

IBEX: A versatile multiplex optical imaging approach for deep phenotyping and spatial analysis of cells in complex tissues

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The diverse composition of mammalian tissues poses challenges for understanding the cell-cell interactions required for organ homeostasis and how spatial relationships are perturbed during disease. Existing methods such as single-cell genomics, lacking a spatial context, and traditional immunofluorescence, capturing only two to six molecular features, cannot resolve these issues. Imaging technologies have been developed to address these problems, but each possesses limitations that constrain widespread use. Here we report a method that overcomes major impediments to highly multiplex tissue imaging. "Iterative bleaching extends multiplexity" (IBEX) uses an iterative staining and chemical bleaching method to enable high-resolution imaging of >65 parameters in the same tissue section without physical degradation. IBEX can be employed with various types of conventional microscopes and permits use of both commercially available and user-generated antibodies in an "open" system to allow easy adjustment of staining panels based on ongoing marker discovery efforts. We show how IBEX can also be used with amplified staining methods for imaging strongly fixed tissues with limited epitope retention and with oligonucleotide-based staining, allowing potential cross-referencing between flow cytometry, cellular indexing of transcriptomes and epitopes by sequencing, and IBEX analysis of the same tissue. To facilitate data processing, we provide an open-source platform for automated registration of iterative images. IBEX thus represents a technology that can be rapidly integrated into most current laboratory workflows to achieve high-content imaging to reveal the complex cellular landscape of diverse organs and tissues.

high-dimensional imaging | tissue immunity | quantitative microscopy | immune system | immunofluorescence

types, presenting a major challenge to understanding the cell-cell interactions required for homeostasis as well as the compositional changes associated with disease. To address this complexity, several multiplexed imaging methods utilizing conventional microscopes and commercially available antibodies have been described to overcome the target detection limitations of conventional immunohistochemistry (IHC) or immunofluorescence (IF) imaging (1–8). The majority of these methods generate high-dimensional datasets through an iterative, multistep process (i.e., a cycle) that includes 1) immunolabeling with antibodies, 2) image acquisition, and 3) fluorophore inactivation or antibody/chromogen removal. While these methods are capable of generating high-dimensional datasets, they are greatly limited by the number of markers visualized per cycle or length of time required for each cycle, or they involve special fluid-handling platforms not generally available to most laboratories (1).

Commercial systems based on the codetection by indexing method (9) have facilitated the acquisition of multiplex imaging data by providing a fully automated instrument for cyclic imaging. Despite this advancement, the proprietary nature of this method imposes constraints on the reagents available for use as well as the number of markers to be imaged for each round. Furthermore, cyclic imaging methods that employ a small number of markers per cycle (i.e., fewer than three) may result in tissue loss due to the stress of repeated fluid exchanges. To this end, imaging techniques such as multiplexed ion beam imaging (10) and imaging mass cytometry (11) enable the capture of multiparameter data without cyclic imaging. However, both of these methods require specialized instrumentation and consumables, with the latter often again limited in breadth to choices made by the supplier, not the investigator. This constrains their capacity for broadly analyzing human or

Significance

Single-cell flow cytometry and genomic methods are rapidly increasing our knowledge of the diversity of cell types in metazoan tissues. However, suitably robust methods for placing these cells in a spatial context that reveals how their localization and putative interactions contribute to tissue physiology and pathology are still lacking. Here we provide a readily accessible pipeline (IBEX) for highly multiplex immunofluorescent imaging that enables a fine-grained analysis of cells in their tissue context. Additionally, we describe extensions of the IBEX workflow to handle hard-to-image tissue preparations and a method to facilitate direct integration of the imaging data with flow cytometry and sequencing technologies.

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experimental animal tissues with respect to lists of validated antibodies, the ability to work across various established protocols for tissue processing, and the capacity for real-time changes to the epitope target list based on data emerging from high-content methods such as single-cell RNA sequencing (scRNA-seq).

To address the increasing need for high-content analysis of tissues for projects such as the Human Cell Atlas and others, the field requires a fully open and extensible method for multiplex imaging. Our laboratory has extensively characterized murine and human immune responses using quantitative multiparameter imaging of fixed frozen samples (12-18). Importantly, this method of tissue fixation preserves tissue architecture and cellular morphology, is archivable and compatible with large-volume imaging (19), and, in its optimal form, eliminates technical challenges posed by formalin-fixed, paraffin-embedded (FFPE) samples. Leveraging this experience and our original single-cycle histocytometry method for multiplex data acquisition (12), we have now developed "iterative bleaching extends multiplexity" (IBEX). This imaging technique reduces the time per cycle, uses a high number of antibodies per cycle, employs widely available reagents and instruments, provides open-source software for image alignment, and minimizes physical damage to the tissue during multiple imaging cycles. Beyond the basic IBEX workflow, we have developed extensions to achieve multiparameter imaging of heavily fixed tissues with limited retention of target epitopes and have incorporated commercially available oligonucleotide-conjugated antibodies to enable direct crosscomparisons to flow cytometry and scRNA-seq data obtained by the cellular indexing of transcriptomes and epitopes by sequencing (CITE-seq) method (20). In addition to describing the specifics of the IBEX procedure, we provide multiple examples of the use of IBEX to analyze both immune and parenchymal cells in a diverse array of mouse and human tissues to illustrate the general applicability of the technique. The IBEX method described here can be rapidly integrated into current laboratory workflows to obtain high-dimensional imaging datasets from a wide range of animal and human tissues.

Results

IBEX Builds and Improves upon Existing Iterative Imaging Techniques.

Iterative imaging methods typically use either fluorophore bleaching or antibody/chromogen removal to obtain multiparameter datasets (1–8). Due to the harsh and variable conditions required to remove chromogens and antibodies with diverse target affinities, we pursued a strategy based on fluorophore bleaching. To achieve an efficient means to increase the number of markers visualized on a single section, we sought an inactivation method that could bleach a wide range of fluorophores in minutes without epitope loss or tissue destruction. While H₂O₂ in alkaline solution has been reported to inactivate Cy3- and Cy5-conjugated antibodies in human FFPE samples (3), we observed significant tissue loss using this formulation over multiple cycles with fixed frozen samples (SI Appendix, Fig. S1A). Adams et al. demonstrated the initial feasibility of using borohydride derivates to bleach fluorophores; however, their fluorophore quenching method required 2 h per cycle and comprised only three distinct imaging channels (6), making direct application for highly multiplex imaging impractical. To expand upon this method, we tested antibodies directly conjugated to fluorophores with excitation and emission spectra spanning from 405 nm to 750 nm. We consistently found that the following fluorophores were inactivated within 15 min of exposure to 1 mg/mL of lithium borohydride (LiBH₄): Pacific blue, Alexa Fluor (AF) 488, fluorescein isothiocyanate (FITC), AF532, phycoerythrin (PE), AF555, eFluor (eF) 570, AF647, eF660, and AF700. Brilliant Violet conjugates BV421 and BV510 bleached within 15 min of exposure to 1 mg/mL of LiBH₄ in the presence of light. In contrast, AF594, eF615, and the nuclear markers JOJO-1 and Hoechst required more than 120 min for significant loss of fluorescence signal (SI Appendix, Table S1), permitting these probes to be used as fiducials for alignment of images emerging from iterative cycles.

To prevent tissue destruction over multiple cycles, we evaluated several different tissue adhesives and found that chrome gelatin alum securely adhered tissues to glass coverslips and slides, permitting more than 15 cycles to be performed with no appreciable loss to the tissue (SI Appendix, Fig. S1 B and C). We next reduced the antibody labeling time from 6 to 12 h to 30 to 45 min by designing programs for a nonheating microwave that facilitates rapid antibody penetration into the section. Finally, although IBEX was designed to simply bleach the fluorophores, it was important to assess whether LiBH4 treatment physically removes antibodies from the tissue, as this would have direct bearing on both the design and order of imaging panels. To examine this issue, mouse lymph node (LN) sections were immunolabeled with various primary antibodies, imaged, treated with LiBH₄, and then incubated with a secondary antibody that would react with the primary antibody if it was still present on the tissue. For most of the antibody isotypes tested, we found almost identical staining patterns with the primary and secondary antibodies, indicating that LiBH₄ acts primarily by fluorophore bleaching without stripping the fluorescently conjugated antibodies themselves (SI Appendix, Fig. S2).

The resulting method, IBEX, reduces the fluorophore inactivation and antibody labeling steps to less than 1 h (Fig. 1A). We first tested IBEX in practical use by examining the image quality that could be obtained from a three-cycle analysis of mouse LNs (Fig. 1B), a tissue with which we had extensive experience using multiparameter, single-cycle staining and image collection. This initial test used six to eight markers per cycle and showed that all fluorophores, except for AF594 and JOJO-1, bleached rapidly in the presence of LiBH₄ treatment with or without light, with no appreciable signal present after 10 min (Fig. 1 C and D). These findings show that the IBEX pipeline performs as designed and allows for the rapid capture of high-quality, multiplexed imaging data over multiple cycles without tissue loss.

A SimpleITK Image Registration Pipeline. The IBEX method yields a series of images that are collected separately. To properly process these data and correctly assign markers to individual cells, it is essential that all of the images be aligned at high resolution. While various registration algorithms have been reported (21, 22), we sought a method that could align large datasets, was flexible in terms of the repeated markers (fiducials) utilized, and provided both a qualitative and quantitative metric for registration. For these reasons, we developed a workflow using SimpleITK, a simplified open-source interface to the Insight Toolkit (ITK) that is compatible with multiple programming languages (23, 24). The SimpleITK workflow is an image intensity-based form of image alignment that relies on a repeated marker channel (fiducial) for registration. Due to the resistance of AF594, JOJO-1, and Hoechst to bleaching, we utilized markers in these fluorophores as fiducials. For a multicycle IBEX experiment, a "fixed" image z-stack was selected, and all other "moving" images were resampled to this image. A crosscorrelation matrix was generated on the repeated marker channels to provide a quantitative means for assessing the quality of image registration (Fig. 24). To test the fidelity of this method, a three-cycle IBEX experiment was performed on mouse spleen sections labeled with the nuclear marker JOJO-1 and membrane label CD4 AF594. For these experiments, JOJO-1 was selected as the fiducial used for image registration; however, CD4 (also repeated in these experiments) showed pixel-to-pixel alignment as reflected in the images (Fig. 2B). Importantly, this platform can readily scale to handle large datasets (>260 GB) comprised of 20 cycles of imaging (SI Appendix, Fig. S1 B-D). The SimpleITK registration workflow thus provides the needed cell-cell registration across x-y-z dimensions obtained via iterative imaging cycles using IBEX.

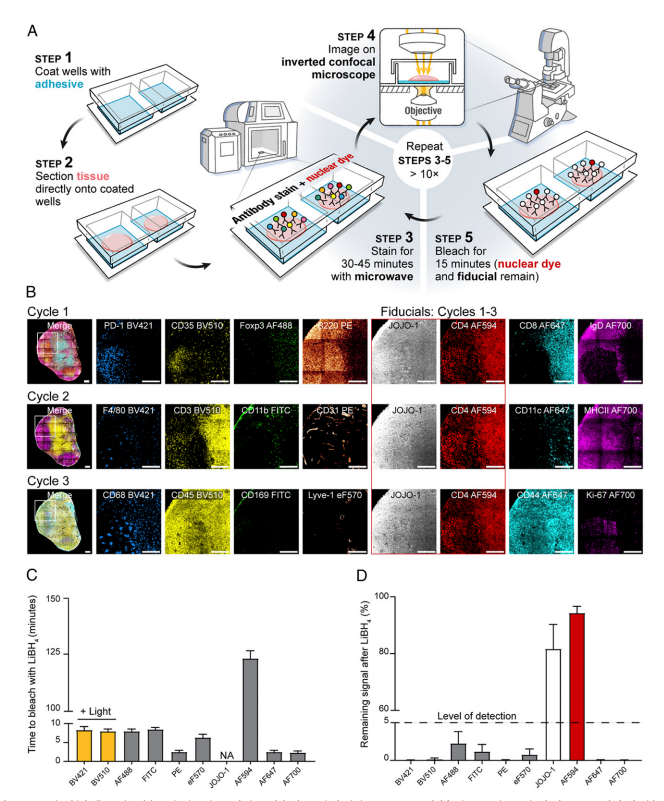


Fig. 1. IBEX is a high-dimensional, iterative imaging technique. (A) Schematic depicting IBEX protocol. (B) Mice were immunized subcutaneously (s.c.) with 25 μ L of SRBCs on day 0 and 7. On day 14, pLN tissue sections were labeled with three separate imaging panels. (Scale bar: 150 μ m.) Light refers to bleaching with LiBH₄ while the sample was illuminated with a metal halide lamp and DAPI filter. (C) Time required to bleach respective fluorophores using LiBH₄. NA, no appreciable loss of signal over multiple hours of LiBH₄ exposure. (D) Percentage of fluorophore signal remaining after 15 min of LiBH₄ treatment. Data are pooled from two similar experiments and presented as mean \pm SEM.

IBEX Is a Versatile Imaging Method. One obstacle to the wide adoption of existing multiplex imaging methods is the need for specialized instruments or custom imaging chambers, a luxury

not afforded to all laboratories. In contrast, IBEX is easily adaptable to diverse microscope systems and has no restrictions on the substrate (e.g., slide, coverslip) used for imaging

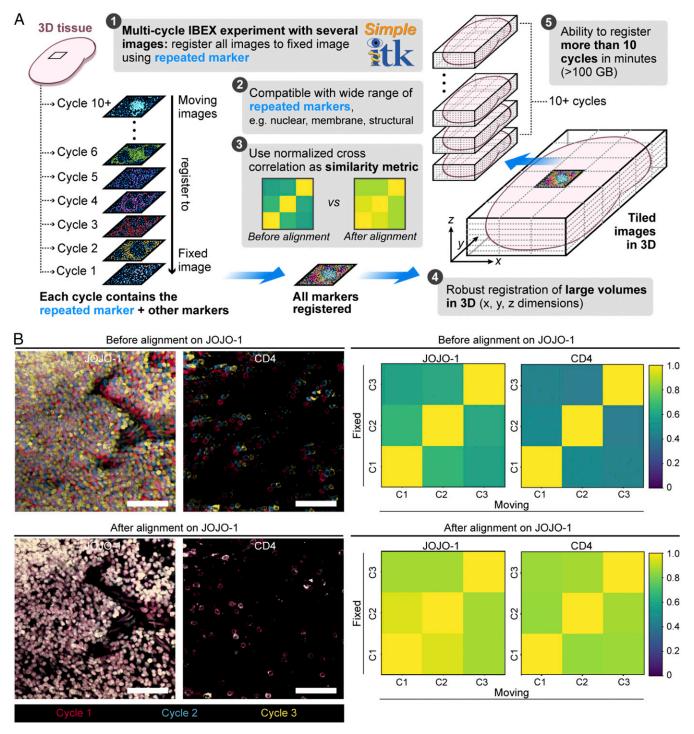


Fig. 2. Image alignment with SimpleITK image registration pipeline. (A) Workflow for SimpleITK image registration pipeline. (B) Confocal images showing JOJO-1 and CD4 from three consecutive IBEX cycles before and after alignment using the nuclear marker JOJO-1 as a fiducial across all three cycles ("C," Right). CD4 was also repeated and shows cell-cell registration after JOJO-1 alignment. (Scale bar: 50 μm.) Cross-correlation similarity matrices before and after alignment with JOJO-1 for JOJO-1 and CD4 channels. All experiments are representative of at least two similar experiments.

(SI Appendix, Fig. S3A). As a proof of concept, immunized mouse LN sections adhered to slides were visualized using an upright confocal microscope (n = 4 cycles, n = 6 markers per cycle) or an inverted fluorescence microscope (n = 4 cycles, n = 4 markers per cycle; SI Appendix, Fig. S3 B and C). These results demonstrate the compatibility of IBEX with a wide range of imaging systems; however, it is worth noting that the microscope system (confocal versus widefield) and configuration (light source, detectors, filter cubes) will dictate the image acquisition time, number of markers per cycle, and sample type that can be effectively imaged (5- vs. 30um tissue thickness).

In the case of animal studies, it is also very useful to be able to integrate antibody staining with imaging of fluorescent marker proteins expressed by engineered cells transferred into animals or expressed in situ. We therefore next investigated whether the IBEX method could be used to image tissues from animals

expressing fluorescent proteins (25). To this end, high-dimensional imaging was performed on thymic tissues from transgenic animals expressing the following fluorescent proteins (FPs): cyan FP (CFP), green FP (GFP), yellow FP (YFP), and red FP (RFP) (26). No appreciable loss in signal was observed after 10 IBEX cycles for any of the FPs examined (*SI Appendix*, Fig. S4 *A* and *B*). The photostable FPs were used as fiducials for a four-cycle IBEX experiment that incorporated the bleachable fluorophores AF647 and AF700, yielding a dataset that provided information on clonality (CFP, GFP, YFP, RFP) of T cells (CD4, CD8, Foxp3) and myeloid cells (CD11c, MHC II) in the thymus (*SI Appendix*, Fig. S4C).

To determine how IBEX performs using sections from a variety of tissues, we performed experiments using three to five cycles of IBEX with sections from murine spleen, thymus, lung, small intestine, and liver (Fig. 3 A and B and Movies S1–S5 and SI Appendix, Table S2). It is important to note that the cycle and marker numbers described here are provided as a proof of concept and do not reflect technical limitations of the method. The antibody panels were designed to capture the major cellular populations and structures present in each organ, and fluorophores were chosen to avoid native tissue autofluorescence. Organ-specific fiducials were selected based on expression throughout the tissue, e.g., EpCAM to mark the epithelium of the small intestine and laminin for the liver sinusoids. Collectively, these data confirm the ability to use IBEX to obtain high-quality, multiplexed imaging datasets from a wide range of tissues.

IBEX Enables Highly Multiplex, Quantitative Imaging. The design principles of IBEX were chosen to enable a very high number of parameters to be attained in the analysis of an individual tissue sample. To determine how extensively the multiplexing capacity of IBEX can be pushed, we first performed 10-cycle, 41-parameter IBEX experiments on LNs obtained from naïve and sheep red blood cell (SRBC)-immunized mice (Fig. 4A and SI Appendix, Table S2 and Movie S6). While epitope loss has been described for other iterative imaging techniques (5), we minimized this problem by increasing the number of markers per cycle and grouping markers present on the same cell in the same cycle. We observed qualitatively similar staining patterns when antibody panels were applied on individual sections alone (serial) versus on the same section iteratively (IBEX; SI Appendix, Fig. S5A and Movie S7). Therefore, quantitative differences observed between the two methods likely reflect biological differences resulting from variations in the magnitude of the immune response in individual LNs and not technical differences associated with epitope loss or steric hindrance (SI Appendix, Fig. S5B and Movie S7).

To assess the quality of data generated by the IBEX method, we employed the open-source, computational histology topography cytometry analysis toolbox (histoCAT) to quantify differences in LN organization resulting from immunization (27). Individual cells were segmented based on membrane and nuclear labels with Ilastik (28) and CellProfiler (29) and then analyzed using the histoCAT graphical user interface (SI Appendix, Fig. S64). The unsupervised clustering algorithm Phenograph (30) identified 29 phenotype clusters shared across the naïve and immunized LNs that could be visualized using the data dimensionality reduction method t-SNE (31) in histoCAT (Fig. 4B). As a testament to the fidelity of cell-cell alignment, phenotype clusters were often characterized by the expression of several different markers present in distinct imaging cycles (SI Appendix, Fig. S6 B and C). The abundance of these cell phenotypes varied from naïve and immunized LNs (SI Appendix, Fig. S6D) and could be manually annotated based on marker expression to reveal an increase in plasma cells (PCs; cluster 10) and germinal center (GC) B cells (cluster 6) in immunized LNs (Fig. 4C). As histoCAT relies on nuclear and membrane-based cell segmentation, it suffers from limitations frequently encountered with this approach: miscalling of phenotypes due to spatial overlap

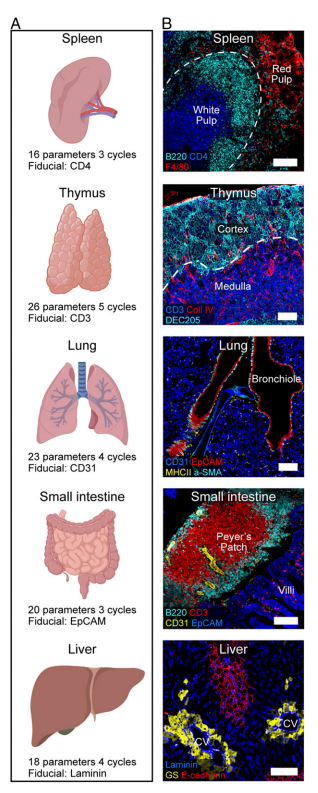


Fig. 3. IBEX in multiple murine organs. (A) IBEX experimental parameters. (B) Confocal images from IBEX experiments in various mouse organs. Liver: central vein (CV) and glutamine synthetase (GS). (Scale bar: $100 \mu m$.) Movies 1-55 show additional details.

and improper segmentation of nonlymphocyte populations (32). The former is evident in the identification of the B cell-specific transcription factor pax5 (33) on CD3⁺CD4⁺PD-1⁺Bcl6⁺

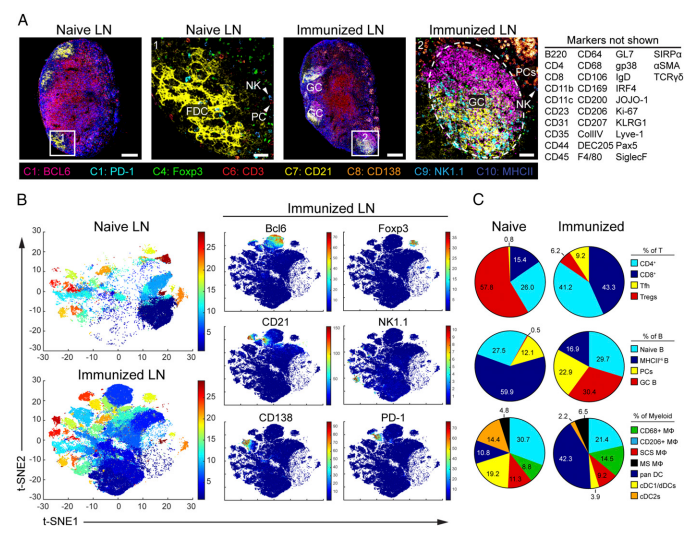


Fig. 4. Visualization and quantification of LN populations using IBEX and histoCAT following immune perturbation. (A) Confocal images of pLNs from naïve and SRBC-immunized mice from 10-cycle ("C" at bottom), 41-parameter IBEX experiments. NK, natural killer; FDC, follicular DC. (Scale bars: left to right, 100 μm, 25 μm, 100 μm, and 50 μm.) (B) t-SNE plots from naïve and immunized LNs identified by Phenograph clustering using segmented cells in histoCAT (naïve, n = 32,091; immune, n = 80,355). Color reflects the cluster ID number (from 1 to 29). Single plots show separation of representative markers into discrete clusters, with color map showing relative expression levels based on Z score-normalized marker intensity values. (C) Phenograph clusters identified by histoCAT were phenotyped based on marker expression and expressed as a proportion of lineage. MΦ, macrophage: SCS, subcapsular sinus: MSM, medullary sinus; dDC, dermal DC. Data are from one experiment and representative of two similar experiments (SI Appendix, Fig. S6 and Movie S6).

T follicular helper (Tfh) cells (cluster 19), an artifact due to the close proximity of these cells within the GC (SI Appendix, Fig. S6C). Nevertheless, the results presented here demonstrate that IBEX-generated images are compatible with established methods for analyzing high-dimensional imaging data.

IBEX Scales to Capture Ultra High-Content Data from Large Areas of **Human Tissues.** In addition to capturing the cellular landscape of a diverse range of murine tissues, the IBEX method scales to enable high-content imaging of human tissues. To this end, we present an application of the IBEX method to visualize tumor-immune interactions in a human pancreatic LN with metastatic lesions as shown by CD138+EpCAM+ cells in the LN capsule and sinuses (Fig. 5A and SI Appendix, Table S3). Interestingly, cancer cells were segregated from lymphocytes by extensive collagen remodeling and recruitment of myeloid cells expressing the secreted protein acidic and rich in cysteine (SPARC), a matricellular glycoprotein involved in extracellular matrix deposition and implicated in metastasis (34). To move to an even deeper analysis of a much larger sample, we used a total of 66 antibodies for the analysis of a >3-mm² human mesenteric LN section. Using this approach, we were able to deeply phenotype a wide range of immune cells while observing no tissue degradation over 20 cycles (Fig. 5B and Movie S8 and SI Appendix, Table S3). Additionally, we observed subcellular resolution for PC-specific markers (membrane, CD138; nuclear, IRF4; cytoplasmic, IgA1, IgA2) present in distinct cycles with no epitope loss, as evidenced by our ability to label the immune marker CD45 with two different antibody clones present in cycles 9 and 19 (Movie S8). The utility of this method is further exemplified by our ability to characterize the complex stroma of the LN, shown to contain nine distinct clusters in scRNA-seq experiments from mouse LNs (35), using a wide range of antibodies visualized in situ: α-SMA, CD21, CD23, CD34, CD35, CD49a, CXCL12, CXCL13, desmin, fibronectin, and vimentin. These data highlight the capacity of IBEX to identify a large number of distinct cell types in clinically relevant samples, while also placing these components in a spatial setting missed by methods employing dissociated single cells.

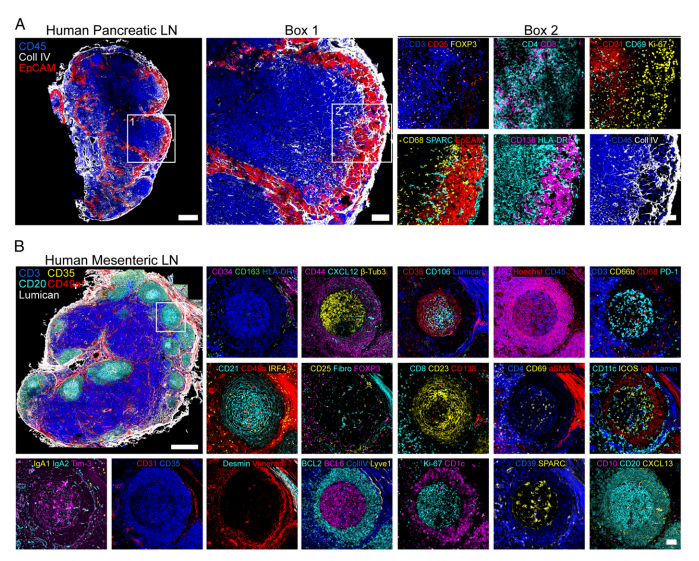


Fig. 5. IBEX scales to capture ultra high-content imaging in human tissues. (A) Confocal images from a pancreatic LN with metastatic lesions from a patient with colorectal cancer (4 cycles, 17 parameters). (Scale bars: Left, 500 μm; box 1, 100 μm; box 2, 50 μm). (B) Confocal images from human mesenteric LN (20 cycles, 66 parameters). (Scale bars: 500 μm or 50 μm.) Fibro, fibronectin; Lamin, laminin. Movie S8 shows additional details.

Extensions of IBEX Workflow. Given the inherent versatility of the IBEX method, we sought to extend our workflow to develop two unique protocols, one that enables detection of low abundance epitopes and another that permits iterative imaging with oligonucleotide-conjugated antibodies corresponding to those used in scRNA-seq experiments. Opal IHC, a method of signal amplification that employs incubation with an unconjugated primary, followed by a horseradish peroxidase (HRP)-conjugated secondary, and deposition of Opal fluorophore in the tissue, is an attractive method for detection of very low levels of specific proteins (36). We first tested this method by staining for endogenous levels of the chemokine CXCL9 in the liver sinusoids of mice (SI Appendix, Fig. S7A), which showed a signal not readily detected with direct or indirect staining methods. Further, because Opal IHC is well described for the imaging of fixed (i.e., FFPE) human tissues (37, 38), we next evaluated whether this method could be expanded upon to achieve multiparameter imaging of tissues from high containment facilities. Due to the extreme fixation conditions required to inactivate select agents such as the Ebola virus (10% formalin for 8 d), the majority of stainable epitopes are lost in these tissues (39, 40). To overcome this significant technical limitation, we developed the Opal-plex method that is based on the IBEX pipeline. Opal-plex extends the usual fluorophore limitations of Opal by combining multiplex Opal IHC with cycles of IBEX-based bleaching to eliminate signal from LiBH₄-sensitive dyes (Opal 570, 650, and 690) while utilizing the LiBH₄-resistant dye (Opal 540) as a fiducial (Fig. 6*A* and *SI Appendix*, Fig. S7*B*). Using this approach, we achieved single-cell resolution of 10 unique markers in heavily fixed mouse LNs (Fig. 6*B* and Movie S9).

We next evaluated whether oligonucleotide-conjugated antibodies, including those used for CITE-seq, are compatible with our IBEX workflow. While immunolabeling with oligonucleotide-conjugated antibodies is well established (9), the use of a large number of commercially available TotalSeq-A antibodies with publicly available oligo-tag sequences, the employment of nonproprietary buffers for hybridization and dehybridization, and the use of a wide spectrum of fluorophore-labeled complementary oligonucleotides provides a truly open-source system with many advantages. In particular, the imaging method described here applies the same antibodies used for scRNA-seq, permitting direct comparison between imaging and CITE-seq datasets while providing a muchneeded spatial context for the cell populations identified. Using this approach, we were able to achieve high-quality tissue staining with

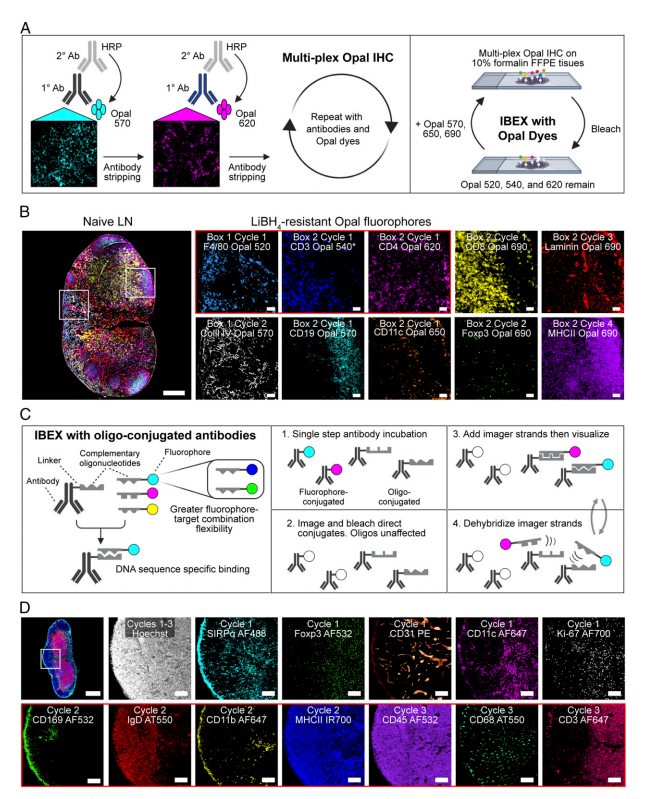


Fig. 6. Incorporation of Opal fluorophores and oligo-conjugated antibodies into IBEX workflow. (A) Opal-plex imaging method. (B) Representative images from a 10-parameter, 4-cycle Opal-plex experiment performed on 5-µm FFPE tissue sections from heavily fixed mouse pLNs. CD3 Opal 540 was present throughout cycles 1 through 4 and served as a fiducial (asterisk). (Scale bars: 50 µm; leftmost panel, 200 µm.) (C) IBEX with oligo-conjugated antibodies method. (D) Confocal images from a 13-parameter, 3-cycle IBEX experiment performed on 20-µm tissue sections from an immunized inguinal mouse LN. Cycle 1: fluorophore-conjugated antibodies. Cycles 2 and 3: oligo-conjugated antibodies, Atto550 (AT550). (Scale bars: 50 µm; Top Left, 400 µm.) Data are representative of three similar experiments (SI Appendix, Fig. S7 and Movie S9).

five unique fluorophores (*SI Appendix*, Fig. S7C). This method can be directly integrated into our IBEX protocol, alongside fluorophore-conjugated antibodies when CITE-seq antibodies to desired targets do not exist, as LiBH₄ bleaching leaves oligonucle-otide binding intact (Fig. 6 C and D). Importantly, the quality of staining achieved with oligonucleotide-conjugated antibodies, even after multiple cycles of LiBH₄ bleaching, is comparable to conventional IF, as quantitative differences, e.g., higher expression of MHCII on dendritic cells (DCs) versus B cells, can still be observed (Fig. 6D and *SI Appendix*, Fig. S7D and Movie S9). In summary, this protocol improves upon existing high-dimensional DNA-based imaging techniques by offering full flexibility in antibody–fluorophore pairing, integrating commercially produced CITE-seq reagents, reducing antibody labeling to one step, and extending the number of fluorophores per cycle.

Discussion

Multiplex imaging of tissues is increasingly important for studies of tumor-immune interactions, for discovery efforts such as the Human Cell Atlas, for better understanding of pathological events in infected or physically damaged tissues, and for placing data from isolated cells in the context of in situ tissue organization. IBEX is a broadly applicable technique that utilizes conventional microscopes and commercially available antibodies to obtain these essential high-dimensional imaging data. IBEX improves upon existing iterative methods by addressing many of the limitations inherent to these techniques. First, we have significantly reduced the fluorophore inactivation step and antibody labeling time from >16 h to <1 h using a rapid chemical bleaching agent and antibody labeling employing a commercial nonheating microwave. Second, our selection of the bleaching agent LiBH₄ provides an efficient means to bleach over 15 unique fluorophores while preserving select fluorophores to serve as repeated markers for registration. Importantly, LiBH₄ treatment does not cause tissue or epitope loss, as evidenced by our ability to obtain highly multiplexed data over several cycles in a wide range of tissues with a very large number of antibodies. Third, and integral to the preservation of tissue integrity through multiple fluid handling cycles, was the use of the tissue adhesive chrome gelatin alum. Importantly, this adhesive adheres delicate tissues to the slide or coverslip surface while maintaining key anatomical features. Finally, the SimpleITK workflow described here represents a significant advancement for the registration of images obtained via cyclic IF methods. In addition to offering flexibility in terms of the repeated markers (membrane, nuclear, structural) used, it provides alignment of markers present on the same cell but not utilized as the fiducial. This is a critical standard for all high-dimensional imaging methods because multiple markers are often required to phenotype a particular cell type, and staining for the relevant epitopes may occur in different imaging cycles.

In addition to developing an efficient method for highly multiplexed imaging, the IBEX workflow, unlike commercial all-in-one systems (9–11), offers flexibility in terms of cellular markers, antibody-fluorophore combinations, and microscope configurations employed. Because the chemistry of bleaching depends on the fluorophore and not the antibody to which it is conjugated, once the bleaching conditions are defined, staining panels can be designed using specific combinations of fluorophores without regard for the target epitopes of the antibodies employed, providing the user with extreme versatility in experimental design. To this end, we report the validation of more than 200 commercially available antibodies conjugated to fluorophores with excitation and emission spectra ranging from 405 nm to 750 nm. Furthermore, we demonstrate that commercially available oligonucleotide-conjugated antibodies can be seamlessly integrated into our IBEX workflow, representing an application of TotalSeq-A antibodies for in situ IHC. Given that the barcode sequences for TotalSeq-A antibodies are disclosed, and a wide range of fluorophore-conjugated oligos are readily available, fluorophore and antibody pairing can be fully customized to match microscope configuration, epitope abundance, and unique tissue characteristics. Taken together, the oligonucleotide-staining method described here provides a completely open method to achieve highly multiplexed IF imaging using the same antibodies employed for flow cytometry and/or CITE-seq, enabling effective cross-referencing of datasets derived from these complementary technologies.

As a proof of concept, we have used the IBEX workflow to examine such issues as the visualization of difficult-to-extract myeloid populations in various tissues, changes in immune cell composition following immune perturbation, and detection of low abundance epitopes. For the first application, we were able to visualize tissue-resident macrophages that are difficult to characterize using other methods such as flow cytometry because of their limited recovery upon enzymatic tissue digestion (12). Using the panels of antibodies outlined here, we were able to deeply phenotype medullary (CD169⁺F4/80⁺CD11b⁺Lyve-1^{+/-}) and subcapsular sinus (CD169⁺F4/80⁻CD11b⁺) macrophages in the LNs (41), as well as alveolar (SiglecF⁺CD11b⁻CD11c⁺) and interstitial (CD11b⁺CD11c⁺MHCII⁺) macrophages of the lung (42). Additionally, we show that the IBEX method can be scaled to capture ultra high-content imaging in human tissues. The ability to survey large areas of human tissue is critically important, as all possible information needs to be extracted to provide maximally useful clinical and research data.

Beyond simple visualization of diverse cell types, we have shown compatibility between IBEX-generated data and downstream single-cell, spatially resolved analysis using the opensource method histoCAT. From the 10-cycle IBEX experiments described above, the histoCAT workflow identified 29 phenotype clusters characterized by the expression of several different markers present in distinct imaging cycles. Importantly, this approach identified well described changes following immunization such as an increase in Tfh cells and GC B cells (43). Finally, incorporation of multiplex Opal IHC into our IBEX workflow facilitated the detection of low abundance epitopes present in conventionally fixed tissues while aiding in the detection of epitopes lost under extreme fixation conditions. The latter represents a significant achievement because few studies have visualized the immunopathology induced by select agents and, to date, the largest number of parameters examined in a single section has been limited to three (44). The ability to use IBEX in its native format and with the Opal-plex variation is especially valuable in the context of the current COVID-19 pandemic. Preliminary data show that these methods work well with highly fixed postmortem samples from such patients.

In summary, IBEX constitutes a versatile technique for obtaining high-content imaging data using conventional microscopes and commercially available antibodies. In addition to providing a valuable resource for studying tissue-based immunity in animal models of disease, ongoing studies have shown the value of the IBEX method to provide a spatially defined assessment of complex cell phenotypes from diverse organs including lung, kidney, heart, and lymphoid tissues from surgical specimens as well as postmortem samples from human COVID-19 patients. We believe that the open nature of the reagents that can be utilized, and the variety of instruments suitable for implementation of IBEX, make it an attractive method for many laboratories seeking to obtain a deeper understanding of cell composition and spatial organization in tissues of interest.

Materials and Methods

Detailed descriptions of animals, immunization and tissue preparations, reagents, imaging protocols with Opal fluorophores and oligonucleotide-conjugated

antibodies, microscopy configurations, and image acquisition and analysis details are reported in SI Appendix, Supplemental Methods.

Mouse and Human Tissues. Murine organs and human LNs (1 cm³ or smaller in size) were fixed with BD CytoFix/CytoPerm (BD Biosciences) diluted in phosphate-buffered saline (PBS) solution (1:4) for 2 d at 4 °C. Following fixation, all tissues were washed briefly (5 min per wash) in PBS and incubated in 30% sucrose for 2 d at 4 °C before embedding in OCT compound (Tissue-Tek). All mice were maintained in specific pathogen-free conditions at an Association for Assessment and Accreditation of Laboratory Animal Care accredited animal facility at the National Institute of Allergy and Infectious Diseases (NIAID). All procedures were approved by the NIAID Animal Care and Use Committee (NIH). Deidentified LN samples were obtained from patients undergoing elective risk-reducing gastrectomies or colon resections for colon adenocarcinoma at the National Cancer Institute (NCI) based on an institutional review board-approved tissue collection protocol (13C-0076).

IBEX Using Inverted Confocal Microscope. Sections (20 to 30 μ m) were cut on a CM1950 cryostat (Leica) and adhered to two-well Chambered Coverglasses (Lab-Tek) coated with 15 μL of chrome alum gelatin (Newcomer Supply) per well. Frozen sections were permeabilized, blocked, and stained in PBS containing 0.3% Triton X-100 (Sigma-Aldrich), 1% BSA (Sigma-Aldrich), and 1%mouse or human Fc block (BD Biosciences). For conventional IF, sections were first blocked for 1 to 2 h at room temperature and then stained for 12 h at 4 °C in a humidity chamber. For microwave-assisted IF, we utilized the PELCO BioWave Pro-36500-230 microwave equipped with a PELCO SteadyTemp Pro-50062 thermoelectric recirculating chiller (Ted Pella). A 2-1-2-1-2-1-2 program was used for immunolabeling, where "2" denotes 2 min at 100 W and "1" denotes 1 min at 0 W. The above program was executed once for blocking and secondary antibody labeling and twice for primary antibody labeling. A complete list of antibodies and tissue-specific panels is provided in SI Appendix, Tables \$1-\$5. Cell nuclei were visualized with JOJO-1 (Thermo Fisher Scientific) or Hoechst (Biotium), and sections were mounted using Fluoromount G (Southern Biotech). Mounting media was thoroughly removed by washing with PBS after image acquisition and before chemical bleaching of fluorophores. Samples were treated with 1 mg/mL of LiBH₄ (STREM Chemicals) prepared in diH₂O for 15 min to bleach all fluorophores except JOJO-1, Hoechst, eF615, and Alexa Fluor 594. To bleach antibodies conjugated to Brilliant Violet 421 (BV421) and Brilliant Violet 510 (BV510) dyes, tissue sections were illuminated using the metal halide lamp with the DAPI filter of the Leica TCS SP8 X inverted confocal microscope. The efficiency of fluorophore bleaching was assessed in real time by viewing the LiBH_a-incubated samples on the microscope. Following efficient bleaching, the LiBH₄ solution was removed, and samples were washed in three exchanges of PBS, restained with the next panel, and mounted with Fluoromount

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G. Fluorophore inactivation with H₂O₂ was conducted as described previously (5), with tissue sections being treated for 1 h at room temperature with 4.5% H₂O₂ (Sigma) prepared in an alkaline solution. Tissue sections were imaged as described in SI Appendix, Supplemental Methods.

Image Alignment and Registration. The alignment of all IBEX panels to a common coordinate system was performed using SimpleITK (23, 24). To facilitate registration, we utilized a common channel present in all panels. As the images may differ by a significant translational motion, we use a Fourier domain-based initialization approach (45) that accommodates this motion. In addition, we utilized SimpleITK's multiscale registration framework with four levels, reducing the resolution by a factor of two per level. Directly using the original voxel sizes, on the order of 10^{-3} mm, in the gradient descent optimizer computations leads to numerical instability. We therefore normalized the voxel dimensions during optimization while preserving anisotropy. The final, optimal transformations are then used to resample all channels from each panel to the common coordinate system. The software repository for SITK_IBEX is provided at https://github.com/niaid/sitk-ibex.

Extensions of IBEX Protocol. Integrating IBEX with Opal dyes or oligonucleotideconjugated antibodies was performed as detailed in SI Appendix, Supplemental Methods. For multiplex Opal IHC, the following steps were repeated to achieve six-plex imaging using the Opal 520, 540, 570, 620, 650, and 690 fluorophores: primary antibody incubation, incubation with HRP-conjugated secondary, labeling with Opal dye, and antibody stripping. After representative images were captured, coverslips were removed and tissue sections were treated with 1 mg/mL of LiBH₄ prepared in diH₂O for 30 min to bleach the Opal 570, 650, and 690 dyes. Cycles of multiplex Opal IHC and IBEX were repeated to achieve the desired number of markers. For integration of oligonucleotide-conjugated antibodies into the IBEX workflow, TotalSeq-A antibodies were incubated with fluorophore-conjugated antibodies, which were imaged first before LiBH₄ bleaching. Oligonucleotides are preserved, and complementary fluorophoreconjugated oligonucleotides are used to reveal immunostaining, then either dehybridized or bleached again in situ across multiple cycles.

Data Availability. All study data are included in the article and supporting information.

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