1 Supplementary Information for

IBEX – A versatile multi-plex optical imaging approach for deep phenotyping and spatial analysis of cells in complex tissues

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34 Supplemental Methods

35 Animals, immunizations, and tissue preparations

36 5-12 week old naïve C57BL/6 and Cxc/9^{-/-} mice were purchased from Jackson Laboratories 37 (Bar Harbor, ME) and maintained at a facility at the NIH. Fluorescent Confetti animals were 38 generated by crossing B6.129P2-Gt(ROSA)26Sortm1(CAG-Brainbow2.1)Cle/J × B6.Cg-Tg(UBC-39 cre/ERT2)1Ejb/2J and LysM-tdTomato mice were generated by crossing LysM-Cre x B6.Cg-40 Gt(ROSA)26Sortm14(CAG-tdTomato)Hze/J (Jackson Laboratory). Fluorescent Confetti animals 41 heterozygous for both transgenes were injected intraperitoneally (i.p.) with tamoxifen 100 µg per 42 gram of body weight in peanut oil (Sigma-Aldrich) on day 0 and day 2 and tissues were collected 43 on day 4 for processing. To evaluate changes in the immune cell composition and LN architecture 44 following immunization, sheep red blood cells (SRBCs, Colorado Serum Company) were prepared 45 by resuspending 3 mL of SRBCs in 10 mL of Hanks' Balanced Salt Solution (HBSS, Sigma-Aldrich). Mice were injected subcutaneously (s.c.) with SRBCs in a volume of 25-50 μ l per site on day 0 and 46 47 day 4 with organs harvested on day 11. Unless stated otherwise, all tissues were treated similarly 48 and embedded as whole organs. Prior to fixation, murine livers and lungs were perfused with PBS 49 in order to remove blood. The ileal portion of the small intestine was excised and prepared using 50 the Swiss roll technique (1). Livers were collected from LysM-tdTomato mice and processed as 51 described previously (2). Lungs were inflated with 10 ml of fixative via a tracheal cannula before 52 harvest (3). Lungs were then tethered to a small weight and fixed overnight in BD 53 CytoFix/CytoPerm (BD Biosciences) diluted in PBS (1:4). Following fixation, all tissues were 54 washed briefly (5 minutes per wash) in PBS and incubated in 30% sucrose for 2 days before 55 embedding in OCT compound (Tissue-Tek).

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57 Additional considerations for IBEX method

58 For optimal results, it is important to be aware of the expiration date of the chrome gelatin alum and to follow these steps for coating chambers and glass slides: 1) spread the adhesive 59 60 evenly over the imaging substrate, 2) dry 1 hour at 60° C, 3) section tissue onto coated chamber or glass slide, 4) dry for 1 hour at 37° C, and 5) work with freshly prepared samples (don't store at 61 62 -20° C). In order to achieve efficient bleaching, always use 1 mg/ml solutions of LiBH4 within 4 63 hours of preparation and wait until large bubbles form before adding to tissue (usually 10 minutes 64 after dissolving in diH₂O). Be careful as LiBH₄ can react violently with water. To avoid flames, 65 always work with small amounts (<10 mg) of LiBH₄. Store LiBH₄ with desiccant and use a new vial 66 of LiBH₄ after 4 weeks of use. As noted in the "IBEX using inverted confocal microscope" section 67 of the Methods section, Brilliant Violet 421 (BV421) and Brilliant Violet 510 (BV510) dyes require 68 LiBH₄ bleaching in the presence of light (metal halide lamp with the DAPI filter). To ensure efficient 69 bleaching of BV421 and BV510 conjugates, we assess bleaching in real time by viewing the LiBH₄-70 incubated samples under the microscope, bleaching a field of view (1-2 minutes), and moving to 71 an adjacent field of view (1-2 minutes). This process is repeated until the entire region of interest 72 is bleached. Of note, the time required for this process is dependent on the imaging area and 73 objective used for bleaching with larger areas requiring longer overall bleaching times. As an example, a tissue section from a mouse LN (1-2 mm) can be readily bleached in 15 minutes using 74 75 this approach. Another consideration is that all images be oriented identically in the Z stack (same 76 begin and end) for proper alignment with SimpleITK. For gross observation and visualization, we 77 found that IBEX-generated images could be aligned using Imaris 9.5.0 (Bitplane) by "adding 78 images" and manually aligning images with the reference manager. Alternatively, the image 79 alignment feature under the image processing menu was used to provide a rough assessment of 80 image registration before submitting IBEX-generated images to the SimpleITK workflow.

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82 Ensuring optimal imaging quality: Antibody validation and controls

83 Selection of antibodies that bind their targets with great specificity while yielding 84 reproducible immunolabeling across different samples is critical for all imaging methods. Wherever 85 possible, we utilize highly cited antibodies previously validated for immunofluorescence of fixed 86 frozen tissues. Upon identifying suitable antibody candidates for our imaging workflow, we note the 87 location (membrane, cytoplasm, nucleus) and tissue distribution of the marker of interest. 88 Additionally, we procure positive and negative control tissues based on the described expression 89 of a particular marker. Finally, we pair the new antibody with previously validated markers that co-90 stain the same cell type. For example, the SPARC antibody (R&D AF941) is reported to label 91 macrophages, fibroblasts, and endothelial cells within human tissues. To validate this antibody, we 92 evaluate the spatial distribution of the anti-SPARC antibody in human LNs co-stained with CD3 93 (negative) and CD11c and CD31 (positive controls). Prior to all iterative imaging, antibodies are 94 tested for their sensitivity to LiBH₄ by pretreating tissue sections with LiBH₄ for 15 minutes. To 95 evaluate whether epitopes are sensitive to LiBH₄, staining patterns for individual antibodies are 96 compared between serial sections with or without LiBH₄ pre-treatment. As an additional control, 97 individual panels are acquired on serial sections in parallel with iterative rounds of imaging. The 98 spatial distribution patterns are then compared between the serially and iteratively acquired images 99 for each antibody to ensure there is no epitope loss or steric hindrance with cyclic imaging. Finally, 100 antibody concentrations vary greatly depending on the tissue, fixation conditions, and imaging 101 system employed. For these reasons, we strongly recommend careful titration of all antibodies prior 102 to IBEX imaging.

103

104 IBEX imaging conditions for inverted confocal microscope

105 Representative sections from different tissues were acquired using an inverted Leica TCS 106 SP8 X confocal microscope equipped with a 40X objective (NA 1.3), 4 HyD and 1 PMT detectors, 107 a white light laser that produces a continuous spectral output between 470 and 670 nm as well as 108 405, 685, and 730 nm lasers. Panels consisted of antibodies conjugated to the following 109 fluorophores and dyes: Hoechst, BV421, BV510, AF488, AF532, JOJO-1, PE, eF570, AF555, 110 AF594, AF647, eF660, and AF700. All images were captured at an 8-bit depth, with a line average of 3, and 1024x1024 format with the following pixel dimensions: x (0.284 μ m), y (0.284 μ m), and z 111 112 (1-1.25 µm). Images were tiled and merged using the LAS X Navigator software (LAS X 113 3.5.5.19976). To ensure proper alignment over distinct imaging cycles, careful attention was paid to the quality of image stitching achieved with the Leica software and z-stacks were set by manual 114 115 inspection of notable features such as unusually shaped nuclei throughout the tissue volume. These unusual features were matched across the z-stack and over multiple cycles of IBEX. 116

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118 Adoption of IBEX to additional imaging systems

119 For adoption of the IBEX protocol to an upright Leica TCS SP8 X confocal microscope, 20-120 30 µm sections were adhered to Super Frost Plus Gold slides (Electron Microscopy Services) 121 coated with 30 µl of chrome alum gelatin. The IBEX protocol was executed as described above 122 with the following exceptions: slides were mounted with a No. 1.5 coverslip (VWR) and antibody 123 panels were designed without BV421 and BV510 conjugated antibodies. These conjugates were 124 omitted because they require the tissue to be immersed in LiBH4 while illuminated with the metal 125 halide lamp, an impossibility for samples mounted on slides. Following image acquisition, 126 coverslipped slides were immersed in PBS until the coverslip floated off. Non-coverslipped slides 127 were incubated with LiBH₄ for 15 minutes, washed extensively in PBS, immunolabeled with the 128 next round of antibodies, mounted with Fluoromount G, and coverslipped. Image acquisition 129 parameters and system configurations were identical to the details listed for the inverted Leica TCS 130 SP8 X confocal microscope with the exception of the 685 nm laser. For adoption of the IBEX 131 protocol to an inverted fluorescent microscope, 5-10 µm sections were adhered to Super Frost Plus 132 Gold slides coated with chrome alum gelatin. Slides were mounted with a coverslip and antibody 133 panels were designed with the following fluorophores and dyes: Hoechst (Biotium), AF488, PE, 134 and AF647. Representative sections were acquired using a Keyence BZ-X800 microscope 135 equipped with a 40X objective, metal halide lamp, and DAPI, GFP, TRITC, and Cy5 filter sets. All 136 images were captured using the auto-exposure settings for each channel at standard resolution 137 yielding an 8-bit image. Haze reduction was applied post-acquisition to improve image quality.

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139 Image analysis and quantification

140 Fluorophore emission was collected on separate detectors with sequential laser excitation 141 of compatible fluorophores (3-4 per sequential) used to minimize spectral spillover. The Channel 142 Dye Separation module within the LAS X 3.5.5.19976 (Leica) was then used to correct for any 143 residual spillover. Threshold identification, voxel gating, surface creation, and masking were 144 performed as previously described (4, 5). For publication quality images, gaussian filters, 145 brightness/contrast adjustments, and channel masks were applied uniformly to all images. Unless 146 stated otherwise, images are presented as maximum intensity projections (MIP) of tiled z-stacks. 147 Quantification of cell surfaces was based on images with unadjusted gamma values. To calculate 148 the amount of signal remaining after LiBH₄ bleaching (Figs. 1D and S7B), the Color Pixel Counter 149 plugin developed by Ben Pichette for FIJI was used (6). The Structural SSIMilarity (SSIM) index, a 150 method for measuring the similarity between 2 images, was used to assess whether LiBH4 151 treatment removed primary antibodies from the tissues (7) (Fig. S2). The surface creation module 152 of Imaris 9.5.0 (Bitplane) was used to segment cells based on the nuclear marker Hoechst (Figs. 153 S1C or S5B) or CFP+, GFP+, YFP+, or RFP+ expression (Fig. S4B). For consistency, 154 segmentation conditions were applied as a batch using identical parameters. Segmented cells were 155 randomly colored and manually inspected to assess the quality of segmentation based on Hoechst 156 or FP staining. For Fig. S5B, segmentation on Hoechst+ cells provided accurate identification of 157 round, regular shaped, lymphocytes but, as expected, struggled with morphologically complex 158 stromal and structural cell segmentation. Nevertheless, this approach provided a relative 159 quantification for the number of surfaces positive for each individual marker. Importantly, these qualitative measures were in agreement with the amount of signal present by visual inspection. 160 161 Surfaces were scored positive for a given marker based on absolute intensity values generated in 162 Imaris and these values were applied uniformly to serial and IBEX-generated images to quantify 163 marker-positive surfaces (Fig. S5B).

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histoCAT analysis of IBEX-generated images

165 Using the workflow described previously (8), IBEX-generated images were segmented 166 using Ilastik (Version 1.3.3) (9) and CellProfiler (Version 3.1.9) (10). Briefly, B220, CD3, and CD45 167 (most of the immune cells) markers were merged into a single membrane and JOJO-1 was used as a nuclear marker. Membrane and nuclear Images were loaded into Ilastik for supervised training 168 169 of pixel segmentation. The trained llastik model classified image pixels into 3 classes (membrane, 170 nuclear, and background) and generated probability maps for CellProfiler for cellular objects 171 segmentation. CellProfiller-generated object/mask images were exported as TIFF files for 172 downstream analysis. To quantify marker expression, individual marker images, along with masks, 173 were loaded into histoCAT and Phenograph clustering with default parameters was performed. 174 Phenograph consistently identified 29 different clusters/phenotypes (Fig. 4B). Hierarchical 175 clustering and heatmap (Fig. S6B) were generated from the Phenograph output using the Seaborn

176 Python package. The high-dimensional single-cell data was projected onto 2 dimensions using the 177 t-SNE module in histoCAT for visualization purposes shown in Fig. 4B. Cellular populations 178 identified in Fig. 4C were manually phenotyped based on the cellular markers expressed within 179 each Phenograph cluster and summarized in the heat map found in Fig. S6B: CD4⁺ T (cluster 3), 180 CD8⁺ T (cluster 2), Tfh (cluster 19), Tregs (cluster 18), naïve B (cluster 5), MHCII^{Hi} B (cluster 13), 181 PCs (cluster 10), GC B (cluster 6), CD68⁺ macrophages (cluster 15), CD206⁺ macrophages (cluster 182 11), SCS macrophages (cluster 17), MS macrophages (cluster 25), pan DC (cluster 4), cDC1/dDCs 183 (cluster 22), and cDC2s (cluster 26). A median of ratios method (11) was performed for the cell 184 count normalization (Fig. S6D). All histoCAT analysis was done on an iMAC, (Retina 5K, 27-inch, 185 2017, 4.2Ghz i7, 64G RAM) with macOS Mojave operating system.

186 Multi-plex Opal IHC on heavily fixed tissues

187 pLNs were collected from naïve mice and fixed at 4° C for 8 days in 10% neutral-buffered 188 formalin (Cancer Diagnostics). Following fixation, samples were washed in PBS to remove formalin 189 and embedded in paraffin blocks. 5 µm sections were cut from paraffin blocks, mounted on slides, 190 and left at 37°C for 16 hours to dry. Slides were incubated at 60°C for 45 minutes, dewaxed using 191 a standard protocol of 10 minutes in xylene (2 times), and rehydrated with graded concentrations 192 of ethanol and water (100% ethanol for 10 minutes, 95% ethanol for 10 minutes, 70% ethanol for 193 5 minutes). Following dewaxing, slides were first rinsed in water and tissues were fixed to the slides 194 by placing in 10% formalin for 15 minutes, rinsed in water, and then placed in TBST (1X TBS + 195 0.5% Tween20 (Thermo)) to prevent drying out. Antigen retrieval was performed using the AR6 196 buffer from the Opal Multi-plex kit (Akoya Biosciences) using a conventional microwave at 100% 197 power for 45 seconds followed by 10% power for 15 minutes. To perform multi-plex Opal IHC, 198 tissue sections were first blocked in Opal antibody diluent/blocking buffer (Akoya Biosciences) for 199 10 minutes. Next, unconjugated primary antibodies were added to the tissue for 12-16 hours at 4° 200 C, 4 hours at room temperature, or 30 minutes using the PELCO BioWave Pro 36500-230 201 microwave described under 'IBEX using inverted confocal microscope' section of the Material and 202 Methods. After primary antibody incubation, samples were washed with TBST and an Opal anti-203 rabbit HRP-conjugated secondary antibody was added at a dilution of 1:5 for 10 minutes at room 204 temperature or with a species matched HRP-conjugated secondary antibody for 1 hour at room 205 temperature. Samples were washed several (5-6) times with TBST and Opal dyes (diluted 1:100 206 in 1X amplification buffer (Akoya Biosciences)) were added to the sample and incubated for 10 207 minutes at room temperature. Following this last step, samples were washed with TBST and 208 antigen retrieval/antibody stripping was performed using the AR6 buffer and conventional 209 microwave treatment described above. This series of steps-primary antibody incubation, 210 incubation with HRP-conjugated secondary, labeling with Opal dye, antibody stripping-was 211 repeated for the following Opal fluorophores to achieve 6-plex imaging: Opal 520, 540, 570, 620, 650, and 690. Slides were mounted and imaged with an upright Leica TCS SP8 X confocal microscope as described above. After representative images were captured, coverslips were removed and tissue sections were treated with 1 mg/mL of LiBH₄ prepared in diH₂O for 30 minutes to bleach the Opal 570, 650, and 690 dyes. Cycles of multi-plex Opal IHC and IBEX were repeated to achieve the desired number of markers. Individual images were aligned and processed as described above. A complete list of antibodies and reagents can be found in Table S4.

218 Chemokine staining for endogenous CXCL9

219 Livers were collected from naïve WT and Cxcl9^{-/-} animals, perfused with 1% PFA through the portal vein, and fixed at 4° C for 12-16 hours in BD CytoFix/CytoPerm diluted 1:4 in PBS. 220 221 Samples were incubated in sucrose for 24 hours at 4° C and frozen in OCT. To visualize 222 endogenous CXCL9 levels, 20 µm sections were rehydrated in PBS, incubated in 0.1% H₂O₂ for 223 30 minutes to quench endogenous peroxidase, and incubated with unconjugated (CXCL9) and 224 fluorescently-conjugated primary antibodies. Tissue sections were washed to remove unbound 225 antibodies, fixed with 10% formalin to cross-link the bound antibodies to the tissue, washed with 226 TBST, and an Opal anti-rabbit HRP-conjugated secondary antibody was added at a dilution of 1:5 227 for 10 minutes at room temperature. Following secondary antibody incubation, samples were 228 washed in TBST and the Opal 650 dye (diluted 1:100 in 1X amplification buffer) was added for 5 229 minutes. Slides were washed, mounted in Fluoromount G, and imaged as described above. A 230 complete list of antibodies and reagents can be found in Table S4.

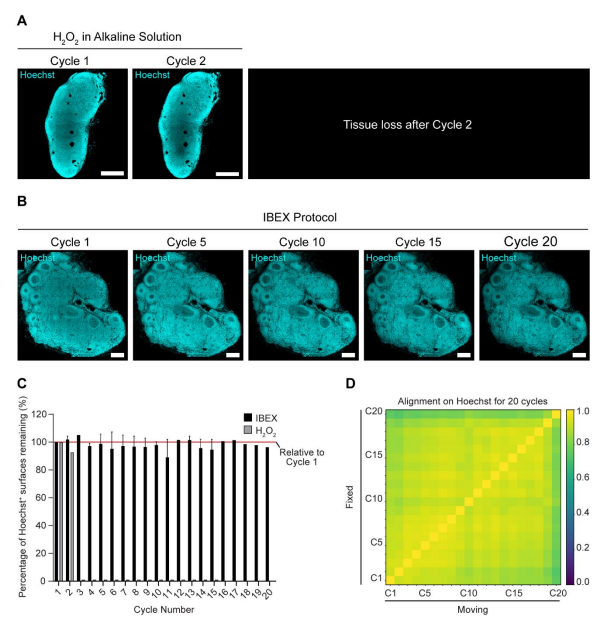
231 Incorporation of TotalSeqA[™] antibodies into IBEX workflow

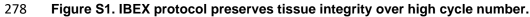
232 LNs from C57BL/6 mice were harvested and processed as described above. Integrating 233 published methods, slides containing 30 µm sections were blocked for 30 minutes at room 234 temperature in Buffer A (PBS supplemented with 5% donkey serum (Jackson Immunoresearch), 235 200 µg/ml sheared salmon sperm (Thermo), 0.2% Triton X-100 (Sigma) and 5 µg/ml Fc-block 236 (Thermo)). Immunolabeling was conducted using Buffer A with the addition of 5 mM EDTA and 237 0.02% Dextran sulfate, TotalSeqA[™] antibodies (5 µg/ml), and fluorophore-conjugated antibodies 238 at the indicated concentrations (Table S1) (12). All primary antibody incubations were performed 239 using the PELCO BioWave Pro 36500-230 microwave as described in the "IBEX using inverted 240 confocal microscope' section of the Materials and Methods. Following antibody incubation, slides 241 were washed using PBS with 0.1% Triton X-100 and post-fixed with 5 mM of BS(PEG)5 (Thermo) 242 for 10 minutes before quenching with 100 mM Ammonium Chloride (Sigma). After washing with 243 PBS, Fluoromount G was applied and fluorescently-conjugated antibodies were imaged in the first 244 cycle. Mounting medium was removed by washing with PBS before bleaching with LiBH₄ for 15 minutes. For in situ visualization of TotalSeqA[™] antibodies, oligonucleotides were designed 245

complementary to the TotalSeqA[™] barcode. These complementary oligonucleotides were synthesized by Integrated DNA Technologies (IDT) and 5'-end conjugated with indicated fluorophores before HPLC purification (Table S5). Direct labeling of TotalSeqA™ antibodies with complementary fluorescent oligonucleotides was achieved by adding 1 µM concentrations of imager oligonucleotides in PBS with 0.1% Triton X-100 and incubating for 30 minutes at room temperature. Tissue sections were washed with 0.5X PBS and 0.1% Triton X-100, mounted, and imaged (Cycle 2 of IBEX). After removal of Fluoromount G, oligonucleotides were dehybridized for 15 minutes by incubation with a 0.1X PBS solution containing 30% Formamide. Following 5-6 exchanges of PBS, the next set of oligonucleotides (Cycle 3) was applied as before.

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276 Figures S1-S7 and Legends





279 (A) Confocal images of mouse inguinal LN iteratively imaged using H_2O_2 in alkaline solution. 280 Tissues lifted after 2 cycles and could not be imaged as a comparison. Scale bar corresponds to 281 500 µm. (B) Confocal images of human mesenteric LN iteratively imaged using IBEX protocol for 20 cycles. Scale bar corresponds to 500 µm. (C) Quantification of percentage of Hoechst+ surfaces 282 283 remaining per cycle relative to the number of surfaces present in cycle 1. Red line is reflective of 284 no tissue loss. Data are pooled from 2 similar experiments: a 15 cycle mouse inguinal LN and a 20 cycle human mesenteric LN. Shown is the mean ± SEM. (D) Cross correlation similarity matrix after 285 alignment with Hoechst across 20 cycle experiment shown in B. 286

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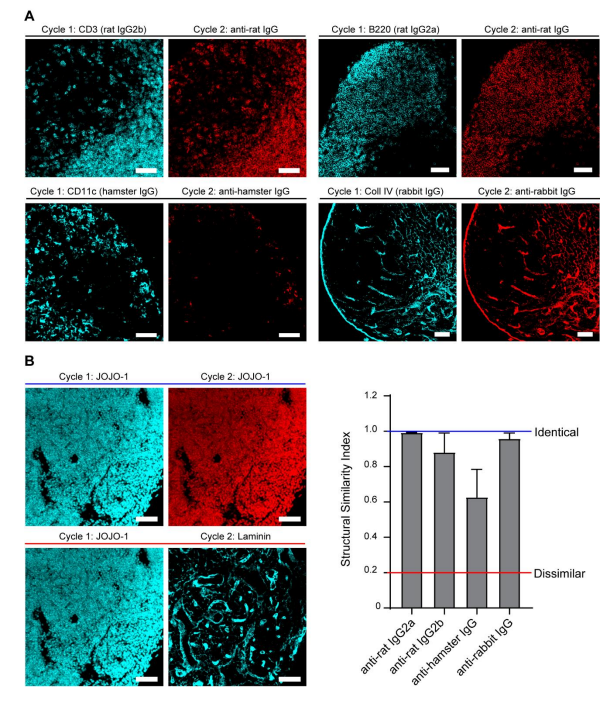
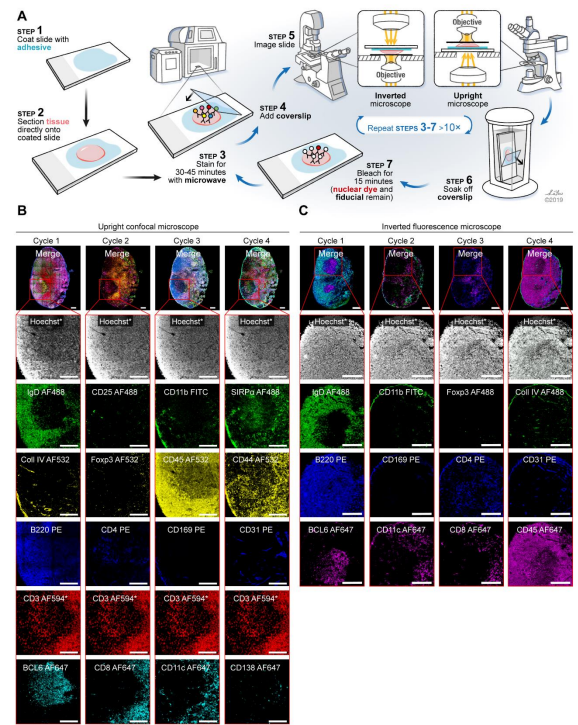




Figure S2. LiBH₄ primarily acts by bleaching fluorophores and not stripping antibodies.

290 Popliteal LN sections were stained with primary antibodies directed against various antigens, 291 bleached with LiBH₄, and then labeled with secondary antibodies directed against the primary 292 antibody isotype. (A) Confocal images of murine LN sections depicting the level of signal present 293 with the primary antibodies as compared to an appropriate secondary antibody after IBEX protocol. 294 (B) Bar graph summarizing the level of similarity between images captured before and after LiBH₄ 295 treatment. Red line represents dissimilar images (Cycle 1 JOJO-1 versus Cycle 1 Laminin) and 296 blue line denotes highly similar images (Cycle 1 JOJO-1 vs Cycle 2 JOJO-1). Data representative 297 of 5 similar experiments. Shown is the mean ±SEM. Scale bar represents 50 µm.



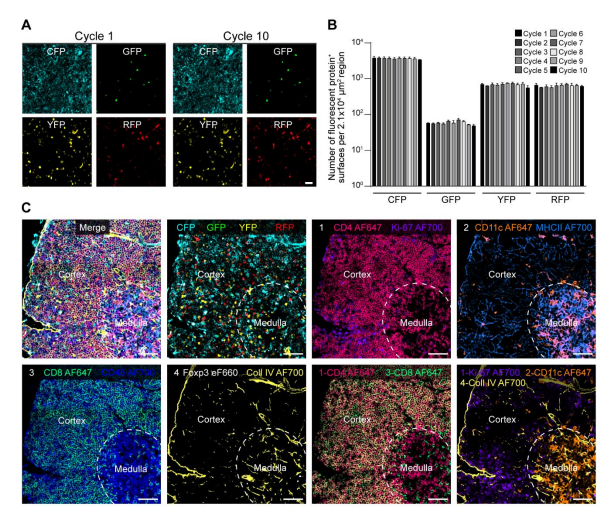
298 *Fiducials: Hoechst and CD3 AF594, scale bar 150 um

299 Figure S3. IBEX can be easily adapted to other microscope configurations.

(A) Schematic depicting IBEX protocol using an inverted or upright microscope. (B) Confocal
 images of popliteal LNs from SRBC-immunized mice. 30 µm tissue sections were labeled with 4
 separate 6 parameter imaging panels. The nuclear dye Hoechst and membrane label CD3 AF594

303 were present throughout cycles 1-4 and served as fiducials. Top panels are a merge of all channels

for each cycle except for Hoechst. Scale bar represents 150 μm. Confocal images were acquired
 by an upright confocal microscope. (C) 5 μm sections from popliteal LNs from SRBC-immunized
 mice were visualized using an inverted epifluorescence microscope. Tissue sections were labeled
 with 4 separate 4 parameter imaging panels with Hoechst serving as a fiducial for cycles 1-4. Scale
 bar represents 150 μm.



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311 Figure S4. Fluorescent proteins are not bleached with IBEX protocol.

Fluorescent Confetti animals were injected i.p. with tamoxifen on day 0 and day 2 to induce the 312 expression of the following fluorescent proteins: membrane CFP, nuclear GFP, and cytoplasmic 313 314 YFP and RFP. On day 4, tissues were collected and processed for confocal microscopy. (A) 315 Example images of signal from fluorescent proteins at Cycle 1 (no LiBH₄) and Cycle 10 (after 9 316 rounds of LiBH₄ bleaching). (B) Quantification of the number of fluorescent protein+ surfaces per 317 2.1x10⁴ μ m² region of interest from 2 independent experiments. Shown is the mean ± SEM. (C) 318 Four cycles of IBEX were applied to 30 µm sections of thymus tissue. AF647 and AF700 conjugated 319 antibodies were used to stain immune and structural markers in the tissue. Representative images 320 showing the compatibility of IBEX with transgenic animals expressing fluorescent proteins. Data 321 are representative of 2 independent experiments. For A and C, images represent a single z slice 322 and scale bar corresponds to 25 μ m (A) or 50 μ m (B).

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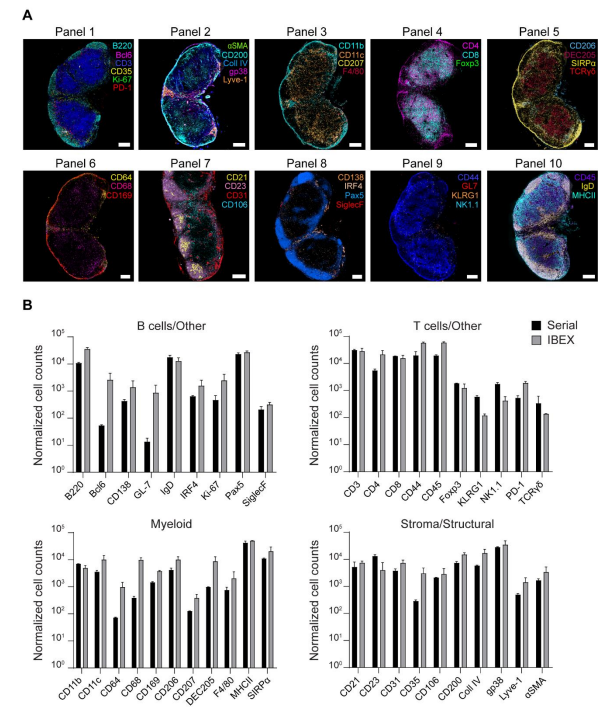
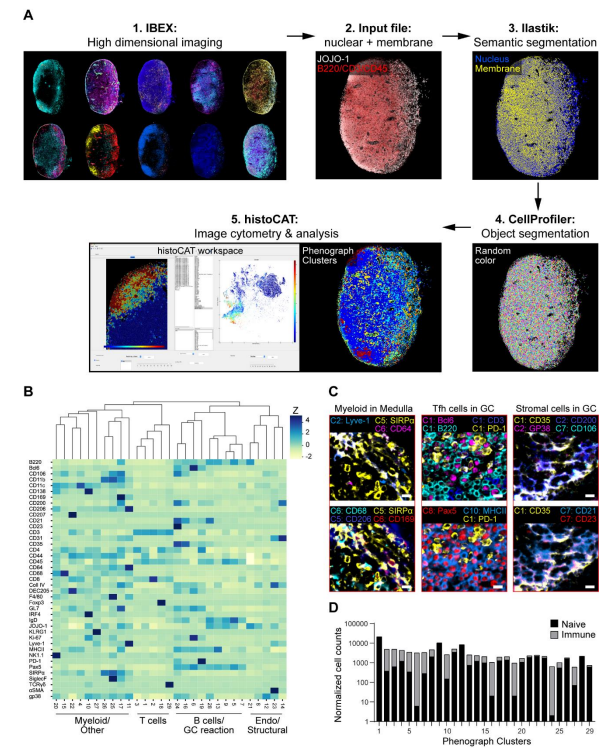




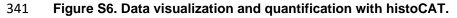
Figure S5. Comparable staining observed by serial and iterative immunofluorescence methods.

(A) Confocal images of inguinal LNs (iLNs) from SRBC-immunized mice. Serial sections were
 stained with the same panels used for the 10 cycle 41 parameter IBEX experiments described in
 Fig. 4A. Scale bar is 200 µm. (B) Cells were segmented on the nuclear marker JOJO-1 and the
 number of surfaces positive for each marker were quantified from images acquired serially as in A
 or iteratively via IBEX method. Data are from 2-3 LNs per group with 2 immunized iLNs for serial

- method and 1 naïve pLN, 1 immunized pLN, and 1 immunized axillary LN (aLN) for IBEX method. Shown is the mean \pm SEM. See Movie S7. 336







(A) histoCAT workflow for image analysis. Step 1: High dimensional imaging of mouse LNs using
 IBEX method. Step 2: Nuclei were defined by JOJO-1 and membranes were generated by
 combining B220, CD3, and CD45 into one composite channel. Step 3: Semantic segmentation was
 performed using Ilastik. Step 4: Objects were further segmented using CellProfiler. Each

346 segmented cell is randomly colored and projected back onto original x-y coordinates. Step 5: Data 347 visualization and analysis with histoCAT. (B) Marker expression heatmap for the phenotypes 348 identified by PhenoGraph clustering in histoCAT using segmented cells from naïve (n = 32,091) 349 and SRBC-immunized mouse LNs (n = 80,355). The heatmap displays relative expression levels based on Z-score normalized marker intensity values, and single cells are hierarchically clustered 350 351 within each phenotype group. Numbers at the bottom of the heatmap indicate corresponding 352 Phenograph cluster IDs with labels corresponding to manually assigned cell populations. 353 Endothelial (Endo). (C) Example images of myeloid, Tfh, and stromal cell populations identified by 354 histoCAT Phenograph clustering from an immunized mouse LN. Scale bar corresponds to 10 µm. 355 (D) Normalized cell counts for Phenograph clusters obtained from naïve and SRBC-immunized 356 mouse LNs. Data are from one experiment and are representative of 2 similar experiments.

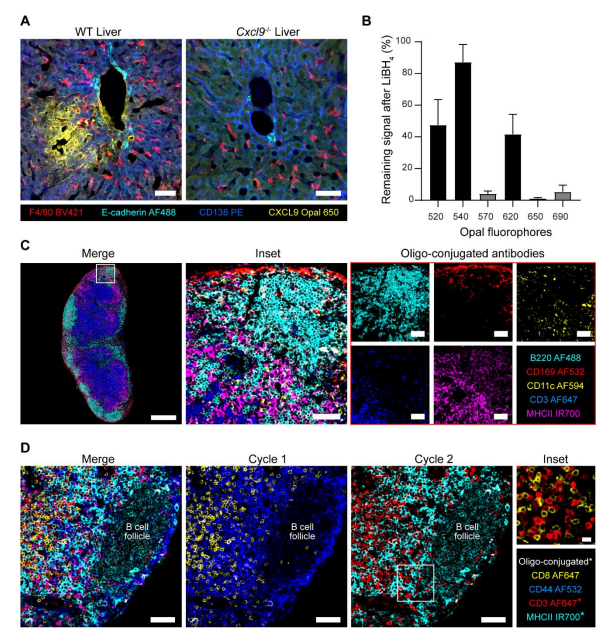


Figure S7. Extensions of IBEX workflow to enable single cell resolution of endogenous chemokines and immune populations using Opal fluorophores and oligonucleotideconjugated antibodies.

(A) Detection of endogenous CXCL9 levels in fixed frozen mouse liver sections stained with 361 indicated markers. Scale bars 50 µm. Confocal images are representative of 3 similar experiments. 362 363 (B) Percentage of fluorophore signal remaining after 30 minutes of LiBH₄ treatment. Data are 364 pooled from 4 experiments with iterative staining of the multiple Opal fluorophores. Shown is the 365 mean ± SEM. (C) Confocal images of mouse inguinal LN labeled with 5 different oligo-conjugated 366 antibodies and complementary fluorescent imager strands. Scale bars (left-most panel 400 µm or 367 100 µm). (D) Representative images from a 2 cycle IBEX experiment performed on inguinal mouse 368 LNs where the first cycle consisted of fluorophore-conjugated antibodies and the second cycle 369 consisted of oligo-conjugated antibodies denoted by the asterisks. Scale bars (50 µm, Inset 10 370 μm).

371 Tables S1-S5

Table S1. Time and method used to bleach fluorescently conjugated antibodies and dyes.

							Time to (Minutes)	
Marker	Clone	Conjugate	Vendor	Cat No.	Isotype	Dilution	LiBH₄	LiBH₄ - Light
β -3 Tubulin	AA10	BV421	BioLegend	657412	Mouse IgG2a, к	1:200	-	<15
β-3 Tubulin	AA10	AF532	BioLegend	Custom	lgG2a, к IgG2a, к	1:100	<15	-
B220	RA3-6B2	BV421	BioLegend	103251	Rat IgG2a, ĸ	1:400	-	<15
B220	RA3-6B2	BV510	BioLegend	103248	Rat IgG2a, ĸ	1:300	-	<15
B220	RA3-6B2	AF488	BD Biosciences	557669	Rat IgG2a, ĸ	1:400	<15	-
B220	RA3-6B2	AF532	Thermo	58-0452-82	Rat IgG2a, ĸ	1:50	<15	-
B220	RA3-6B2	PE	BD Biosciences	553090	Rat IgG2a, ĸ	1:400	<15	-
B220	RA3-6B2	eF570	Thermo	41-0452-80	Rat IgG2a, ĸ	1:200	<15	-
B220	RA3-6B2	eF615	Thermo	42-0452-82	Rat IgG2a, ĸ	1:200	>120	-
B220	RA3-6B2	AF647	BioLegend	103226	Rat IgG2a, ĸ	1:400	<15	-
B220	RA3-6B2	AF700	BioLegend	103232	Rat IgG2a, ĸ	1:50	<15	-
BCL2	100	AF647	BioLegend	658705	Mouse IgG ₁	1:25	<15	-
Bcl6	K112-91	AF647	BD Biosciences	561525	Mouse IgG ₁ , K	1:50	<15	-
CD1c	L161	PE	BioLegend	331506	Mouse IgG ₁ ,	1:50	<15	-
CD1d	1B1	PE	BD Biosciences	553846	к Rat IgG2b, к	1:100	<15	-
CD3	17A2	BV421	BioLegend	100228	Rat IgG2b, K	1:400	<15	- <15
CD3	17A2	BV510	BioLegend	100234	Rat IgG2b, K	1:50	-	<15
CD3	17A2	AF488	BioLegend	100210	Rat IgG2b, K	1:200	<15	-
CD3	17A2	AF532	Thermo	58-0032-80	Rat IgG2b, ĸ	1:50	<15	-
CD3	17A2	PE	BioLegend	100205	Rat IgG2b, ĸ	1:200	<15	-
CD3	17A2	AF594	BioLegend	100240	Rat IgG2b, ĸ	1:400	>120	-
CD3	17A2	AF647	BD Biosciences	557869	Rat IgG2b, K	1:400	<15	-
CD3	UCHT1	AF532	Thermo	58-0038-42	Mouse IgG ₁ ,	1:50	<15	-
CD3	UCHT1	AF594	BioLegend	300446	Mouse IgG ₁	1:200	>120	-
CD4	GK1.5	BV421	BioLegend	100443	Rat IgG2b, ĸ	1:200	-	<15
CD4	GK1.5	BV510	BioLegend	100449	Rat IgG2b, к	1:50	-	<15
CD4	GK1.5	PE	BD Biosciences	553730	Rat IgG2b, ĸ	1:200	<15	-
CD4	GK1.5	AF594	BioLegend	100446	Rat IgG2b, ĸ	1:200	>120	-
CD4	RM4-5	AF488	BD Biosciences	557667	Rat IgG2a, ĸ	1:100	<15	-
CD4	RM4-5	AF532	Thermo	58-0042-80	Rat IgG2a, ĸ	1:50	<15	-
CD4	RM4-5	eF570	Thermo	41-0042-82	Rat IgG2a, κ	1:100	<15	-
CD4	RPA-T4	AF532	Thermo	58-0049-42	Mouse IgG ₁ , K	1:25	<15	-
CD4	RPA-T4	AF700	BioLegend	300526	Mouse IgG ₁ ,	1:25	<15	-
CD8	53-6.7	BV421	BioLegend	100738	Rat IgG2a, ĸ	1:200	-	<15
CD8	53-6.7	BV510	BioLegend	100752	Rat IgG2a, ĸ	1:200	-	<15
CD8	53-6.7	AF488	BioLegend	100723	Rat IgG2a, ĸ	1:200	<15	-
CD8	53-6.7	PE	BD Biosciences	553032	Rat IgG2a, ĸ	1:400	<15	-
CD8	53-6.7	AF594	BioLegend	100758	Rat IgG2a, ĸ	1:200	>120	-
CD8	53-6.7	AF647	BioLegend	100724	Rat IgG2a, ĸ	1:200	<15	-
CD8	SK1	AF488	BioLegend	344716	Mouse IgG ₁ , κ	1:50	<15	-
CD10	FR4D11	PE	Caprico Biotechnologies	103926	Mouse IgG ₁ ,	1:50	<15	-
CD11b	5C6	FITC	Bio-Rad	MCA711F	Rat IgG2b	1:100	<15	-
CD11b	5C6	PE	Bio-Rad	MCA711PE	Rat IgG2b	1:100	<15	-
CD11b	M1/70	AF488	BioLegend	101217	Rat IgG2b, ĸ	1:100	<15	-
CD11c	N418	AF488	Thermo	MCD11c20	Hamster IgG	1:50	<15	-
CD11c	N418	AF594	BioLegend	117346	Hamster IgG	1:50	>120	-
CD11c	N418	AF647	BioLegend	117312	Hamster IgG	1:100	<15	-
CD11c	B-Ly6	AF700	BD Biosciences	561352	Mouse IgG ₁ ,	1:25	<15	-
CD20	L26	AF488	Thermo	53-0202-82	Mouse IgG2b, κ	1:200	<15	-
CD21	7E9	Pacific Blue	BioLegend	123413	Rat IgG2a, ĸ	1:200	<15	-
CD21	Bu32	AF532	BioLegend	Custom	Mouse IgG ₁ ,	1:400	<15	-
CD23	B3B4	AF647	BioLegend	101611	⊼ Rat IgG2a, к	1:50	<15	-
CD23	EBVCS-5	AF532	BioLegend	NA, Custom	Mouse IgG ₁ ,	1:25	<15	-
CD25	PC61.5	AF488	Thermo	53-0251-82	Rat IgG1, λ	1:50	<15	-
CD25	M-A251	AF647	BioLegend	356127	Mouse IgG ₁ ,	1:50	<15	-
0020	101-7231	AI 047	DioLegeniu	330127	wouse igo1,	1.50	<13	l -

CD31	MEC13.3	AF488	BioLegend	102514	Rat IgG2a, к	1:100	<15	-
CD31	MEC13.3	PE	BD Biosciences	553373	Rat IgG2a, ĸ	1:200	<15	-
CD31	MEC13.3	AF594	BioLegend	102520	Rat IgG2a, к	1:100	>120	-
CD31	MEC13.3	AF647	BioLegend	102516	Rat IgG2a, к	1:100	<15	-
CD31	WM59	AF700	BioLegend	303133	Mouse IgG ₁ , κ	1:25	<15	-
CD34	QBEND/10	PE	Thermo	MAI-10205	Mouse IgG ₁	1:50	<15	-
CD35	8C12	BV510	BD Biosciences	740132	Rat IgG2a, к	1:600	-	<15
CD35	E11	PE	BioLegend	333406	Mouse IgG₁, к	1:800	<15	-
CD38	HIT2	AF700	BioLegend	303524	Mouse IgG ₁ , κ	1:25	<15	-
CD39	A1	PE	BioLegend	328208	Mouse IgG ₁ , K	1:50	<15	-
CD44	IM7	AF488	BioLegend	103016	Rat IgG2b, ĸ	1:100	<15	-
CD44	IM7	AF532	BioLegend	Custom	Rat IgG2b, ĸ	1:100	<15	-
CD44	IM7	AF647	BioLegend	103018	Rat IgG2b, ĸ	1:100	<15	-
CD44	IM7	AF700	BioLegend	103026	Rat IgG2b, κ	1:50	<15	-
CD45	30-F11	BV421	BioLegend	103134	Rat IgG2b, κ	1:100	-	<15
CD45	30-F11	BV510	BioLegend	103138	Rat IgG2b, ĸ	1:100	-	<15
CD45	30-F11	AF488	BioLegend	103122	Rat IgG2b, κ	1:200	<15	-
CD45	30-F11	AF532	Thermo	58-0451-82	Rat IgG2b, κ	1:200	<15	-
CD45	30-F11	AF647	BioLegend	103124	Rat IgG2b, κ	1:400	<15	-
CD45	30-F11	AF700	BioLegend	103128	Rat IgG2b, к	1:50	<15	-
CD45	HI30	AF532	Thermo	58-0459-41	Mouse IgG ₁ , к	1:50	<15	-
CD45	F10-89-4	PE/iFluor594	Caprico Biotechnologies	1016185	Mouse IgG2a, к	1:50	<15	-
CD49a	TS2/7	PE	BioLegend	328304	Mouse IgG1,	1:50	<15	-
CD54	HA58	AF647	BioLegend	353114	к Mouse IgG ₁ ,	1:50	<15	-
CD64	X54-5/7.1	AF647	BioLegend	139322	к Mouse IgG ₁ ,	1:50	<15	-
CDeeb	C10E5	AE647	Dial agend	205110	K Mauga JaMa ::	1.05	.4 5	
CD66b CD68	G10F5 FA-11	AF647 BV421	BioLegend BioLegend	305110 137017	Mouse IgM, κ Rat IgG2a	1:25 1:200	<15	- <15
CD68 CD68	FA-11 FA-11	AF488	BioLegend	137017	Rat IgG2a Rat IgG2a	1:200	- <15	<15
CD68	KP1	AF400 AF647	Santa Cruz	sc-200060	Mouse IgG ₁ ,	1:100	<15	-
CD69	+	None	R&D	AF2386	к Goat IgG	1:50	-	-
CD69	- H1.2F3	PE	BioLegend	104508	Hamster IgG	1:50	- <15	1.
CD69 CD69	FN50	AF647	BioLegend	310918	Mouse IgG1,	1:100	<15	-
CD94	DX22	PE	BioLegend	305506	к Mouse IgG ₁ ,	1:50	<15	-
CD103	2E7	AF488	BioLegend	121407	к Hamster IgG	1:100	<15	-
	429							-
CD106		AF488	BioLegend	105710	Rat IgG2a, ĸ	1:50	<15	
CD106 CD106	429 STA	eF660 PE	Thermo BioLegend	50-1061-80 305806	Rat IgG2a, κ Mouse IgG ₁ ,	1:50 1:50	<15 <15	-
			°,		к		<15	
CD117 CD117	2B8 104D2	BV421 AF488	BioLegend BioLegend	105827 313234	Rat IgG2b, κ Mouse IgG ₁ ,	1:50 1:50	- <15	<15 -
CD129	201.2	D\/424	PD Pionsisses	562610	к Rat lgG2a, к	1.50	+	~1E
CD138 CD138	281-2 281-2	BV421 AF647	BD Biosciences BioLegend	562610 142526	Rat IgG2a, K Rat IgG2a, K	1:50 1:50	- <15	<15
CD138 CD138	281-2 MI15	PE	BioLegend	356504		1:200	<15	-
					Mouse IgG ₁ , K			
CD138	MI15	AF647	BioLegend	356523	Mouse IgG ₁ , K	1:200	<15	-
CD163	GH1/61	AF647	BioLegend	333620	Mouse IgG ₁ , κ	1:100	<15	-
CD166	EPR2759	AF488	AbCAM	Ab197543	Rabbit mAb	1:100	<15	-
CD169	3D6.112	FITC	Bio-Rad	MCA884F	Rat IgG2a, ĸ	1:50	<15	-
CD169	3D6.112	PE	Biolegend	142404	Rat IgG2a, ĸ	1:300	<15	-
CD169	3D6.112	AF594	BioLegend	142416	Rat IgG2a, ĸ	1:100	>120	-
CD200	Ox-90	BV421	BD Biosciences	565547	Rat IgG2a, ĸ	1:100	-	<15
CD206	C068C2	BV421	BioLegend	141717	Rat IgG2a, ĸ	1:100	-	<15
CD207 Clec9a	eBioL31 8F9	PE AF488	Thermo BioLegend	12-2075-80 Custom	Rat IgG2a Mouse	1:50 1:25	<15 <15	-
Collagen IV	-	None	AbCAM	19808	IgG2a, к Rabbit IgG	1:200	-	-
Collagen IV Collagen IV	-	None	AbCAM	Ab6586	Rabbit IgG	1:200	+	-
CXCL12	79018	AF532	R&D	MAB350-500	Mouse IgG ₁	1:200	<15	-
CXCL13	-	AF532	R&D	(Unconjugated) AF801	Goat IgG	1:25	<15	-
01/05/	004067	150/5		(Unconjugated)	D. (). CT	1.55		
CXCR6 Cytokeratin	221002 C-11	AF647 AF647	Novus BioLegend	FAB2145R 628604	Rat IgG2b Mouse IgG1,	1:50 1:200	<15 <15	-
Cytokeratin	AE1/AE3	eF660	Thermo	50-9003-82	к Mouse IgG ₁	1:100	<15	-
	I AEI/AES	eroou	THEITIO	30-3003-02	wouse igo1	1.100	<10	-
DCAMKL1		None	AbCAM	Ab37994	Rabbit IgG	1:50		-

DC-SIGN	9E9A8	AF647	BioLegend	330112	Mouse	1:50	<15	-
DEC205	NLDC-145	AF647	BioLegend	138204	lgG2a, к Rat lgG2a, к	1:50	<15	-
Desmin	-	None	AbCAM	Ab15200	Rabbit IgG	1:200	-	-
Desmin	Y66	AF488	AbCAM	Ab185033	Rabbit IgG	1:200	<15	-
E-cadherin	DECMA-1	AF647	BioLegend	147308	Rat IgG1, K	1:100	<15	-
EpCAM	G8.8	BV510	BD Biosciences	563216	Rat IgG2a, ĸ	1:100	-	<15
EpCAM	G8.8	AF594	BioLegend	118222	Rat IgG2a, ĸ	1:400	>120	-
EpCAM	G8.8	AF647	BioLegend	118212	Rat IgG2a, ĸ	1:200	<15	-
EpCAM	9C4	AF594	BioLegend	324228	Mouse IgG2b, к	1:500	>120	-
EpCAM	9C4	AF647	BioLegend	324212	Mouse IgG2b, к	1:100	<15	-
F4/80	BM8	BV421	BioLegend	123132	Rat IgG2a, к	1:50	-	<15
F4/80	BM8	PE	Thermo	12-4801-82	Rat IgG2a, ĸ	1:100	<15	-
F4/80	BM8	AF647	BioLegend	123122	Rat IgG2a, к	1:50	<15	-
F4/80	BM8	AF700	BioLegend	123130	Rat IgG2a, к	1:50	<15	-
Fibronectin	2F4	AF532	Novus	NBP2- 22113AF532	Mouse IgG ₁	1:25	<15	-
Foxp3	FJK-16s	AF488	Thermo	53-5773-82	Rat IgG2a, к	1:50	<15	-
Foxp3	FJK-16s	AF532	Thermo	58-5773-82	Rat IgG2a, к	1:50	<15	-
Foxp3	FJK-16s	PE	Thermo	12-5773-82	Rat IgG2a, к	1:50	<15	-
Foxp3	FJK-16s	eF570	Thermo	41-5773-82	Rat IgG2a, ĸ	1:50	<15	-
Foxp3	FJK-16s	eF660	Thermo	50-5773-82	Rat IgG2a, к	1:50	<15	-
FOXP3	236A/E7	eF570	Thermo	41-4777-82	Mouse IgG₁, κ	1:50	<15	-
GL-7	GL7	PE	BD Biosciences	561530	Rat IgM, к	1:100	<15	-
Glutamine synthetase	-	None	AbCAM	Ab49873	Rabbit IgG	1:200	-	-
gp38	8.1.1	AF488	BioLegend	127405	Hamster IgG	1:50	<15	-
HLA-DR	L243	AF488	BioLegend	307619	Mouse IgG2a, к	1:200	<15	-
Hoechst	-	-	Biotium	40046	-	1:5,000	-	-
ICOS	CS98.4A	AF488	BioLegend	313514	Hamster IgG	1:25	<15	-
lgA	-	AF555	Southern Biotech	1040-32	Goat IgG	1:500	<15	-
gA1	B3506B4	AF647	SouthernBiotech	9130-31	Mouse IgG ₁ , κ	1:500	<15	-
lgA2	A9604D2	AF488	SouthernBiotech	9140-30	Mouse IgG1, к	1:500	<15	-
lgD	11-26c.2a	AF488	BioLegend	405718	Rat IgG2a, к	1:400	<15	-
lgD	11-26c.2a	AF594	BioLegend	405740	Rat IgG2a, к	1:400	>120	-
lgD	11-26c.2a	AF700	BioLegend	405729	Rat IgG2a, к	1:50	<15	-
lgD	IA6-2	AF488	BioLegend	348216	Mouse IgG2a, к	1:25	<15	-
IgM	EPR5539- 65-4	AF647	AbCAM	Ab200629	Rabbit mAb	1:100	<15	-
IRF4	3E4	FITC	Thermo	11-9858-82	Rat IgG1, ĸ	1:50	<15	-
IRF4	3E4	PE	Thermo	12-9858-82	Rat IgG1, ĸ	1:50	<15	-
JOJO-1	-	-	Thermo	J11372	-	1:10,000	-	-
Keratin 14	Poly9060	-	BioLegend	906004	Chicken IgY	1:50	-	-
Keratin 18 Ki-67	1G11C4 B56	CoraLite488 AF488	ProteinTech BD Biosciences	CL488-66187 558616	Mouse IgG ₁ Mouse IgG ₁ ,	1:50 1:50	<15 <15	-
Ki-67	B56	AF700	BD Biosciences	561277	κ Mouse IgG ₁ ,	1:50	<15	-
KLRG1	2F1	AF488	BD Biosciences	561619	к Hamster	1:50	<15	-
					lgG₂, κ			
Laminin 1 + 2 Lumican	-	None AF532	AbCAM R&D	Ab7463 AF2846	Rabbit IgG Goat IgG	1:100 1:50	- <15	-
1 1 60	149	ΛE400	Piol occord	(Unconjugated)	Pot laCO	1.50	×1E	
Ly-6G	1A8	AF488	BioLegend AbCAM	127626	Rat IgG2a, ĸ	1:50	<15	-
Lysozyme	-	-		Ab2408	Rabbit IgG	1:50	-	-
Lyve-1 Lyve-1	ALY7 -	eF570 AF532	Thermo R&D	41-0443-82 AF2089 (Upgeniugated)	Rat IgG1, κ Goat IgG	1:100 1:100	<15 <15	-
MARCO	t <u>.</u>	t _	Thermo	(Unconjugated) PA5-64134	Rabbit IgG	1:25	-	+.
MARCO MHC-II	- M5/114.15.2	- BV421	BioLegend	107631	Rat IgG2b, K	1:400	-	- <15
MHC-II	M5/114.15.2	AF647	BioLegend	107618	Rat IgG2b, K	1:600	<15	-
MHC-II	M5/114.15.2	AF700	BioLegend	107622	Rat IgG2b, ĸ	1:100	<15	-
NF-H/NF-M	SM1-35	AF488	BioLegend	835614	Mouse IgG ₁ ,	1:50	<15	-
NK1.1	PK136	BV421	BioLegend	108731	моиse IgG1, к Mouse	1:50	-	<15
					lgG2a, к			-
p53	PAb 240	PE	Novus	NB200-103PE	Mouse IgG ₁ , K	1:50	<15	
Pax5	1H9	AF647	BioLegend	649704	Rat IgG2a, ĸ	1:100	<15	-
PD-1	29F.1A.12	BV421	BioLegend	135217	Rat IgG2a, ĸ	1:100	-	<15
PD-1 PD-1	29F.1A.12	PE	BioLegend	135206	Rat IgG2a, ĸ	1:100	<15	-
- 1-1	EH12.2H7	PE	BioLegend	329906	Mouse IgG1,	1:200	<15	-
RoRyt	AFKJS-9	-	Thermo	14-6988-82	к Rat IgG2a	1:200	-	-

SiglecF	E50-2440	PE	BD Biosciences	552126	Rat IgG2a, к	1:100	<15	-
SiglecF	1RNM44N	AF700	Thermo	56-1702-80	Rat IgG2a, ĸ	1:50	<15	-
SIRPα	P84	AF488	BioLegend	144024	Rat IgG1, κ	1:50	<15	-
SIRPα	P84	AF647	BioLegend	144027	Rat IgG1, ĸ	1:200	<15	-
αSMA	1A4	AF488	Thermo	53-9760-80	Mouse IgG2a, к	1:500	<15	-
αSMA	1A4	eF660	Thermo	53-9760-82	Mouse IgG2a, к	1:500	<15	-
SPARC	-	AF532	R&D	AF941 (Unconjugated)	Goat IgG	1:50	<15	-
TCRγδ	GL3	AF488	BioLegend	118128	Hamster IgG	1:50	<15	-
TCRγδ	GL3	PE	BioLegend	118108	Hamster IgG	1:100	<15	-
ΤCRγδ	B1	PE	BioLegend	331210	Mouse IgG ₁ , κ	1:100	<15	-
Tim-3	344823	AF532	R&D	MAB2365 (Unconjugated)	Rat IgG2a	1:25	<15	-
Tim-4	21H12	BV421	BD Biosciences	742773	Rat IgG1, κ	1:100	-	<15
Tryptase	AA1	-	AbCAM	Ab2378	Mouse IgG ₁	1:50	<15	-
Vα7.2	3C10	AF647	BioLegend	351726	Mouse IgG ₁ , κ	1:50	<15	-
Vimentin	O91D3	AF532	BioLegend	Custom	Mouse IgG2a	1:200	<15	-
anti-chicken IgY	-	FITC	Thermo	SA1-72000	Donkey IgG	1:200	<15	-
anti-goat IgG	-	AF488	Thermo	A-11055	Donkey IgG	1:400	<15	-
anti-hamster IgG	-	AF647	Thermo	A-21451	Goat IgG	1:400	<15	-
anti-rabbit IgG	-	AF488	Thermo	A-11034	Goat IgG	1:400	<15	-
anti-rabbit IgG	-	AF532	Thermo	A-11009	Goat IgG	1:400	<15	-
anti-rabbit IgG	-	AF555	Thermo	A-21428	Goat IgG	1:400	<15	-
anti-rabbit IgG	-	AF594	Thermo	A-11037	Goat IgG	1:400	>120	-
anti-rabbit IgG	-	AF647	Thermo	A-21245	Goat IgG	1:400	<15	-
anti-rabbit IgG	-	AF700	Thermo	A-21038	Goat IgG	1:400	<15	-
anti-rabbit IgG	-	AF750	Thermo	A-21039	Goat IgG	1:400	<15	-
anti-rabbit IgG	-	AF647	Thermo	A-31573	Donkey IgG	1:400	<15	-
anti-rabbit IgG	-	AF488	Thermo	Z25302	Goat Fab ₂	-	<15	-
anti-rabbit IgG	-	AF532	Thermo	Z25303	Goat Fab ₂	-	<15	-
anti-rabbit IgG	-	AF555	Thermo	Z25305	Goat Fab ₂	-	<15	-
anti-rabbit IgG	-	AF594	Thermo	Z25307	Goat Fab ₂	-	>120	-
anti-rabbit IgG	-	AF647	Thermo	Z25308	Goat Fab ₂	-	<15	-
anti-rat IgG	-	AF647	Thermo	A-21247	Goat IgG	1:400	<15	-

Table S2. IBEX panels for individual organs (See Figs. 3, 4A, and Movies S1-S7).

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379 Spleen: 16 parameters, 3 cycles

Cycle	Marker	Clone	Conjugate	Vendor	Catalog Number	Dilution
1	CD8	53-6.7	BV421	BioLegend	100738	1:200
	Autofluorescence	-	BV510*	-	-	-
	Autofluorescence	-	AF488**	-	-	-
	B220	RA3-6B2	PE	BD Biosciences	553090	1:400
	CD4	GK1.5	AF594	BioLegend	100446	1:200
	Foxp3	FJK-16s	eF660	Thermo	50-5773-82	1:50
	IgD	11-26c.2a	AF700	BioLegend	405729	1:50
2	F4/80	BM8	BV421	BioLegend	123132	1:50
	Autofluorescence	-	BV510*	-	-	-
	Collagen IV	-	None	AbCAM	19808	1:200
	Goat anti-rabbit IgG	-	AF488	Thermo	A-11034	1:400
	CD169	3D6.112	PE	Biolegend	142404	1:300
	CD4	GK1.5	AF594	BioLegend	100446	1:200
	CD11c	N418	AF647	BioLegend	117312	1:100
	MHC-II	M5/114.15.2	AF700	BioLegend	107622	1:100
3	CD68	FA-11	BV421	BioLegend	137017	1:200
	Autofluorescence	-	BV510*	-	-	-
	CD45	30-F11	AF488	BioLegend	103122	1:50
	CD31	MEC13.3	PE	BD Biosciences	553373	1:200
	JOJO-1	-	AF532	Thermo	J11372	1:20,000
	CD4	GK1.5	AF594	BioLegend	100446	1:200
	CD3	17A2	AF647	BD Biosciences	557869	1:400
	Ki-67	B56	AF700	BD Biosciences	561277	1:50

380

381 Thymus: 26 parameters, 5 cycles

Cycle	Marker	Clone	Conjugate	Vendor	Catalog Number	Dilution
1	CD138	281-2	BV421	BD Biosciences	562610	1:50
	CD11b	5C6	FITC	Bio-Rad	MCA711F	1:100
	β-3 Tubulin	AA10	AF532	BioLegend	Custom	1:100
	B220	RA3-6B2	PE	BD Biosciences	553090	1:400
	CD3	17A2	AF594	BioLegend	100240	1:400
	CD106	429	eF660	Thermo	50-1061-80	1:50
	Collagen IV	-	None	AbCAM	19808	1:200
	Goat anti-rabbit IgG	-	AF700	Thermo	A-21038	1:400
2	CD8	53-6.7	BV421	BioLegend	100738	1:200
	CD25	PC61.5	AF488	Thermo	53-0251-82	1:50
	Foxp3	FJK-16s	AF532	Thermo	58-5773-82	1:50
	CD4	RM4-5	eF570	Thermo	41-0042-82	1:100
	CD3	17A2	AF594	BioLegend	100240	1:400
	DEC205	NLDC-145	AF647	BioLegend	138204	1:50
	Ki-67	B56	AF700	BD Biosciences	561277	1:50
3	CD68	FA-11	BV421	BioLegend	137017	1:200
	SIRPα	P84	AF488	BioLegend	144024	1:50
	CD31	MEC13.3	PE	BD Biosciences	553373	1:200
	CD3	17A2	AF594	BioLegend	100240	1:400
	CD11c	N418	AF647	BioLegend	117312	1:100
	MHC-II	M5/114.15.2	AF700	BioLegend	107622	1:100
4	CD206	C068C2	BV421	BioLegend	141717	1:100
	αSMA	1A4	AF488	Thermo	53-9760-80	1:500
	CD44	IM7	AF532	BioLegend	Custom	1:100
	CD169	3D6.112	PE	Biolegend	142404	1:300
	CD3	17A2	AF594	BioLegend	100240	1:400
	CD45	30-F11	AF700	BioLegend	103128	1:50
5	JOJO-1	-	AF532	Thermo	J11372	1:20,000
	ΤCRγδ	GL3	PE	BioLegend	118108	1:100
	CD3	17A2	AF594	BioLegend	100240	1:400
	Cvtokeratin	C-11	AF647	BioLegend	628604	1:200

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383 Lung: 23 parameters, 4 cycles

Cycle	Marker	Clone	Conjugate	Vendor	Catalog Number	Dilution
1	CD206	C068C2	BV421	BioLegend	141717	1:100
	CD11b	5C6	FITC	Bio-Rad	MCA711F	1:100
	CD4	RM4-5	AF532	Thermo	58-0042-80	1:50
	Lyve-1	ALY7	eF570	Thermo	41-0443-82	1:100

	CD31	MEC13.3	AF594	BioLegend	102520	1:100
	CD11c	N418	AF647	BioLegend	117312	1:100
	SiglecF	1RNM44N	AF700	Thermo	56-1702-80	1:50
2	CD68	FA-11	BV421	BioLegend	137017	1:200
	Ly-6G	1A8	AF488	BioLegend	127626	1:50
	β-3 Tubulin	AA10	AF532	BioLegend	Custom	1:100
	B220	RA3-6B2	PE	BD Biosciences	553090	1:400
	CD31	MEC13.3	AF594	BioLegend	102520	1:100
	EpCAM	G8.8	AF647	BioLegend	118212	1:200
	MHC-II	M5/114.15.2	AF700	BioLegend	107622	1:100
3	CD138	281-2	BV421	BD Biosciences	562610	1:50
	KLRG1	2F1	AF488	BD Biosciences	561619	1:50
	IgA	-	AF555	Southern Biotech	1040-32	1:500
	CD31	MEC13.3	AF594	BioLegend	102520	1:100
	CD44	IM7	AF647	BioLegend	103018	1:100
	Collagen IV	-	None	AbCAM	19808	1:200
	Goat anti-rabbit IgG	-	AF700	Thermo	A-21038	1:400
4	αSMA	1A4	AF488	Thermo	53-9760-80	1:500
	JOJO-1	-	AF532	Thermo	J11372	1:20,000
	CD169	3D6.112	PE	Biolegend	142404	1:300
	CD31	MEC13.3	AF594	BioLegend	102520	1:100
	CD8	53-6.7	AF647	BioLegend	100724	1:200
	CD45	30-F11	AF700	BioLegend	103128	1:50

385 Small Intestine: 20 parameters, 3 cycles

Cycle	Marker	Clone	Conjugate	Vendor	Catalog Number	Dilution
1	CD8	53-6.7	BV421	BioLegend	100738	1:200
	CD35	8C12	BV510	BD Biosciences	740132	1:600
	CD4	RM4-5	AF532	Thermo	58-0042-80	1:50
	Foxp3	FJK-16s	eF570	Thermo	41-5773-82	1:50
	EpCAM	G8.8	AF594	BioLegend	118222	1:400
	CD3	17A2	AF647	BD Biosciences	557869	1:400
	IgD	11-26c.2a	AF700	BioLegend	405729	1:50
2	CD117	2B8	BV421	BioLegend	105827	1:50
	B220	RA3-6B2	BV510	BioLegend	103248	1:300
	KLRG1	2F1	AF488	BD Biosciences	561619	1:50
	DCAMKL1	-	None	AbCAM	Ab37994	1:50
	Goat anti-rabbit IgG	-	AF532	Thermo	A-11009	1:400
	IgA	-	AF555	Southern Biotech	1040-32	1:500
	EpCAM	G8.8	AF594	BioLegend	118222	1:400
	CD31	MEC13.3	AF647	BioLegend	102516	1:100
	Ki-67	B56	AF700	BD Biosciences	561277	1:50
3	CD45	30-F11	BV421	BioLegend	103134	1:100
	CD11b	5C6	FITC	Bio-Rad	MCA711F	1:100
	JOJO-1	-	AF532	Thermo	J11372	1:20,000
	Lyve-1	ALY7	eF570	Thermo	41-0443-82	1:100
	EpCAM	G8.8	AF594	BioLegend	118222	1:400
	CD11c	N418	AF647	BioLegend	117312	1:100
	MHC-II	M5/114.15.2	AF700	BioLegend	107622	1:100

386

387 Liver: 18 parameters, 4 cycles

Cycle	Marker	Clone	Conjugate	Vendor	Catalog Number	Dilution
1	CD4	GK1.5	BV421	BioLegend	100443	1:200
	Autofluorescence	-	BV510*	-	-	-
	Autofluorescence	-	AF488**	-	-	-
	LysM-tdTomato*	-	565-595 nm	-	-	-
	Laminin 1 + 2	-	None	AbCAM	Ab7463	1:100
	Goat anti-rabbit IgG	-	AF594	Thermo	A-11037	1:400
	E-cadherin	DECMA-1	AF647	BioLegend	147308	1:100
2	CD8	53-6.7	BV421	BioLegend	100738	1:200
	Autofluorescence	-	BV510*	-	-	-
	Autofluorescence	-	AF488**	-	-	-
	Desmin	-	None	AbCAM	Ab15200	1:200
	Goat anti-rabbit IgG Fab ₂	-	AF532	Thermo	Z25303	-
	Laminin 1 + 2	-	None	AbCAM	Ab7463	1:100
	Goat anti-rabbit IgG	-	AF594	Thermo	A-11037	1:400
	CXCR6	221002	AF647	Novus	FAB2145R	1:50
	CD44	IM7	AF700	BioLegend	103026	1:50
3	NK1.1	PK136	BV421	BioLegend	108731	1:50
	Autofluorescence	-	BV510*	-	-	-
	Autofluorescence	-	AF488**	-	-	-
	CD3	17A2	AF532	Thermo	58-0032-80	1:50
	Laminin 1 + 2	-	None	AbCAM	Ab7463	1:100

	Goat anti-rabbit IgG	-	AF594	Thermo	A-11037	1:400
	B220	RA3-6B2	AF647	BioLegend	103226	1:400
	CD45	30-F11	AF700	BioLegend	103128	1:50
4	Tim-4	21H12	BV421	BD Biosciences	742773	1:100
	Autofluorescence	-	BV510*	-	-	-
	CD11b	M1/70	AF488	BioLegend	101217	1:100
	Glutamine Synthetase	-	None	AbCAM	Ab49873	1:200
	Goat anti-rabbit IgG Fab ₂	-	AF532	Thermo	Z25303	-
	CD1d	1B1	PE	BD Biosciences	553846	1:100
	Laminin 1 + 2	-	None	AbCAM	Ab7463	1:100
	Goat anti-rabbit IgG	-	AF594	Thermo	A-11037	1:400
	CD11c	N418	AF647	BioLegend	117312	1:100
	MHC-II	M5/114.15.2	AF700	BioLegend	107622	1:100

389 Naïve and immunized LNs: 41 parameters, 10 cycles

Cycle	Marker	Clone	Conjugate	Vendor	Catalog Number	Dilution
1	PD-1	29F.1.A12	BV421	BioLegend	135218	1:100
	CD35	8C12	BV510	BD Biosciences	740132	1:500
	B220	RA3-6B2	AF488	BD Biosciences	557669	1:400
	JOJO-1	-	-	Thermo	J11372	1:20,000
	CD3	17A2	AF594	BioLegend	100240	1:100
	Bcl6	K112-91	AF647	BD Biosciences	561525	1:50
	Ki-67	B56	AF700	BD Biosciences	561277	1:50
2	CD200	OX-90	BV421	BD Biosciences	565547	1:50
	αSMA	1A4	AF488	Thermo	53-9760-80	1:200
	JOJO-1	-	-	Thermo	J11372	1:20,000
	Lyve-1	ALY7	eF570	Thermo	41-0443-82	1:100
	CD3	17A2	AF594	BioLegend	100240	1:100
	gp38	8.1.1	-	BioLegend	127401	1:50
	Goat anti-hamster IgG	-	AF647	Thermo	A-21451	1:400
	Collagen IV	-	-	AbCAM	19808	1:200
	Goat anti-rabbit IgG	-	AF700	Thermo	A-21038	1:400
3	F4/80	BM8	BV421	BioLegend	123132	1:50
	CD11b	5C6	FITC	Thermo	MA5-16529	1:100
	JOJO-1	-	-	Thermo	J11372	1:20,000
	CD207	eBioL31	PE	Thermo	12-2075-80	1:50
	CD3	17A2	AF594	BioLegend	100240	1:100
	CD11c	N418	AF647	BioLegend	117312	1:100
4	CD8	53-6.7	BV421	BioLegend	100738	1:200
	JOJO-1	-	-	Thermo	J11372	1:20,000
	CD4	GK1.5	PE	BD Biosciences	553730	1:200
	CD3	17A2	AF594	BioLegend	100240	1:100
	Foxp3	FJK-16s	eF660	Thermo	50-5773-82	1:50
5	CD206	C068C2	BV421	BioLegend	141717	1:100
	SIRPα	P84	AF488	BioLegend	144024	1:50
	JOJO-1	-	-	Thermo	J11372	1:20,000
	ΤCRγδ	GL3	PE	BioLegend	118108	1:100
	CD3	17A2	AF594	BioLegend	100240	1:100
	DEC205	NLDC-145	AF647	BioLegend	138204	1:50
6	CD68	FA-11	BV421	BioLegend	137017	1:200
	JOJO-1	-	-	Thermo	J11372	1:20,000
	CD169	3D6.112	PE	Biolegend	142404	1:300
	CD3	17A2	AF594	BioLegend	100240	1:100
	CD64	X54-5/7.1	AF647	BioLegend	139322	1:50
7	CD21	7E9	Pacific Blue	BioLegend	123413	1:200
	CD106	429	AF488	BioLegend	105710	1:50
	JOJO-1	-	-	Thermo	J11372	1:20,000
	CD31	MEC13.3	PE	BD Biosciences	553373	1:200
	CD3	17A2	AF594	BioLegend	100240	1:100
	CD23	B3B4	AF647	BioLegend	101611	1:50
3	CD138	281-2	BV421	BD Biosciences	562610	1:50
	IRF4	3E4	FITC	Thermo	11-9858-82	1:50
	JOJO-1	1-	•	Thermo	J11372	1:20,000
	SiglecF	E50-2440	PE	BD Biosciences	552126	1:100
	CD3	17A2	AF594	BioLegend	100240	1:100
	Pax5	1H9	AF647	BioLegend	649704	1:100
9	NK1.1	PK136	BV421	BioLegend	108731	1:50
	KLRG1	2F1	AF488	BD Biosciences	561619	1:56
	JOJO-1	-	-	Thermo	J11372	1:20,000
	GL-7	GL7	PE	BD Biosciences	561530	1:100
	CD3	17A2	AF594	BioLegend	100240	1:100
	CD44	IM7	AF647	BioLegend	103018	1:100
10	MHC-II	M5/114.15.2	BV421	BioLegend	107631	1:400
	IgD	11-26c.2a	AF488	BioLegend	405718	1:400
	JOJO-1	-	-	Thermo	J11372	1:20,000
	CD3	17A2	AF594	BioLegend	100240	1:100
	CD45	30-F11	AF647	BioLegend	103124	1:400

390 Table S3. IBEX panels for human LNs (See Fig. 5, Movie S8).

Clone Cycle Marker Conjugate Vendor Catalog Number Dilution Hoechst Biotium 40046 1:5000 SK1 AF488 CD8 BioLegend 344716 1:50 CD3 FOXP3 UCHT1 AF532 Thermo 58-0038-42 1:50 236A/E7 eF570 Thermo 41-4777-82 1:50 EpCAM 9C4 AF594 BioLegend 324228 1:500 CD25 M-A251 AF647 BioLegend 356127 1:50 CD4 RPA-T4 AF700 BioLegend 300526 1:25 2 Hoechst Biotium 40046 1:5000 -CD45 HI30 AF532 Thermo 58-0459-41 1:50 PD-1 EH12.2H7 PE BioLegend 329906 1:200 EpCAM 9C4 AF594 BioLegend 324228 1:500 CD69 FN50 AF647 BioLegend 310918 1:100 3 Biotium 40046 1:5000 Hoechst --HLA-DR L243 AF488 BioLegend 307619 1:200 SPARC Goat IgG AF532 R&D AF941 1:50 (Unconjugated) EpCAM 9C4 AF594 BioLegend 324228 1:500 KP1 Santa Cruz Sc-200060 CD68 AF647 1:100 Collagen IV AbCAM None Ab6586 1:200 Goat anti-rabbit IgG AF700 Thermo A-21038 1:400 4 Hoechst -AF532 Biotium 40046 1:5000 CD21 Bu32 BioLegend Custom 1:400 EpCAM 9C4 AF594 BioLegend 324228 1:500 CD138 MI15 AF647 356523 1:200 BioLegend Ki-67 B56 AF700 BD Biosciences 1:50 561277

391 Metastatic pancreatic LN: 17 parameters, 4 cycles

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393 Mesenteric LN: 66 parameters, 20 cycles

Cycle	Marker	Clone	Conjugate	Vendor	Catalog Number	Dilution	
1	Hoechst	-	-	Biotium	40046	1:5000	
	CD20	L26	AF488	Thermo	53-0202-82	1:200	
	SPARC	Goat IgG	Goat IgG AF532 R&D		AF941 1:50		
		-			(Unconjugated)		
	CD10	10 FR4D11 PE Caprico		103926	1:50		
				Biotechnologies			
	CD3	UCHT1	AF594	BioLegend	300446	1:100	
	BCL2	100	AF647	BioLegend	658705	1:25	
	Collagen IV	-	None	AbCAM	Ab6586	1:200	
	Goat anti-rabbit IgG	-	AF700	Thermo	A-21038	1:400	
2	Hoechst	-	-	Biotium	40046	1:5000	
	IgD	IA6-2	AF488	BioLegend	348216	1:25	
	CD21	Bu32	AF532	BioLegend	NA, Custom	1:600	
	CD138	MI15	PE	BioLegend	356504	1:200	
	CD3	UCHT1	AF594	BioLegend	300446	1:100	
	BCL6	K112-91	AF647	BD Biosciences	561525	1:25	
	CD31	WM59	AF700	BioLegend	303133	1:25	
3	Hoechst	-	-	Biotium	40046	1:5000	
	HLA-DR	L243	AF488	BioLegend	307620	1:100	
	CD23	EBVCS-5	AF532	BioLegend	NA, Custom	1:25	
	CD1c	L161	PE	BioLegend	331506	1:50	
	CD3	UCHT1	AF594	BioLegend	300446	1:100	
	CD163	GH1/61	AF647	BioLegend	333620	1:100	
	CD11c	B-Ly6	AF700	BD Biosciences	561352	1:25	
4	Hoechst	-	-	Biotium	40046	1:5000	
	CD8	SK1	AF488	BioLegend	344716	1:25	
	CD4	RPA-T4	AF532	Thermo	58-0049-42	1:25	
	FOXP3	236A/E7	eF570	Thermo	41-4777-82	1:25	
	CD3	UCHT1	AF594	BioLegend	300446	1:100	
	CD25	M-A251	AF647	BioLegend	356128	1:50	
	Ki-67	B56	AF700	BD Biosciences	561277	1:50	
5	Hoechst	-	-	Biotium	40046	1:5000	
	ICOS	CS98.4A	AF488	BioLegend	313514	1:25	
	CXCL13	Goat IgG	AF532	R&D	AF801	1:25	
					(Unconjugated)		
	PD-1	EH12.2H7	PE	BioLegend	329906	1:100	
	CD3	UCHT1	AF594	BioLegend	300446	1:100	
	CD69	FN50	AF647	BioLegend	310918	1:25	
6	Hoechst	-	-	Biotium	40046	1:5000	
	CD117	104D2	AF488	BioLegend	313234	1:50	
	Lyve-1	Goat IgG	AF532	R&D	AF2089	1:100	
					(Unconjugated)		

	CD25	E11	DE	Piol ogond	222406	1.900
	CD35 CD3	E11 UCHT1	PE AF594	BioLegend BioLegend	333406 300446	1:800
	CD3 CD68		AF594 AF647	BioLegend		
		KP1		Santa Cruz	sc-20060	1:100
-	CD38	HIT2	AF700	BioLegend	303524	1:25
	Hoechst	-	-	Biotium	40046	1:5000
	Clec9a	8F9	AF488	BioLegend	Custom	1:25
	Tim-3	344823	AF532	R&D	MAB2365	1:25
	1054	1054.054	55	D : 1	(Unconjugated)	1.05
	IRF4	IRF4.3E4	PE	BioLegend	646404	1:25
	CD3	UCHT1	AF594	BioLegend	300446	1:100
	DC-SIGN	9E9A8	AF647	BioLegend	330112	1:50
3	Hoechst	-	-	Biotium	40046	1:5000
	CXCL12	79018	AF532	R&D	MAB350-500	1:25
					(Unconjugated)	
	ΤCRγδ	B1	PE	BioLegend	331210	1:100
	CD3	UCHT1	AF594	BioLegend	300446	1:100
	Va7.2	3C10	AF647	BioLegend	351726	1:50
)	Hoechst	-	-	Biotium	40046	1:5000
	α-SMA	1A4	AF488	Thermo	53-9760-82	1:100
	CD45	HI30	AF532	Thermo	58-0459-42	1:25
	CD106	STA	PE	BioLegend	305806	1:50
	CD3	UCHT1	AF594	BioLegend	300446	1:100
	CD44	IM7	AF647		103018	1:50
10		1/1/1	71 047	BioLegend	40046	1:5000
J	Hoechst	Contine	- AF532	Biotium	40046 AF2745	1:5000
	Lumican	Goat IgG	AF332	R&D	(Unconjugated)	1.50
	CD24		PE	Thormo		1.50
	CD34	QBEND/10		Thermo Biol occord	MAI-10205	1:50
	CD3	UCHT1	AF594	BioLegend	300446	1:100
4	CD54	HA58	AF647	BioLegend	353114	1:50
1	Hoechst	-	-	Biotium	40046	1:5000
	NF-H/NF-M	SM1-35	AF488	BioLegend	835614	1:50
	CD3	UCHT1	AF594	BioLegend	300446	1:100
	CD66b	G10F5	AF647	BioLegend	305110	1:25
2	Hoechst	-	-	Biotium	40046	1:5000
	CD166	EPR2759	AF488	AbCAM	Ab197543	1:125
	Fibronectin	2F4	AF532	Novus	NBP2-22113AF532	1:25
	CD39	A1	PE	BioLegend	328208	1:50
	CD3	UCHT1	AF594	BioLegend	300446	1:100
	Cytokeratin	AE1/AE3	eF660	Thermo	50-9003-82	1:100
13	Hoechst	-	-	Biotium	40046	1:5000
	MARCO	Rabbit IgG	-	Thermo	PA5-64134	1:25
	Zenon Fab	-	AF532	Thermo	Z25303	NA
	CD94	DX22	PE	BioLegend	305506	1:50
	CD3	UCHT1	AF594	BioLegend	300446	1:100
	EpCAM	9C4	AF647	BioLegend	324212	1:100
4	Hoechst	-	-	Biotium	40046	1:5000
	β-Tubulin 3	TUJ1	AF532	BioLegend	Custom	1:50
	CD49a	TS2/7	PE	•	328304	1:50
				BioLegend		
	CD3	UCHT1	AF594 AF647	BioLegend	300446	1:100
_	IgM	EPR5539-65-4	-	AbCAM	Ab200629	1:100
15	Hoechst	-	-	Biotium	40046	1:5000
	IgA2	A9604D2	AF488	SouthernBiotech	9140-30	1:500
	p53	PAb 240	PE	Novus	NB200-103PE	1:50
	CD3	UCHT1	AF594	BioLegend	300446	1:100
	IgA1	B3506B4	AF647	SouthernBiotech	9130-31	1:500
6	Hoechst	-	-	Biotium	40046	1:5000
	Laminin 1+2	Rabbit IgG		AbCAM	Ab7463	1:100
	Zenon Fab	-	AF488	Thermo	Z25302	NA
	Lysozyme	Rabbit IgG	1 -	AbCAM	Ab2408	1:50
	Zenon Fab	-	AF532	Thermo	Z25303	NA
	CD3	UCHT1	AF594	BioLegend	300446	1:100
7	Hoechst	-	-	Biotium	40046	1:5000
	Keratin 18	1G11C4	CoraLite488	ProteinTech	CL488-66187	1:50
	Vimentin	O91D3	AF532	BioLegend	Custom	1:200
	CD3	UCHT1	AF594	BioLegend	300446	1:100
8	Hoechst	-	-	Biotium	40046	1:5000
	Desmin	Y66	AF488	AbCAM	Ab185033	1:200
	CD3	UCHT1	AF594	BioLegend	300446	1:100
	Tryptase	AA1	-	AbCAM	Ab2378	1:50
	Zenon Fab	-	AF700	Thermo	Z25011	NA
9	Hoechst	-	-	Biotium	40046	1:5000
-	Donkey anti-goat IgG	-	AF488	Thermo	A11055	1:1000
	CD45	- F10-89-4	PE/iFluor594	Caprico	1016185	1:50
	0040	1 10-03-4	FE/IFI001594	Biotechnologies	1010105	1.50
	CD2				200446	1.100
	CD3	UCHT1	AF594	BioLegend	300446	1:100
	Donkey anti-rabbit IgG	-	AF647	Thermo	A31573	1:1000
20	Hoechst	-	-	Biotium	40046	1:5000
	Keratin 14	Poly9060	-	BioLegend	906004	1:50
	Donkey anti-chicken IgY CD3	- UCHT1	FITC AF594	Thermo BioLegend	SA1-72000 300446	1:200

	Goat anti-rat IgG	-	AF647	Thermo	A21247	1:1000
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Table S4. Reagents used for multi-plex Opal IHC experiments (See Figs. 6B, S7A, and Movie

- S9).

Marker	Clone	Conjugate	Vendor	Cat No.	Isotype	Dilution	Time to bleach with LiBH₄
CD3e	D4V8L	None	CST	99940	Rabbit IgG	1:100	-
CD4	D7D2Z	None	CST	25229	Rabbit IgG	1:100	-
CD8	D4W2Z	None	CST	98941	Rabbit IgG	1:200	-
CD11c	D1V9Y	None	CST	97585	Rabbit IgG	1:100	-
CD19	D4V4B	None	CST	90176	Rabbit IgG	1:300	-
CD138	281-2	PE	BioLegend	142504	Rat IgG2a, к	1:100	-
Collagen IV	-	None	AbCam	19808	Rabbit IgG	1:50	-
CXCL9	-	None	R&D	AF-492-NA	Goat IgG	1:50	-
E-cadherin	DECMA-1	AF488	Thermo	53-3249-82	Rat IgG1, к	1:100	-
F4/80	D2S9R	None	CST	70076	Rabbit IgG	1:100	-
F4/80	BM8	BV421	BioLegend	123132	Rat IgG2a, к	1:50	-
Foxp3	D608R	None	CST	12653	Rabbit IgG	1:200	-
Laminin	-	None	AbCam	Ab7463	Rabbit IgG	1:100	-
MHCII	-	None	AbCam	Ab180779	Rabbit IgG	1:100	-
Anti-rabbit IgG		HRP			Goat IgG	1:5	-
-	-	Opal 520	Akoya Biosciences	FP1487001KT	-	-	>30 minutes
-	-	Opal 540	Akoya Biosciences	FP1494001KT	-	-	>30 minutes
-	-	Opal 570	Akoya Biosciences	FP1488001KT	-	-	30 minutes
-	-	Opal 620	Akoya Biosciences	FP1495001KT	-	-	>30 minutes
-	-	Opal 650	Akoya Biosciences	FP1496001KT	-	-	30 minutes
-	-	Opal 690	Akoya Biosciences	FP1497001KT	-	-	30 minutes

Table S5. Reagents used in combined IBEX oligonucleotide-based staining panel. (See Figs. 6D, S7C-D, and Movie S9).

Cycle	Antibody/Dye	Clone	Vendor	Cat No.	Imaging oligo sequence
1	Hoechst	-	Biotium	40046	
	SIRPa AF488	P84	BioLegend	144024	-
	Foxp3 AF532	FJK-16s	Thermo	58-5773-82	-
	CD31 PE	MEC13.3	BD Biosciences	553373	-
	CD11c AF647	N418	BioLegend	117312	-
	Ki-67 AF700	B56	BD Biosciences	561277	-
2	Hoechst	-	Biotium	40046	-
	CD169 AF532*	3D6.112	BioLegend	142425	ATGACTGTCGTCAATTG
	IgD Atto 550*	11.26c.2a	BioLegend	405745	GGACAACGGATATGATG
	CD11b AF647*	M1/70	BioLegend	101265	ACAAATGAGCCTTCATG
	MHCII IR700*	M5/114.15.2	BioLegend	107653	ATCATACTGGTGACCTG
3	Hoechst	-	Biotium	40046	-
	CD45 AF532*	30-F11	BioLegend	103159	TCTGCTCCATAGCCATG
	CD68 Atto 550*	FA-11	BioLegend	137031	TCCCGTGAAAGAAAGTG
	CD3 AF647*	17A2	BioLegend	100251	ATCGAGCGGACATACTG
Non-	B220 AF488*	RA3-6B2	BioLegend	103263	ATTATGAGGTGTAGGTG
IBEX**	CD11c AF594*	N418	BioLegend	117355	GCAAGCGTCCATAACTG

*Denotes fluorescent label provided by complementary fluorescent oligonucleotides.

**Additional reagents tested (Figure S7C).

407 Movies S1-S9 Legends

408 Movie S1. High dimensional imaging of the spleen using IBEX. Confocal images of mouse
409 spleen tissue from a 3 cycle 16 parameter IBEX experiment with CD4 serving as a fiducial. See
410 Fig. 3. Data are representative of 2 similar experiments.

411 **Movie S2. High dimensional imaging of the thymus using IBEX.** Confocal images of mouse 412 thymus tissue from a 5 cycle 26 parameter IBEX experiment with CD3 serving as a fiducial. See

413 Fig. 3. Data are representative of 2 similar experiments.

414 Movie S3. High dimensional imaging of the lung using IBEX. Confocal images of mouse lung
415 tissue from a 4 cycle 23 parameter IBEX experiment with CD31 serving as a fiducial. See Fig. 3.
416 Data are representative of 2 similar experiments.

417 **Movie S4. High dimensional imaging of the small intestine using IBEX.** Confocal images of 418 mouse small intestine tissue from a 3 cycle 20 parameter IBEX experiment with EpCAM serving as 419 a fiducial. See Fig. 3. Data are representative of 2 similar experiments.

420 **Movie S5. High dimensional imaging of the liver using IBEX.** Confocal images of liver tissue 421 from a LysM-tdTomato mouse. A 4 cycle 18 parameter IBEX experiment was performed with 422 Laminin serving as a fiducial. See Fig. 3. Data are representative of 2 similar experiments.

423 Movie S6. High dimensional imaging of naïve and immunized LNs using IBEX. Confocal
424 images of pLNs from naïve and SRBC-immunized mice from 10 cycle 41 parameter IBEX
425 experiments. See Fig. 4A. Data are representative of 2 similar experiments.

426 Movie S7. Comparable staining observed by serial and iterative immunofluorescence 427 methods. Confocal images of inguinal LN (iLN) or pLNs from SRBC-immunized mice 428 demonstrating qualitatively similar staining patterns when antibody panels were applied on 429 individual sections alone (serial) versus on the same section iteratively (IBEX). See Figs. 4 and S5. 430 Data are representative of 2 similar experiments.

Movie S8. IBEX scales to capture ultra-high content imaging in large human tissues.
Confocal images of human LN tissue section with metastatic lesions from a 4 cycle 17 parameter
IBEX experiment or human mesenteric LN from a 20 cycle 66 parameter IBEX experiment. See
Fig. 5. Data are representative of similar experiments in normal and diseased human LNs.

435 Movie S9. Extensions of IBEX workflow to include Opal fluorophores and oligo-conjugated
 436 antibodies. Representative confocal images from a 10 parameter 4 cycle IBEX experiment
 437 incorporating Opal fluorophores performed on heavily fixed mouse pLN tissue sections. Confocal

- 438 images from a 13 parameter 3 cycle IBEX experiment performed on mouse inguinal LN sections.
- 439 Markers were visualized using either fluorescently-conjugated antibodies (Cycle 1) or oligo-
- 440 conjugated antibodies and complementary fluorescent oligos (Cycles 2-3). See Figs. 6 and S7.
- 441 Data are representative of 2-4 similar experiments.

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