million (TPM), while Ki-67 and CD8a expression was 15 and 19 TPM, respectively³⁷). We performed whole-slide scanning with five-color spectral multiplexing (DAPI and four other targets) (Fig. 4a). Immuno-SABER amplification also enables improved imaging throughput by allowing short exposure times. For the abundant proteins IgA and IgM, linear amplification provided bright enough signal to permit 1–20 ms exposure times at 20 \times magnification. For moderately available proteins (Ki-67 and CD8a), branched SABER enabled 2–10 ms camera exposures and fast high-quality scans at subcellular resolution (Fig. 4a).

Next, we utilized Exchange-SABER to increase the multiplexing level and image different cell types in germinal centers with six markers (CD8a, CD3e, Ki-67, PD-1, IgA and IgM) (Fig. 4b). SABER amplification was applied simultaneously for all targets via hybridization of concatemers to the orthogonal bridge strands on the antibodies. Simultaneous branching was applied to the four less abundant targets (CD3e, CD8a, Ki-67 and PD-1) following primary concatemer hybridization, and three markers were imaged per round in two rounds as depicted in Fig. 4b. Figure 4c shows a germinal center with interfollicular zone. With our exchange protocol, efficient imager removal can be performed in 10 min without displacing the concatemers (Supplementary Fig. 7).

Exchange-SABER imaging in mouse retina cryosections. Next, we validated Immuno-SABER in 40- μ m-thick cryosections and demonstrated ten-target protein imaging in mouse retina (Fig. 5). We first screened antibodies against a list of targets that have defined staining patterns, including cone arrestin, SV2, visinin-like protein 1 (VLP1), rhodopsin, calretinin, protein kinase C alpha (PKC α), glial fibrillary acidic protein (GFAP), vimentin, collagen IV and calbindin. We then conjugated DNA bridge strands to those antibodies, and validated the specificity and affinity of DNA-conjugated antibodies by comparing the staining patterns from conventional staining using unmodified primary antibodies followed by indirect immunofluorescence (Supplementary Fig. 8a). As controls for exchange imaging, we compared the pre- and post-washing images for SV2, and found the imager strands were efficiently removed after washing (Supplementary Fig. 8b) without sample damage or

signal loss (correlation coefficient between images >0.95 in three wash rounds (Supplementary Fig. 8c)).

The multiplexed imaging result showed that all targets were successfully captured with expected staining patterns (Fig. 5a). Interestingly, we found that two calcium-binding proteins, VLP1 and calretinin, together identify three populations of cells, VLP1⁺calretinin⁻, VLP1⁺calretinin⁺ and VLP1⁺calretinin⁺ cells (Fig. 5b). This observation was confirmed by validating the antibody specificity through comparing the staining patterns of three different antibodies against the same targets and by conventional indirect immunofluorescence using unconjugated primary antibodies (Supplementary Fig. 8d–f).

Fast multiplexed super-resolution imaging by Expansion-SABER. The spatial resolution of conventional fluorescence microscopy is limited by the diffraction of light. A variety of super-resolution techniques have been developed to overcome this limitation. Expansion microscopy (ExM), which improves the practical resolution by physically expanding hydrogel-embedded samples, enables subdiffraction imaging of tissues without specialized super-resolution instruments³¹. However, one challenge for ExM is the dilution of fluorescence signals during physical expansion, which creates a need for signal amplification. Both indirect immunofluorescence and HCR have been used to achieve higher signals in ExM with limited multiplexing capability²⁸. A technique termed magnified analysis of the proteome (MAP) was developed to achieve higher multiplexing by combining expansion with repeated antibody staining and retrieval, demonstrating seven rounds of sequential labeling; however, owing to the slow permeation of the antibodies into the thick expanded tissue samples, each round of primary and secondary antibody staining takes 2–9 d, making the approach slow and laborious for higher levels of multiplexing³⁸.

To solve this problem, we combined ExM with SABER (Expansion-SABER). We modified the 5' end of the SABER concatemers with an acrydite moiety that could be incorporated into polyacrylate hydrogels. Using the original ExM protocol³¹ with Immuno-SABER for SV2 in mouse retina cryosections, we obtained approximately threefold expansion (Fig. 6a), which is slightly lower than the expansion factor reported before (~4.5-fold) owing to the shrinkage of the gel in the ionic gel re-embedding solution or in the imaging buffer with 0.5× PBS (as was also observed for the expansion-FISH protocol^{31,39}). Higher levels of expansion can be attained by more recent expansion protocols that can achieve 10- to 20-fold expansion^{40,41}. To validate the subdiffraction resolution, we imaged the presynaptic marker Bassoon and the post-synaptic marker Homer1b/c (which are typically ~160 nm apart⁴²) in fixed primary mouse hippocampal neuron culture (Fig. 6b and Supplementary Fig. 9a). While Bassoon and Homer1b/c were readily separated after expansion, they strongly overlapped without expansion. We next demonstrated Exchange-SABER in expansion samples by visualizing six targets (vimentin, collagen IV, rhodopsin, calretinin, GFAP and SV2) in mouse retina cryosections (Fig. 6c). Individual Muller cells labeled by vimentin became clearly distinguishable after expansion (Supplementary Fig. 9b). As the expansion gel samples reached ~350 µm in thickness, we increased both the incubation time and wash time for imagers (to 45 min and 1.5 h, respectively) to achieve optimal imaging quality. We were able to perform six-target imaging in two rounds of imager exchange in 6 h—substantially faster than the MAP protocol, which would take >3 d (ref. ³⁸). To further increase the speed, we incorporated an alternative fluorophore removal protocol using imager strands with a disulfide bond between the DNA sequence and the fluorophore, which allow quick removal of the fluorescence signal by reducing agents (for example, TCEP)⁴³. This approach shortened the signal removal time from 1.5 h to only 10 min (Supplementary Fig. 9c,d).

Discussion

Immuno-SABER provides a new highly multiplexed amplification capability to increase the detection sensitivity and tune it independently for multiple targets. To validate its suitability for tissue imaging, we addressed three main considerations. (1) Specific labeling of targets in commonly used sample preparations. We used SABER to label various protein targets (nuclear, cytoplasmic and membrane) in cell and tissue preparations (FFPE sections, cryosections and expansion samples), and reproduced the expected labeling patterns (Figs. 2, 4, 5 and 6 and Supplementary Figs. 2, 3 and 8). (2) Efficient access to the target and quantifiable amplification without sacrificing resolution. SABER probes can penetrate into whole-mount preparations and reach targets at depths of up to 100 µm (Fig. 2 and Supplementary Fig. 2). Linear amplification with primary antibodies in tissues yielded 5- to 20-fold signal enhancement (typically at a similar level to indirect immunofluorescence for the same target) (Figs. 2 and 3 and Supplementary Fig. 4). Branched SABER generated ~50-fold signal amplification (>5 -fold higher signal than indirect immunofluorescence) and iterative SABER produced over 180-fold signal amplification (surpassing TSA) (Fig. 3 and Supplementary Fig. 5). At these levels of amplification, SABER did not have a significant impact on the resolution for conventional microscopy and tissue imaging (Fig. 2 and Supplementary Figs. 2 and 3). This observation still holds for super-resolution imaging with ExM (Fig. 6 and Supplementary Fig. 9). We also validated that the signal level can be further controlled by concatemer length and the concatemers (<700 nucleotides for linear and <450 nucleotides for branching) do not hinder access to antigens (Supplementary Fig. 4). (3) Multiplexing, through the use of orthogonal concatemers, we showed multiplexed amplification for ten protein targets in the same sample (Fig. 5) and simultaneous use of linear and branched SABER (Fig. 4). The ability to program and control the fold amplification of signal for each target is valuable for multiplexed imaging of target proteins expressed with a high dynamic range. To lead the way toward higher multiplexing, we performed *in situ* testing of a pool of 32 sequences from our 50 *in silico*-designed sequences³⁰, and detected minimal crosstalk for only 1 of the 1,024 pairs (Supplementary Fig. 6). Owing to the simplicity of sequence design, it is straightforward to upscale the orthogonal pool size and reach much higher multiplexing levels.

Immuno-SABER is compatible with widely available sample preparation and microscopy platforms (such as standard wide-field and confocal microscopes or slide scanners), promising easy adoption by research and clinical laboratories. SABER enables higher throughput in two ways. First, it saves sample preparation time by allowing application of multiple primary antibodies in one step, thus alleviating the need for multiple time-consuming cycles of antibody staining. This advantage is particularly important for thicker tissue samples such as whole-mount and ExM preparations (where we imaged six targets in approximately threefold expanded tissues within 3–6 h in contrast to the ~4–5 d that would be required in protocols such as MAP³⁸). Second, SABER saves image acquisition time as high signal level supports reduced camera exposures. A 10- to 100-fold improvement in image acquisition can deliver one to two orders of magnitude enhancement of throughput, which is needed for discovery-oriented profiling of cells and tissues and large-scale mapping projects.

To implement Immuno-SABER for new targets, we recommend aiming to conjugate a limited number (1–3 copies) of bridge oligonucleotides per antibody to preserve the antigen recognition efficiency of the antibody and to validate the specificity of the staining by comparison to unconjugated controls. For relative quantifications, a baseline control can be utilized (on the same sample (Supplementary Fig. 4) or on comparable samples (Fig. 3)). As in any signal amplification method, increasing amplification level is expected to yield a wider spread in the population, while moving the mean to higher intensities. For branched amplification, it is

theoretically possible that the small differences in the initial binding might lead to larger variations. However, at the population level, these factors should have an even effect on the sample (unless there is critical internal heterogeneity, like the target being more buried in some locations, which would bias antibody binding for all immunofluorescence studies), as suggested by the consistency of the stainings in Supplementary Fig. 4. Advantageously, Immuno-SABER avoids in situ enzymatic reactions, which are hard to reproduce in a quantitative manner as the enzyme activity is hard to control. For SABER, every step relies on simple DNA–DNA binding that can be primarily modulated by salt and formamide concentration, as well as temperature.

In summary, Immuno-SABER is a simple and effective method for multiplexed and sensitive in situ protein detection with individually programmable signal amplification. Immuno-SABER enables higher throughput for imaging assays ranging from super-resolution studies to centimeter-scale tissue mapping efforts. The SABER strategy is also directly applicable for RNA and DNA targets as we recently demonstrated in our SABER-FISH work³⁰. We expect that the unified SABER framework will be useful to a broad range of researchers, complementing single-cell RNA sequencing analysis with the functional information of protein expression, and will prove valuable for a wide spectrum of potential applications including tissue atlases^{44,45}, tumor and disease profiling, in situ single-cell validation for bulk assays such as flow or mass cytometry, and CITE-Seq⁴⁶, as well as digital pathology and biomarker screening and discovery.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, statements of code and data availability and associated accession codes are available at <https://doi.org/10.1038/s41587-019-0207-y>.

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Author contributions

S.K.S. and Y.W. conceived the study, performed experiments, analyzed the data and wrote the manuscript. J.Y.K. contributed to conceptualization of the study, protocol optimization and experimental design. A.Z., Y.Z., W.X., S.Y. and M.L. provided experimental assistance. K.K., C.Y. and M.C. performed data analysis and visualization. B.J.B. contributed to protocol optimization. S.W.L. provided the retina samples for staining. G.P. contributed to contextualization of SABER application, and provided guidance and tissues for the FFPE study. E.S.B. contributed to the expansion study. P.S.K. contributed to the neuronal study. G.M.C. provided scientific guidance and contributed to study supervision. P.Y. conceived and supervised the study, and wrote the manuscript. All authors edited and approved the manuscript.

Competing interests

S.K.S., Y.W., J.Y.K., B.J.B. and P.Y. are inventors for a US patent application based on this work (PCT/US2018/013019). P.Y. is a co-founder of Ultivue, Inc. and NuProbe Global.

Additional information

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Methods

Technical details for the experiments are given below. We also provide generalized, step-by-step protocols for antibody–DNA conjugation, purification of conjugated antibodies and performing Immuno-SABER in cells, FFPE samples and fixed cryosections in the Supplementary Protocols.

Additionally, we will keep an updated ‘Frequently Asked Questions’ section available at <http://www.immuno-saber.net/> and <http://www.saber-imaging.net/>.

Primer exchange reaction sequences and preparation of SABER concatemers.

In vitro extension of primers. Concatemer extensions were prepared as described previously³⁰. Typically, 100- μ l reactions were prepared in 1 \times PBS (diluted from a 10 \times stock; Thermo Fisher, AM9625) with final concentrations of 10 mM MgSO₄, 400–1,000 U ml^{−1} Bst LF polymerase (NEB, M0275L; or McLab, BPL-300), 600 μ M each of dATP, dCTP and dTTP (NEB, M0275L), 100 nM of Clean.G (5′-CCCCGAAAGTGGCCTCGGGCCTTTGGCCCCAGGCCACTTTCG-3′) hairpin²⁹, 50 nM–1.5 μ M hairpin, and water to 90 μ l. The Clean.G hairpin has a 5′ stretch of C. Preincubation with Clean.G helps to get rid of the impurities in the dHTP mixtures (made up of dATP, dTTP and dCTP), which may have small amounts of dGTP contamination²⁹. After Clean.G incubation for 15 min at 37°C, 10 μ l of 10 μ M primer was added to obtain 1 μ M final concentration, and the reaction was incubated for another 1–3 h at 37°C followed by 20 min at 80°C to heat inactivate the polymerase. Reaction products can be stored at −20°C for several months. In our demonstrations, PER products were diluted into concatemer hybridization solutions for binding to the bridge sequences. Alternatively, concatemers can be purified and concentrated using a MinElute (Qiagen, 28004) kit with distilled water elution to reduce volume and salt concentration from the reaction condition. Primer sequences and details of the extension conditions utilized for Supplementary Fig. 6 are listed in Supplementary Table 1. These primer sequences were presented as the 3′ tail of the complements of the bridge strands given in Supplementary Table 2 in the following format: 5′-bridge*-TT (spacer nucleotides)–9-nucleotide primer sequence-3′, where the asterisk denotes the reverse complement. For the extensions in Supplementary Fig. 6, the 25-nucleotide tester* bridge was used for all the primers. The strands were obtained from IDT. Primers were synthesized and provided with standard desalting. Hairpins were ordered with 3′ inverted dT modification to ensure they cannot be extended. Owing to the modification, they were ordered with HPLC purification, which is critical for their performance. Details of the primer and hairpin design criteria are described in our previous work^{29,30}. For primary concatemers, we utilize concatemers of lengths of up to 600–700 nucleotides on the basis of empirical experience³⁰.

Gel electrophoresis. After extension, for internal quality control, the lengths of the concatemers were evaluated by diluting 1 μ l of in vitro reaction with 19 μ l water. For quality control, samples were then run on 1–2% E-Gel EX agarose gels (Thermo Fisher, G402001) for 10 min on the E-gel apparatus (Invitrogen, iBase) alongside a 1 kb Plus DNA Ladder (Invitrogen) and imaged with the SybrGold channel on a Typhoon FLA 9000 scanner.

For the comparison gel in Supplementary Fig. 6, unpurified concatemers were run using 6% Tris–borate–EDTA (TBE)–urea PAGE gels (Thermo Fisher) at 55°C. The gel was prerun for 1 h before loading the samples. One hundred and sixty nanograms of Quick-Load Purple low-molecular-weight DNA ladder (NEB, N0557S) was loaded as a size reference. The reaction products were diluted 1:7 with 2 \times urea loading dye and denatured at 95°C for 5 min. Nine microliters from each sample was loaded on the gel. Both samples and the ladder were denatured. Samples were run for 20 min at 75 V and at 130 V for 1 h. Gels were stained with 1:10,000 SybrGold in 0.5 \times TBE for 30 min and scanned on a Typhoon FLA 9000 scanner.

Imager strands. SABER imager strands are 20-nucleotide DNA oligomers with fluorophores at the 5′ end. The imagers used for our experiments also carry a 3′ inverted dT modification, which is optional and is not essential for SABER experiments. Imagery were designed to bind the dimers of the primer unit sequence to achieve the stable but easily reversible binding that is necessary for DNA-exchange imaging. Hence the format of the imager sequence is as follows: 5′-fluorophore-TT-primer*-T-primer*-T-inverted dT-3′. They were ordered from IDT with a 5′ fluorophore (Atto488, Atto565, Alexa Fluor 647 or Alexa Fluor 750), 3′ inverted dT modification and HPLC purification. They are named as i.primerID#*. Sequences are listed in Supplementary Table 3.

Branching primers. For stable hybridization of the secondary (branching) concatemers onto the primary concatemers, trimers of the unit repeat sequence were used as bridges, creating a 30-nucleotide hybridization sequence. Hence, branching primers are designed in the format: 5′-p.1*-T-p.1*-T-p.1*-TTT-p.2-3′, where p.1 is the primer used for the primary concatemer and p.2 is the primer for the secondary concatemer.

Similarly, for iterative branching, the tertiary concatemer is designed to use the trimers of primer 2 as the bridge, in the format 5′-p.2*-T-p.2*-T-p.2*-T-p.2*-T-p.2-3′, where p.3 is the primer for the tertiary concatemer and T's are spacer nucleotides.

Antibody–DNA conjugation and purification. *Conjugation.* The conjugation involves crosslinking of thiol-modified DNA oligonucleotides to lysine residues on antibodies in a non-sequence-specific way. In brief, 25 μ l of 1 mM 5′-thiol-modified DNA oligonucleotides (IDT) was activated by 100 mM dithiothreitol (DTT) (Thermo Fisher, 20291) for 2 h at room temperature in the dark, and purified using NAP5 columns (GE Healthcare Life Sciences, 17-0853-02) to remove excess DTT. Antibodies formulated in PBS only (or with sodium azide) were concentrated using 0.5-ml 50-kDa Amicon Ultra Filters (EMD Millipore, UFC510096) to 2 mg ml^{−1} and reacted with maleimide–PEG₂–succinimidyl ester crosslinkers (Thermo Fisher, 22102) for 1.5 h at 4°C (100 μ g antibodies: 2.5–3.4 μ l of 0.85 mg ml^{−1} cross-linker). Antibodies were then purified using 0.5-ml 7-kDa Zeba desalting columns (Thermo Fisher, 89883) to remove excess cross-linkers. Activated DNA oligonucleotides were incubated with antibodies (molar ratio of antibody to single-stranded DNA of ~1:11) overnight at 4°C. The final conjugated antibodies were washed using 2-ml 50-kDa Amicon Ultra Filters six times to remove non-reacted DNA oligonucleotides. The list of bridge sequences used for conjugation is provided in Supplementary Table 2. The list of antibodies and the corresponding bridge sequences (IDT) used for each staining, as well as the capture and toehold strands for purification is provided in Supplementary Table 4. Conjugated antibodies were diluted at ratios of between 1:50 and 1:200 for immunostaining. A detailed step-by-step protocol for general use is available in the Supplementary Protocols.

Purification. To increase the staining efficiency, conjugated antibodies can optionally be purified using a DNA-toehold-mediated affinity pulldown protocol (Supplementary Fig. 1a). For this, 200 μ l of high-capacity streptavidin agarose (Thermo Fisher, 20357) was centrifuged, washed three times using 500 μ l of PBS, and incubated with 10 μ l of 1 mM biotin-labeled binding sequences in 300 μ l PBS with 0.1% Triton X-100 for 30 min at room temperature. The agarose was then washed twice with PBS with 0.1% Triton X-100, followed by blocking with 250 μ l of blocking buffer (2% BSA + 0.1% Triton in PBS) for 1 h at room temperature with rotation. The agarose was then centrifuged and resuspended with 200 μ l of incubation buffer (1% BSA + 0.1% Triton in PBS) containing the DNA-conjugated antibodies, followed by rotation at 4°C for 1 h. The sample was centrifuged at 4°C and washed twice with 200 μ l incubation buffer. The bound antibodies were recovered by adding 20 μ l of 1 mM toehold strands (listed in Supplementary Table 4) in 200 μ l of incubation buffer. After centrifugation, the supernatant was collected and the agarose was washed three times with 300 μ l of washing buffer (PBS + 0.1% Triton); supernatant was collected after each wash. The supernatant was pooled together and the buffer was exchanged using 2-ml 50-kDa Amicon Ultra Filters six times to remove toehold DNA oligonucleotides. Binding sequences and toehold sequences were designed using NUPACK (<http://www.nupack.org/>)^{36,31,32} and are provided in Supplementary Table 4.

SDS–PAGE. To examine DNA–antibody conjugation, antibodies were denatured in LDS sample buffer (Thermo Fisher, NP0007) without reducing reagents (for example, DTT or 2-mercaptoethanol) at 90–95°C for 3 min, and left to cool down to room temperature. The samples were run on 3–8% Tris–acetate PAGE gels (Thermo Fisher, EA03752BOX) in NuPAGE Tris–Acetate SDS Running Buffer (Thermo Fisher, LA0041) at 80 V for 30 min and 120 V for 3.5 h. The gels were stained with SimplyBlue safe stain (Thermo Fisher, LC6060) according to the manufacturer's manual, and imaged using a Biorad Gel Doc EZ imager system. Note that BSA should be avoided in the purification step if the sample needs to be examined using PAGE gels.

Microtubule staining in cell culture and full-width half-maximum analysis.

Cell culture and staining. BS-C-1 cells (ATCC CCL-26, African green monkey) were grown in DMEM (Gibco, 10564) supplemented with 10% (vol/vol) serum (Gibco, 10437), 50 U ml^{−1} penicillin and 50 μ g ml^{−1} streptomycin (Gibco, 15070) and cultured at 37°C in the presence of 5% CO₂. They were plated on eight-well Ibidi glass-bottom μ -slides (Ibidi, 80826) and grown to 50–60% confluency. For the cell culture experiments in Figs. 2 and 3, cells were fixed with 4% paraformaldehyde (PFA) for 45 min, quenched with 100 mM NH₄Cl in PBS for 20 min and washed with PBS for 5 min. Cells were then permeabilized and blocked in 2% nuclease-free BSA (AmericanBIO, CAS 9048-46-8) with 0.1% Triton in PBS for 30 min. Samples were incubated with DNA-conjugated primary antibodies diluted in incubation buffer (0.1% Triton X-100, 2% nuclease-free BSA, 0.2 mg ml^{−1} sheared salmon sperm DNA, 0.05% dextran sulfate (Millipore, S4030) and 4 mM EDTA (Ambion, AM9260G) in PBS) overnight at 4°C, and then washed three times with PBS with 0.1% Triton X-100 and 2% BSA for 10 min. Samples were then washed twice with PBS for 5 min and post-fixed using 5 mM BS(PEG)₂ (Thermo Fisher, 21581) in PBS for 30 min, followed by quenching in 100 mM NH₄Cl in PBS for 5 min. The incubation with the primary concatemer was performed at 37°C in 20% formamide (deionized; Ambion, AM9342), 10% dextran sulfate and 0.1% (vol/vol) Tween-20 in 2 \times saline sodium citrate (SSC) buffer with 0.2 mg ml^{−1} sheared salmon sperm DNA for 3 h. The 650-nucleotide primary concatemers prepared in vitro by PER were diluted in this buffer at a 133 nM final primer concentration (primer concentration in the PER mix is considered a proxy for the concatemer concentration after the reaction, as all primers are expected to be extended by the

catalytic hairpins, which are provided in excess). After concatemer hybridization, the samples were washed for 5 min at room temperature with 45% formamide in PBS and three times for 10 min each with PBS + 0.1% Triton X-100 at 37 °C. Branching hybridization was performed overnight at 37 °C in 30% formamide, 10% dextran sulfate and 0.1% (vol/vol) Tween-20 in 2× SSC with 0.2 mg ml⁻¹ sheared salmon sperm DNA and the 450-nucleotide secondary concatemers at a final concentration of 133 nM. Samples were washed for 5 min at room temperature with 45% formamide in PBS and three times for 10 min each with PBS + 0.1% Triton X-100 at 37 °C. Imagers were hybridized at a 1–1.5 μM final concentration in PBS + 0.1% Triton X-100 for 1 h at room temperature (hybridization duration with the imagers can be decreased for faster preparation), followed by a 5-min wash with PBS + 0.1% Triton X-100 and two 5-min washes with PBS. Samples were stained with 4 μg ml⁻¹ DAPI (Invitrogen, D1306) in PBS for 10 min and washed twice for 1 min with PBS. Imaging was performed in SlowFade with DAPI (Invitrogen, S36938) embedding medium.

Imaging. For the images shown in Fig. 2a and Supplementary Figs. 2a and 3e, a Leica SP5 confocal microscope with a 63×/1.3 NA glycerol objective was used. A white-light laser (470–670 nm) was used at 650 nm for excitation of Alexa Fluor 647, and a photomultiplier tube (PMT) was used for detection.

Full-width half-maximum analysis. The best focus planes for isolated microtubules were manually selected from z stacks. Line plots (2.5 μm) were drawn across isolated microtubules in this single plane image using Fiji³³. Small rectangular background areas were manually drawn to obtain the background values around selected microtubule lines. Average background was subtracted from the line-plot values. Gaussian curves were fit to the background-subtracted values; FWHM was calculated on the basis of the fits and average values were obtained by analyzing 30–45 line plots from five images per condition using a Python script. A similar calculation was performed for 200-nm dark red FluoSphere beads (Invitrogen, F8807). Distributions are displayed as box plots. Line plots with multiple discernible peaks were discarded. A two-sample *t* test was performed to check for statistical significance.

Lamin B staining in cell culture for amplification quantification. *Staining.* HeLa cells (human) were cultured and stained following the general protocol as described above for microtubules, with the following changes. Before the secondary antibody incubation, blocking was performed in 2% donkey serum with 1.5% BSA, 0.1% Triton and 0.2 μg ml⁻¹ sheared salmon sperm DNA for 15 min. Secondary antibodies were also diluted in the same buffer, and stained for 1.5 h at room temperature. Post-fixation was done for 10 min with 4% PFA. The following conditions were used for primary concatemer hybridization: 125 nM for 2 h at 37 °C in 20% formamide, 10% dextran sulfate and 0.1% (vol/vol) Tween-20 in 2× SSC with 0.2 mg ml⁻¹ sheared salmon sperm DNA. Before and after the concatemer hybridization, a block and wash was done for 10 min with 0.1% Triton X-100 in PBS, 0.2 μg ml⁻¹ salmon sperm DNA and 3.3% dextran sulfate for 10 min. Imager hybridization 1 was performed with 1 μM Alexa Fluor 647–i.27* for 75 min at room temperature. For linear amplification samples, imager hybridization 2 was performed similarly with Alexa Fluor 647–i.28*. Branch concatemer hybridization was performed overnight at 37 °C with 100 nM extended concatemer in the same buffer as the primary concatemer hybridization. For branching samples, imager hybridizations 2 and 3 were done with Alexa Fluor 647–i.30* and Alexa Fluor 647–i.25*, respectively. Sequences and extension conditions are listed in Supplementary Table 5.

Imaging. Twenty-one- to 26-plane z stacks were imaged using a Zeiss Axio Observer Z1 with a 100×/1.4 NA oil objective. Alexa Fluor 647 (excitation at 590–650 nm, detection at >665 nm) images were acquired with 10% of the LED power at 150-ms exposure, and DAPI (excitation at 300–400 nm, detection at 420–470 nm) with 20% power at 100 ms exposure.

Signal quantification. Maximum projections of the z stacks were created using Fiji. DAPI images were acquired and used to create nuclear masks in MATLAB and Image Processing Toolbox. Mean Alexa Fluor 647 signal per pixel was calculated within the mask region for each cell.

SABER application, quantification and analysis on human tonsil formalin-fixed paraffin-embedded sections. *Preparation of formalin-fixed paraffin-embedded tonsil samples and antigen retrieval.* Human specimens were retrieved from the archives of the Pathology Department of Beth Israel Deaconess Medical Center under the discarded and excess tissue protocol as approved in Institutional Review Board (IRB) Protocol 2017P000585. We have complied with all relevant ethical regulations. Five-micrometer sections were cut with a rotary microtome, collected in a water bath at 30 °C, transferred to positively charged glass slides and baked at 60 °C for 2 h. For antigen unmasking, slides were placed on a PT-Link instrument (Agilent), which allows the entire pretreatment process of deparaffinization, rehydration and epitope retrieval (with citrate buffer) to be combined into a single step. Slides were held at 4 °C in PBS until staining. Antigen-retrieved FFPE sections used in the multicolor experiments in Fig. 4 were acquired from Ultivue, Inc.

Staining of antigen-retrieved formalin-fixed paraffin-embedded tonsil samples.

After antigen retrieval, sections were optionally stored in PBS at 4 °C for up to 2 weeks. For staining, sections were washed in PBS for 15 min and outlined with a hydrophobic pen (ImmEdge Hydrophobic Barrier PAP Pen; Vector Laboratories, H4000) or enclosed in a removable chamber (Ibidi, 80381). At this stage, mild 1-h prebleaching with 1% H₂O₂ in PBS can optionally be applied (for TSA). This step is not necessary for Immuno-SABER. We incorporated this step for the Ki-67 stainings in Fig. 3 and Supplementary Fig. 5, to keep the conditions similar to TSA preparations, which benefit from a prebleaching step. Samples were blocked for 1 h with PBS containing 2% BSA and 0.1–0.3% Triton X-100, with three buffer exchanges. DNA-conjugated primary antibodies were diluted in the blocking solution supplemented with 0.2 μg ml⁻¹ sheared salmon sperm DNA, 0.05% dextran sulfate and optionally 4 mM EDTA, and incubated with the samples overnight at 4 °C in a humidified chamber. The antibodies and respective bridge sequences used for the experiments are listed in Supplementary Table 4. Depending on the antibody, this step can be shortened to 1 h when performed at room temperature. Excess antibodies were removed by washing at room temperature three times for 15 min with PBS containing 2% BSA and 0.1–0.3% Triton X-100, and twice for 5 min with PBS. Bound antibodies were then post-fixed with 5 mM BS(PEG)₅ in PBS for 30 min at room temperature, followed by quenching in 100 mM NH₄Cl in PBS for 5 min, and washed for 15 min with PBS with 0.1% Triton X-100 at room temperature. Post-fixation is critical to ensure that the antibodies are not washed away during further labeling and imaging. The incubation with the primary concatemer was performed at 37 °C in 20–30% formamide, 10% dextran sulfate and 0.1% (vol/vol) Tween-20 in 2× SSC with 0.2 mg ml⁻¹ sheared salmon sperm DNA for between 1 h and overnight. Concatemers prepared *in vitro* by PER were diluted in this buffer at a final concentration of 66–150 nM (1:15 to 1:7.5 dilution). For multiplexing, all primary concatemers were incubated simultaneously. After concatemer hybridization, the samples were washed for 5 min at room temperature with 45–50% formamide in PBS and three times for 10 min each with PBS + 0.1% Triton X-100 at 37 °C. For cases where further amplification was desired, branching hybridizations were performed similarly (at 37 °C in 30% formamide, 10% dextran sulfate and 0.1% Tween-20 in 2× SSC with 0.2 mg ml⁻¹ sheared salmon sperm DNA for between 1 h and overnight). Bridge and primer sequences for each target and experiment are given in Supplementary Table 4. For the unamplified sample, the unextended primer with a single imager-binding site (equivalent to two repeats of the primer sequence) was incubated at the same concentration instead of the extended concatemer (for CD8a, bc42_2*–TT–p.25–A–p.25–A; for Ki-67, bc42_0*–TT–p.30–A–p.30–A; sequences are given in Supplementary Table 5).

For different tissue types and combinations of targets for multiplexing, experimental conditions may need to be optimized to achieve the best signal level. For primary concatemers, we recommend using sequences of ≤650 nucleotides, for secondary concatemers sequences of ≤450 nucleotides and for tertiary concatemers sequences ≤250 nucleotides on the basis of empirical experience. For iterative amplification, the wash temperatures were raised to 42 °C and the 45–50% formamide in PBS wash step was performed once as a final wash only at the end of all iterations (rather than after each amplification round).

Fluorophore hybridization and dehybridization. Imagers were hybridized at a final concentration of 1–1.5 μM in PBS + 0.1% Triton X-100 for 1–2 h at room temperature, followed by a 5-min wash with PBS + 0.1% Triton X-100 at 37 °C and two 5-min washes with PBS at room temperature. Lengthened hybridization times were used for convenience; for faster protocols, hybridization duration for the imagers can be decreased with similar performance. Samples were stained with 1 μg ml⁻¹ DAPI in PBS for 10 min and washed twice for 1 min with PBS. For multiplexing experiments, samples were imaged in PBS shortly after preparation and coverslips were temporarily secured with Fixogum (Marabu). Applying coverslips to the samples for imaging is optional depending on the instrument type. Imagers were removed with a 10-min incubation at room temperature in 50% formamide in PBS, followed by two 5-min washes with PBS at room temperature. A new round of imager hybridization was performed as above.

For experiments with a single round of imager hybridization (no multiplexing or only spectral multiplexing), samples were embedded in SlowFade with DAPI, secured with nail polish and imaged on the same day or embedded with ProLong Diamond (Invitrogen, P36971) and incubated at room temperature overnight for curing (Supplementary Note 2).

Tyramide signal amplification. For FFPE samples, the anti-rabbit IgG Alexa Fluor 647 Tyramide SuperBoost kit (Life Technologies, B40916) was used according to the manufacturer's recommendations. The optional step of 1-h prebleaching with 1% H₂O₂ in PBS was applied. Tyramide–Alexa Fluor 647 was incubated for 2.5 min, 7.5 min and 10 min. Longer incubation times (≥7.5 min) were observed to cause increased blurring of the signal, making shorter incubation times more favorable (as suggested by the manufacturer).

Fluorescence imaging. FFPE samples in Figs. 2 and 3, and Supplementary Figs. 2, 3, 4, 5b,c and 7a were imaged with an Olympus VS-120 system equipped with an Orca R2 monochrome (16-bit) camera using a 20×/0.75 NA air objective with a pixel size of 0.320 μm, and single-plane tile scans were acquired. For Fig. 3 staining,

exposure times of 35 ms and 30 ms were used for DAPI and Ki-67, respectively. For the CD8a staining in Figs. 1 and 2, an exposure time of 350 ms was used.

For the expanded views in Fig. 3f and Supplementary Fig. 5f a Leica SP5 confocal microscope with 63×/1.3 NA glycerol objective was used. A white light laser (470–670 nm) was used at 650 nm for excitation of Alexa Fluor 647, and a PMT was used for detection.

Multiplex FFPE samples in Fig. 4 and Supplementary Fig. 7b were imaged with a Perkin Elmer Vectra Polaris microscope in whole-slide scanning configuration with a 20×/0.80 NA air objective. For Fig. 4c, imagers were first hybridized for three of the targets (as depicted in the figure), followed by nuclear DAPI staining, washing, coverslipping and whole-section imaging in five channels (three markers plus DAPI and autofluorescence). This was followed by dehybridization as described above. The second set of imagers was similarly hybridized and the sample was reimaged. Images obtained in .qptiff format were converted to .tif with Imapris software and aligned using the 'Align by line ROI' plugin on Fiji using the DAPI fluorescence in each cycle; overlays for display were assembled in Adobe Photoshop.

Signal quantification. For quantification of signal amplification in CD8a labeling of tonsil samples in Fig. 2 and Supplementary Fig. 3b,d, ROIs covering 0.03–1.2 mm² tile scan areas were selected after manual inspection to exclude areas with autofocusing errors or sectioning imperfections, and a CellProfiler routine was used first to remove masked autofluorescent structures identified in an independent channel (using global robust thresholding) and then to calculate mean fluorescence signal per pixel for cell regions masked via thresholding of the CD8a signal⁵⁴. For masking of the labeled structures, edges were enhanced by the Canny edge-finding method with automatic threshold calculation. Then, global robust background thresholding was applied. The upper outlier fraction and correction factor for thresholding were adjusted manually to compensate for the difference in overall signal level under different amplification conditions. The background was calculated by averaging the CD8a fluorescence signal of the tissue regions (ensured by the presence of DAPI) outside of the masked signal regions (after dilation of the masked pixels). The final intensity value was obtained by subtracting the mean background value for each ROI from the average signal value for masked cells. Fold amplification of signal was calculated by dividing the mean background-subtracted fluorescence by the unamplified sample (for linear SABER; Fig. 2c,e) or by the linear amplification sample (for branched SABER; Supplementary Fig. 3b,d). Figures were prepared for display using OMERO for ROI selection and scaling⁵⁵.

Nuclear segmentation, analysis of single-nuclei intensity distribution and quantification of fold amplification of signal. We annotated the contours of all the nuclei for 80 patches of 128 × 128 pixels. This dataset was then augmented eightfold using reflections and rotations, resulting in a dataset of 640 images. For each image, an equal number of non-contour pixels were randomly selected to represent the complementary class in the machine learning algorithm. We trained a variant of the U-Net model⁵⁶, adding batch normalization⁵⁷ and residual learning⁵⁸, to classify pixels in two classes (class 1, nuclei contours; class 2, background and nuclei interiors). We used our own implementation in TensorFlow (hyperparameters are given in Supplementary Note 3). The approximate center of each nucleus was then identified through the regional maxima of a Gaussian-blurred version of the inverted contour probability class. These regional maxima were used as seed points for applying a marker-controlled watershed transform. Background objects were eliminated on the basis of mean intensity and area. For estimation of the fold amplification of signal, mean signal intensity from all the nuclei in each tissue sample was summed to obtain the total Ki-67 signal level. Then, the total signal for each sample was divided by the total signal for the unamplified condition. Total values were not corrected for variation in the number of cells in each section (tissue samples are consecutive sections from the same tissue). Histograms that show the distribution of Ki-67 signal across individual cells were plotted using Python.

SABER application and quantification on mouse retina cryosections. Sample preparation. All animal procedures complied with all relevant ethical regulations and were in accordance with the National Institute for Laboratory Animal Research Guide for the Care and Use of Laboratory Animals as approved by the Harvard Medical School Committee on Animal Care. Animals were given a lethal dose of sodium pentobarbital (120 mg kg⁻¹) (MWI, 710101) and enucleated immediately. Eyes were removed and fixed in PFA for 15–30 min. Following dissection, retinas were immersed in 30% sucrose overnight before freezing in TFM (EMS, 72592) and cryosectioning at ~30–40 µm. Eight-well Ibidi glass-bottom µ-slides were treated with 0.3 mg ml⁻¹ poly-D-lysine for at least 30 min, followed by three washes in PBS. Retina sections were immobilized onto the glass and stored at –20 °C. Sections were washed three times with Tris-buffered saline (TBS) + 0.3% Triton X-100 for 10 min. Samples were then permeabilized and blocked in 5% normal donkey serum (Jackson ImmunoResearch, 017-000-001) with 0.2 mg ml⁻¹ sheared salmon sperm DNA and 0.1–0.3% Triton X-100 in PBS for 2 h. Samples were incubated with DNA-conjugated primary antibodies diluted in the incubation buffer (0.1% Triton X-100, 5% normal donkey serum, 0.2 mg ml⁻¹ sheared salmon sperm DNA, 0.05% dextran sulfate and 5 mM EDTA in PBS) overnight at 4 °C, and

then washed three times with PBS with 0.1% Triton X-100, 1% normal donkey serum and 5 mM EDTA for 30 min. Samples were then washed twice with PBS for 5 min and post-fixed using 5 mM BS(PEG)₂ in PBS for 30 min, followed by washing in PBS for 2 min and quenching in 100 mM NH₄Cl in PBS or 1× TBS for 10 min.

Immuno-SABER. Extended concatemers were diluted in the 1:7.5 to 1:20 range (depending on the target density) in incubation buffers for overnight hybridization at room temperature. The following two concatemer hybridization buffers were used: buffer 1 (40% formamide, 10% dextran sulfate, 0.1% Triton X-100, 0.02% sodium azide and 5 mM EDTA in PBS) and buffer 2 (30% formamide, 10% dextran sulfate, 0.1% Triton X-100, 0.02% sodium azide and 5 mM EDTA in PBS). Buffer 1 was used for incubation with primary concatemers and buffer 2 was used for branching concatemers. Excess concatemers were removed by washing with 45% formamide, 0.1% Triton X-100 and 5 mM EDTA in PBS for 30 min and twice with 30% formamide, 0.1% Triton X-100 and 5 mM EDTA in PBS for 30 min at room temperature. For branched conditions, 45% formamide was replaced with 40% formamide for the first wash. For the multiplexed imaging experiment, all primary concatemers were incubated simultaneously. Bridge and primer sequences for each target and experiment are given in Supplementary Table 4.

To quantify linear SABER amplification for cone arrestin, the amplification samples were incubated with SABER concatemers extended from tester*–TT–p.28. The unamplified samples were hybridized with the unextended primer tester*–TT–p.28–a–p.28, which carries one imager-binding site (equivalent to two repeats of the primer sequence) instead of the extended concatemer (at the same final concentration). For quantification of branched SABER for cone arrestin, the branching sample was first incubated with the primary concatemers extended from tester*–TT–p.28, followed by incubation with secondary concatemers extended from the 28*–T–28*–T–28*–TTT–p.25 primer.

For iterative SABER quantification for SV2, the unamplified sample was hybridized with the unextended primer bc42_3*–TT–p.27–a–p.27. The linear amplification sample was hybridized with the primary concatemer extended from bc42_3*–TT–p.27. The branching sample additionally hybridized with the secondary concatemer 27*–T–27*–T–27*–TTT–p.28. The iterative amplification sample was additionally hybridized to the tertiary concatemer extended from 28*–T–28*–T–28*–TTT–p.32. Sequences are listed in Supplementary Table 5.

Fluorescence imaging. Fluorophore-labeled imager strands were diluted to 250 nM–1 µM final concentration range in 0.1% Triton X-100 in PBS, and were incubated with samples for 30 min, followed by washing using 0.1% Triton X-100 in 0.5× PBS three times. Samples were left in PBS during image acquisition. For multiplexing, imager strands were dehybridized by incubating three times with 0.1× PBS and 30% formamide for 10 min, followed by three 5-min washes with 1× PBS to remove the residual formamide before rehybridization of imager strands as above.

For ten-color multiplexing, the entire experiment was done using seven exchange rounds (round 1, calbindin and vimentin; round 2, GFAP and PKCα; round 3, collagen IV and calretinin; round 4: rhodopsin; round 5, VLP1; round 6, SV2; round 7: cone arrestin) with an average of 1.5 h per round (including 30 min for imager hybridization, 15 min for washing of excess imager strands, 15 min for imaging and 30 min for imager strand displacement). It should be noted that the duration for each step is sample dependent, with thicker samples requiring a longer time to ensure complete penetration and signal removal. To ensure the best signal and simplify the multiplexed experimental design, we allowed excess time for each step; however, it is possible to shorten incubation and washing times upon optimization.

All images for mouse retina sections were acquired using a Zeiss Axio Observer with LSM 710 scanning confocal system with a 20×/0.8 NA air objective. The images were 512 × 512 pixels or 1,024 × 1,024 pixels and acquired at acquisition speed 7. Each image was acquired with frame averaging of two. Atto488 was visualized using a 488-nm laser; Atto565 was visualized using a 546-nm laser; and Alexa Fluor 647 was visualized using a 633-nm laser. To remove imager strands, samples were washed three times with 30% formamide and 0.1% Triton X-100 in 0.1× PBS. The samples were left in PBS during imaging. Acquired images were scaled and colored for display using Fiji⁵³ and Photoshop.

Tyramide signal amplification. For retina cryosections, Alexa Fluor 647 TSA kit 6 with HRP–goat anti-mouse IgG (Life Technologies, T20916) was used following the manufacturer's recommendations, without the optional prebleaching step and with a 7.5-min tyramide incubation.

Quantification. For quantification of signal amplification in retina samples (SV2 and cone arrestin), the mask regions were selected manually using Fiji, and mean fluorescence intensity was calculated. The background was calculated by averaging the fluorescence signal of six randomly selected regions outside the retinas. The final fold-amplification values were obtained by subtracting the background value from the average signal value for each condition and normalizing that by the unamplified condition (Fig. 2d and Supplementary Fig. 5d) or the linear condition (Supplementary Fig. 3c,d).

SABER application on whole-mount retina samples. *Sample preparation and staining.* Whole-mount retina staining was performed on the free-floating samples, which allowed reagents to penetrate from both sides of retinas. The sample preparation was conducted with a similar protocol as above but with longer incubation and wash times. DNA-conjugated primary antibodies were incubated for 40 h at 4 °C, and washed three times for 1 h. DNA extensions were incubated for 40 h at room temperature, and washed three times for 1 h. Bridge and primer sequences for each target are given in Supplementary Table 4. Fluorescent oligonucleotides were incubated for 2 h at room temperature, followed by three 30 min washes with 0.5× PBS. For imaging, retinas were flattened by creating four incisions and mounted on a glass slide. After flattening, the thickness of whole-mount retina samples was typically ~160 to 180 µm.

Imaging. The images were acquired using a Zeiss Axio Observer with an LSM 710 scanning confocal system with a 10×/0.45 NA air objective. A z stack of 98 sections with 1-µm spacing was taken for each target.

PER sequence crosstalk analysis. For microtubule staining in Supplementary Fig. 6, BS-C-1 cells were grown in glass-bottomed 96-well plate (Ibidi, 89626) with 5,000 cells per well. Cells were fixed with 4% PFA for 15 min, and quenched with 50 mM NH₄Cl in PBS for 7 min. Cells were then permeabilized and blocked in 0.1% Triton X-100, 0.1% Tween-20, 2% nuclease-free BSA (AmericanBIO, CAS 9048-46-8) and 0.2 mg ml⁻¹ sheared salmon sperm DNA (Thermo Fisher, AM9680) in PBS for 1 h. Samples were incubated with DNA-conjugated primary antibodies diluted in incubation buffer (0.05% Triton X-100, 0.05% Tween-20, 2% nuclease-free BSA, 0.2 mg ml⁻¹ sheared salmon sperm DNA, 0.05% dextran sulfate (Millipore, S4030) and 5 mM EDTA in PBS) overnight at 4 °C, and then washed with washing buffer (0.05% Triton X-100, 0.05% Tween-20, 2% nuclease-free BSA and 5 mM EDTA in PBS) five times (1–2 min for the first two washes and 10 min for the other three washes). Samples were washed with PBS twice and post-fixed using 5 mM BS(PEG)₂ in PBS for 1 h, followed by quenching in TBS for 10 min. Concatemer and imager hybridizations were performed as described for the retina cryosections.

For cognate wells, 20 nM of the corresponding imager strands were incubated, while for crosstalk wells, all other imager strands were incubated with 20 nM for each imager strand. The samples were imaged using a Zeiss Axio Observer Z1 with a 20×/0.8 NA air objective. For signal quantification, the bright-field images were acquired and used to create masks in MATLAB. Average fluorescence signals were calculated within the mask region. The background was calculated as the average fluorescence signals outside the cells. The final fluorescence intensity was the average fluorescence intensity within cell masks minus the background intensity.

Expansion microscopy. The PER primer sequences were modified with acrydite at the 5' end (IDT), and extended as above. Mouse retina samples were stained with DNA-conjugated antibodies, followed by concatemer hybridization as above for retina cryosections. After washing away excess concatemers, a layer of expandable gel was formed according to the original expansion microscopy protocol³¹. In brief, samples were incubated in monomer solution (1× PBS, 2 M NaCl, 8.625% (wt/vol) sodium acrylate, 2.5% (wt/vol) acrylamide and 0.15% (wt/vol) N,N'-methylenebisacrylamide) with ammonium persulfate (APS) and teramethylethylenediamine (TEMED) on ice in open air for 20 min. A gelation chamber was then constructed by placing a #1 coverglass on each side of the tissue section. The specimens were transferred to a humidified incubator and left at 37 °C for 2 h. The samples were then digested using proteinase K (New England BioLabs, P8107S) at a 1:100 dilution in digestion buffer (50 mM Tris pH 8, 1 mM EDTA, 0.5% Triton X-100 and 0.8 M guanidine HCl) at 37 °C overnight. The digested samples were then expanded in excess volumes of deionized water. To prevent expanded samples from shrinkage, they were re-embedded in a non-expandable gel (3% acrylamide, 0.15% N,N'-methylenebisacrylamide with 0.05% APS, 0.05% TEMED). The gel was placed on a bind-silane-treated #1.5 coverglass and immersed in the gel solution on ice for 20 min, followed by gelation at 37 °C for 1.5 h. Five microliters of bind-silane reagent (GE, GE17-1330-01) was diluted with 8 ml of ethanol, 1.8 ml of deionized water and 200 µl of acetic acid. For coating, coverslips were washed with deionized water followed by 100% ethanol and then incubated with bind-silane and air-dried. After re-embedding, imager strands were applied as for normal cryosections, but with longer washing (6×10 min), and were imaged as above in regular retina tissue section imaging. Dehybridization times were also extended to six 10-min incubations.

For the primary neuron culture, the culture was grown on a 12-mm-diameter round #1 coverslips, and stained with anti-Bassoon and anti-Homer1b/c antibodies, followed by DNA-conjugated anti-mouse and anti-rabbit secondary antibodies. The primary antibodies were stained in 5% normal donkey serum with 0.1% Triton X-100, and the secondary antibodies were stained in 5% normal donkey serum with 0.1% Triton X-100, 5 mM EDTA and 0.2 mg ml⁻¹ sheared

salmon sperm DNA. Concatemer incubations were done as for retina cryosections. The expandable gel (19% (wt/vol) sodium acrylate, 10% (wt/vol) acrylamide, 0.05% (wt/vol) N,N'-methylenebisacrylamide in PBS) was formed by placing the coverslip against a parafilm sheet with 20 µl of expansion gel solution in between. The gel was then digested and expanded as above. The gel was transferred to a coverslip dish (Ibidi, 81148) and was incubated with the imager strands (Atto488-i.30* and Atto565-i.26*) in 0.5× PBS without re-embedding and left in 0.5× PBS during confocal imaging, which gave a similar expansion factor of approximately threefold. The imager strands were incubated for 30 min followed by three 10-min washes using 0.5× PBS. After imaging, the imager strands were washed five times for 10 min, and imaging was performed as described above for retina cryosections.

For TCEP reduction of fluorophores in expanded samples, the samples were incubated with 50 mM TCEP (Sigma, 646547) diluted in 1× PBS and the fluorescence signal was monitored in a time course of 10 min.

Statistics and reproducibility. Cell experiments in Fig. 2b and Supplementary Figs. 2a–c, 3e,f and 4 were performed a number of times with similar results before the final data were quantified for a single experiment. All retina histology experiments (Figs. 2d–f, 5 and 6, and Supplementary Figs. 2d, 3c,d, 5d and 9) were conducted at least twice with similar results, except the controls in Supplementary Fig. 8, which were performed as a single dataset with internal controls. The FFPE stainings in Figs. 2 and 3, and Supplementary Figs. 3b–d, 5 and 7a were separately tested in at least two independent optimization experiments, before the quantification on one dataset from consecutive sections from the same source sample. For FFPE sections, single-, double- and triple-antibody stainings were performed separately on multiple occasions before the four-color and six-color multiplexed experiments in Fig. 4 and Supplementary Fig. 7b were performed on the same experiment day with consecutive sections from the same source tissue. The in vitro reactions in Supplementary Fig. 6a were individually optimized and each was tested multiple times before the combined gel run. Cell experiments in Supplementary Fig. 6 were repeated twice with similar results. The plots in Supplementary Figs. 1c, 2b, 8b and 9d are specific to the image shown, and are representative of the overall tendency of the measurements in the respective dataset. Further internal controls through the optimization phase, including negative controls with omission of the antibodies or other components, and signal patterns matching expected morphologies, further increase our confidence in the consistency and reproducibility of the technique in multiple contexts.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The data that support the findings of this study will be provided by the corresponding authors upon reasonable request.

Code availability

The deep learning algorithm and test dataset for automated identification of nuclear contours in tonsil tissues are available at <https://github.com/HMS-IDAC/UNet>. The MATLAB code for nuclear segmentation is available at <https://github.com/HMS-IDAC/SABERProbMapSegmentation>.

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Reporting Summary

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Statistics

For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

- | | |
|-------------------------------------|---|
| n/a | Confirmed |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> The statistical test(s) used AND whether they are one- or two-sided
<i>Only common tests should be described solely by name; describe more complex techniques in the Methods section.</i> |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> A description of all covariates tested |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals) |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> For null hypothesis testing, the test statistic (e.g. F , t , r) with confidence intervals, effect sizes, degrees of freedom and P value noted
<i>Give P values as exact values whenever suitable.</i> |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes |
| <input type="checkbox"/> | <input checked="" type="checkbox"/> Estimates of effect sizes (e.g. Cohen's d , Pearson's r), indicating how they were calculated |

Our web collection on [statistics for biologists](#) contains articles on many of the points above.

Software and code

Policy information about [availability of computer code](#)

Data collection	Commercial softwares licensed by microscopy companies were utilized: Zeiss Zen 2012 (for LSM 710), Leica LAS AF (for Leica SP5), Zeiss Zen 2.3 Pro Blue edition (for LZeiss Axio Observer Z1), Olympus VS-ASW (for Olympus VS120), PerkinElmer Phenochart (version 1.0.2) .
Data analysis	Open-source Python (3.6.5), TensorFlow (1.12.0), and Deep Learning packages have been utilized for machine learning-based nuclei identification (the algorithm and code is available at https://github.com/HMS-IDAC/UNet). We used Matlab (2017b) for watershed-based nuclear segmentation using the identified nuclear contours. Python 3.6 was used for the FWHM calculations, as well as plotting of histograms. We used MATLAB and the Image Processing Toolbox R2016a (The MathWorks, Inc., Natick, Massachusetts, United States) for quantifications in mouse retina sections and for Supplementary Fig. 4. We utilized Cell Profiler 3.1.5 for the quantifications of signal amplification in FFPE samples in Figure 2 and 3. FIJI (version 2.0.0-rc-69/1.52n) was utilized for ROI selections and format conversions. HMS OMERO (version 5.4.6.21) was used for viewing images and assembling figure panels.

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research [guidelines for submitting code & software](#) for further information.

Data

Policy information about [availability of data](#)

All manuscripts must include a [data availability statement](#). This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

Data and Software Availability: The data and essential custom scripts for image processing will be made available from the corresponding authors P.Y. (py@hms.harvard.edu), S.K.S. (Sinem.Saka@wyss.harvard.edu), and Y.W. (yuwang01@fas.harvard.edu) upon request. The deep learning algorithm and test dataset for automated identification of nuclear contours in tonsil tissues is available on <https://github.com/HMS-IDAC/UNet>. The MATLAB code for nuclear segmentation is

Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

- ☒ Life sciences ☐ Behavioural & social sciences ☐ Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see [nature.com/documents/nr-reporting-summary-flat.pdf](https://www.nature.com/documents/nr-reporting-summary-flat.pdf)

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size	Each FFPE experiment batch were performed on consecutive sections from the same source, each containing over 600,000 cells. Due to large number of single cells with tens of distinct germinal center morphologies being present in each section, ROIs from different parts of a whole section was used for quantification of signal improvement for each condition (consecutive sections were used for all the conditions of one quantification experiment). Number of ROIs are noted in the respective figure legends. For quantifications in retina samples, due to conserved staining morphology and low sample-to-sample variability n = 6 z-stacks were acquired from at least 2 retina sections. For Supplementary Fig. 4, minimum 5 z-stacks were acquired for each condition to collect images of 18-45 cells. Number of cells are reported in the graphs.
Data exclusions	Parts of the FFPE tissue sections were excluded from analysis due to automated imaging related aberrations (out-of-focus areas) or tissue preparation aberrations (folding of the thin sections at the edges, or uneven thickness at the edge areas). For FWHM calculations in Supplementary Fig. 2, ROIs that yield lineplots with more than one automatically detected peak were discarded to avoid deviations due to multiple peaks. For Supplementary Fig. 4 cells in the samples were excluded when an external bright fluorescent particle (dust speck, dye aggregate etc.) coincided with the nuclei (as confirmed by manual inspection of the images). The exclusion criteria were pre-established.
Replication	Each FFPE experiment batch were performed on consecutive sections from the same source, each containing over 600,000 cells. For evaluation and quantification of our method, multiple biological replicates were not accumulated in order to avoid the error that would be introduced by the natural biological and preparation variation, and to avoid unnecessary use of human tissue material. In the case of the mouse retina quantifications a minimum of two distinct retinal sections were imaged, and each experiment was performed at least twice. For Supplementary Fig. 4 dataset, 16 different conditions were prepared and each were imaged multiple times (before linear, after linear, before branch, after branch). Although the data was not pooled together for the statistics reported in the figure, low cell-to-cell variability was observed and high consistency was seen across the samples for comparable conditions, suggesting low sample to sample variability.
Randomization	Randomization was not necessary for this study.
Blinding	Blinding was not possible as experimental conditions were mostly evident from the image data.

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

n/a	Involved in the study
<input type="checkbox"/>	<input checked="" type="checkbox"/> Antibodies
<input type="checkbox"/>	<input checked="" type="checkbox"/> Eukaryotic cell lines
<input checked="" type="checkbox"/>	<input type="checkbox"/> Palaeontology
<input type="checkbox"/>	<input checked="" type="checkbox"/> Animals and other organisms
<input type="checkbox"/>	<input checked="" type="checkbox"/> Human research participants
<input checked="" type="checkbox"/>	<input type="checkbox"/> Clinical data

Methods

n/a	Involved in the study
<input checked="" type="checkbox"/>	<input type="checkbox"/> ChIP-seq
<input checked="" type="checkbox"/>	<input type="checkbox"/> Flow cytometry
<input checked="" type="checkbox"/>	<input type="checkbox"/> MRI-based neuroimaging

Antibodies

Antibodies used

The full list is also available in Supplementary Information, Supplementary Table 4.

Ki-67 Cell Signaling #9129, clone: D3B5 (formulated in PBS, Lot: 2), diluted 1:100-1:250 after conjugation

CD8a Cell Signaling #85336 clone: D8A8Y (formulated in PBS, Lot: 4) diluted 1:150 after conjugation

PD-1 Cell Signaling #43248, clone: EH33 (formulated in PBS, Lot: 2), diluted 1:150 after conjugation

IgA Jackson ImmunoResearch #109-005-011 (Lot: 134868), diluted 1:150 after conjugation

CD3e Cell Signaling #85061 clone: D7A6E(TM) XP(R) (formulated in PBS, Lot:2), diluted 1:150 after conjugation

IgM Jackson ImmunoResearch #709-006-073 (Lot: 133627), diluted 1:150 after conjugation

Lamin B Santa Cruz sc-6216 clone:C-20, (Lot: E1115), diluted 1:100

Alpha-Tubulin ThermoFisher #MA1-80017 (multiple lots), diluted 1:50 after conjugation

Cone arrestin Millipore #AB15282 (Lot: 2712407), diluted 1:100 after conjugation

GFAP ThermoFisher #13-0300 (Lot: rh241999), diluted 1:50 after conjugation

SV2 HybridomaBank, Antibody Registry ID: AB_2315387, in house production, diluted 1:25 after conjugation

PKCα Novus #NB600-201, diluted 1:50 after conjugation

Collagen IV Novus #NB120-6586, diluted 1:50 after conjugation

Rhodopsin EnCor Bio #MCA-A531, diluted 1:50 after conjugation

Calbindin EnCor Bio #MCA-5A9, diluted 1:25 after conjugation

Vimentin Cell Signaling #5741S, diluted 1:50 after conjugation

Calretinin EnCor Bio #MCA3G9, diluted 1:50 after conjugation

VLP1 EnCor Bio #MCA-2D11, diluted 1:25 after conjugation

Bassoon Enzo ADI-VAM-#PS003, diluted 1:500

Homer1b/c ThermoFisher #PA5-21487, diluted 1:250

Supplementary

Anti-rabbit IgG (to detect Ki-67 and Homer1b/c indirectly) Jackson ImmunoResearch # 711-005-152 (Multiple lots), 1:90 after conjugation

Anti-mouse IgG (to detect Bassoon indirectly) Jackson ImmunoResearch #715-005-151 (Multiple lots), diluted 1:100 after conjugation

Anti-goat IgG (to detect Lamin B indirectly) Jackson ImmunoResearch # 705-005-147 (Lot: 125860), diluted 1:75 after conjugation

Alternative antibodies used to validate colocalization of VLP1 and Calretinin in Supplementary Fig. 8d-f:

Calretinin (SantaCruz #SC-365956; EnCor Bio #CPCA-Calret; EnCor Bio #MCA-3G9 AP), VLP1 (EnCor Bio #RPCA-VLP1; EnCor Bio #CPCA-VLP1; EnCor Bio #MCA-2D11). All diluted 1:100.

Fluorophore-conjugated secondary antibodies used for reference imaging:

anti-rat-Alexa647 (ThermoFisher #A-21472, 1:200), anti-rabbit-Alexa488 (ThermoFisher #A-21206, 1:200), anti-rabbit-Atto488 (Rockland #611-152-122S, Lot:33901, 1:500), anti-mouse-Alexa647 (ThermoFisher #A-31571, 1:400), anti-goat-Alexa647 (ThermoFisher #A-21447, 1:200), anti-rabbit-Alexa647 (Jackson ImmunoResearch, 711-605-152, Lot: 125197, 1:300).

Validation

All antibodies used are from commercial sources as described. Only antibodies that have been validated by the vendor with in vitro and in situ experiments (for IHC and IF, with images available on the websites) and/or heavily used by the community with publication in several references were used. The validation and references for each are publicly available on the respective vendor websites that can be reached via the catalog numbers listed above. In our experiments, IF patterns matched the distribution of cell types these antibodies were expected to label based on the literature both before and after conjugation with DNA strands.

Eukaryotic cell lines

Policy information about [cell lines](#)

Cell line source(s)

BS-C-1 cells and HeLa cells

Authentication

Cell lines were not authenticated (not relevant for the experiment or results)

Mycoplasma contamination

Cell lines were not tested for mycoplasma contamination (not relevant for the experiment or results)

Commonly misidentified lines
(See [ICLAC](#) register)

No commonly misidentified cell lines were used.

Animals and other organisms

Policy information about [studies involving animals](#); [ARRIVE guidelines](#) recommended for reporting animal research

Laboratory animals	Wild-type CD1 mice (male and female) age P13 or P17 were used for retina harvest.
Wild animals	The study did not involve wild animals.
Field-collected samples	The study did not involve samples collected from the field.
Ethics oversight	All animal procedures were in accordance with the National Institute for Laboratory Animal Research Guide for the Care and Use of Laboratory Animals and approved by the Harvard Medical School Committee on Animal Care.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Human research participants

Policy information about [studies involving human research participants](#)

Population characteristics	We have only used exempt tissue sections for technical demonstration, since we do not derive any biological conclusions, the population characteristics is not relevant for this methodological study.
Recruitment	Not relevant for this study.
Ethics oversight	Human specimens were retrieved from the archives of the Pathology Department of Beth Israel Deaconess Medical Center under the discarded/excess tissue protocol as approved in Institutional Review Board (IRB) Protocol #2017P000585. Informed consent was waived on the basis of minimal risk to participants (which is indirect and not based on prospective participation by patients).

Note that full information on the approval of the study protocol must also be provided in the manuscript.