Spatial mapping of protein composition and tissue organization: a primer for multiplexed antibody-based imaging

John W. Hickey1,2,6, Elizabeth K. Neumann2,3,26, Andrea J. Radtke1,4,26✉, Jeannie M. Camarillo5, Rebecca T. Beuschel8, Alexandre Albanese6,7,25, Elizabeth McDonough8, Julia Hatler9, Anne E. Wiblin10, Jeremy Fisher11, Josh Croteau12, Eliza C. Small13, Anup Sood8, Richard M. Caprioli2,3,14, R. Michael Angelo1, Garry P. Nolan1, Kwanghun Chung6,7,15,16,17,18, Stephen M. Hewitt5 and Sinem K. Saka6,7,15,16,17,18, Stephen M. Hewitt

Tissues and organs are composed of distinct cell types that must operate in concert to perform physiological functions. Efforts to create high-dimensional biomarker catalogs of these cells have been largely based on single-cell sequencing approaches, which lack the spatial context required to understand critical cellular communication and correlated structural organization. To probe in situ biology with sufficient depth, several multiplexed protein imaging methods have been recently developed. Though these technologies differ in strategy and mode of immunolabeling and detection tags, they commonly utilize antibodies directed against protein biomarkers to provide detailed spatial and functional maps of complex tissues. As these promising antibody-based multiplexing approaches become more widely adopted, new frameworks and considerations are critical for training future users, generating molecular tools, validating antibody panels, and harmonizing datasets. In this Perspective, we provide essential resources, key considerations for obtaining robust and reproducible imaging data, and specialized knowledge from domain experts and technology developers.

Mammalian tissues comprise a diverse array of cells that possess unique functional attributes and activation states. Many existing methods—immunohistochemistry (IHC), immunofluorescence (IF), transcriptomics, mass spectrometry—and cytometry—capture this heterogeneity and complexity by studying tissues and organ systems at single-cell resolution. All of these approaches represent a trade-off between spatial information and coverage depth. Flow- or droplet-based single-cell methods, such as multiparameter flow and mass cytometry, single-cell RNA sequencing (scRNA-seq), and cellular indexing of transcriptomes and epigenomes by sequencing (CITE-seq) can define unique cell subsets with incredible granularity. These technologies have greatly expanded our understanding of cell types and states and offer new ways for multiparametric stratification of samples or patients while identifying potential targets for clinical research. However, they typically require tissue dissociation and do not provide a spatial context for cell-to-cell interactions present in normal tissues and altered in disease. Furthermore, these methods fail to retrieve all cell types owing to a combination of factors including, but not limited to, differences in dissociation procedures for individual cell types within a tissue, cell loss during sorting, and low sampling owing to cost or sequencing depth. While recent imaging- or sequencing-based methods probe the spatial transcriptome at single-cell resolution, in situ protein detection overwhelmingly relies on antibodies. Hence antibodies have been at the core of several new multiplexing approaches that allow detection of spatial...
cellular organization and composition of tissues at the protein level\(^{17,18}\) (Supplementary Table 1).

These multiplexed imaging technologies enable detailed interrogation and characterization of cell types of interest—lymphocytes, stromal cells, structural markers—in human tissues beyond the spectral limitations of conventional fluorescence microscopy (typically <5 targets). As the number of protein biomarkers detected via multiplexing increases (typically >10–50 parameters) (Fig. 1a), the assays become increasingly more complex. Therefore, additional effort is required for multiplexed panel design, antibody validation, and careful data acquisition to avoid artifacts while maintaining reproducibility. Future analyses will require robust workflows that yield high-quality images and the computational tools needed to optimally mine these data. Here, we provide a summary of several multiplexed antibody-based imaging approaches and outline strategies for sample and custom reagent preparation, rigorous antibody validation, and multiplex panel building. We then discuss unique challenges surrounding the processing, analysis, and storage of imaging data. Finally, we share a perspective on the future of the field that highlights the utility of these methods and provides practical guidelines for widespread adoption for the methods discussed here.

**Multiplexed antibody-based imaging methods**

Multiplexed antibody-based imaging methods can be classified on the basis of the mode of antibody tagging (such as metal tag, fluorophore, DNA oligonucleotide barcode, or enzyme) and detection...
modality (for example mass spectrometry, spectroscopy, fluorescence, or chromogen deposition), with each approach providing distinct advantages and disadvantages (Supplementary Table 1 and Fig. 1b). Detailed descriptions of these methodologies have been discussed elsewhere\(^1,10,20\), so we focus on the practical aspects of highly multiplexed imaging experiments. Of these antibody-based imaging modalities, fluorescence-based multiplexed imaging is the most established given the wide availability of reagents and imaging systems (Fig. 1b). Typically, fluorescence experiments are limited to visualizing 4–7 antigens within a single imaging cycle owing to spectral overlap of selected fluorophores and availability of labeled commercial antibodies. Hyperspectral methods enable data collection beyond this limit, achieving single-pass multiplexed imaging of up to 21 channels by utilizing fluorophores with diverse excitation and emission spectra, advanced instruments, and methods that enable compensation for spectral overlap\(^4\) (Supplementary Table 1).

Even higher-dimensional datasets can be obtained through an iterative, multistep process (or cycle) that includes (Fig. 1b) (1) immunolabeling with fluorescent or oligonucleotide-barcoded antibodies, (2) direct image acquisition (fluorescent antibody-stained material), or reaction with fluorescent complementary oligonucleotides for a subset of the tagged antibodies, followed by image acquisition, and (3) fluorophore inactivation or removal of antibodies or hybridized oligonucleotide probes. This process circumvents spectral overlap by removing fluorescent signals after each cycle. Using iterative staining, imaging, and bleaching/antibody-removal methods, such as tissue-based cyclic immunofluorescence (t-CyCIF), iterative indirect immunofluorescence imaging (4i), iterative bleaching extend multiplexity (IBEX), multiepitope-ligand cartography (MELC), and multiplexed immunofluorescence (MxF; Cell DIVE), it is possible to detect >60 targets in the same tissue section using off-the-shelf antibodies with fluorescent secondaries or fluorcently conjugated primaries\(^11,12\). Critical considerations for the implementation of such methods are the extended experimental time due to lengthy antibody incubation steps, potential for epitope loss, tissue degradation, and incomplete fluorophore inactivation over successive cycles. For these reasons, it is important to include appropriate controls as outlined here and described previously\(^22–24,29\).

Alternative methods that rely on DNA-barcoding of antibodies, such as DNA exchange imaging (DEI)\(^39\), immunostaining with signal amplification by exchange reaction (immuno-SABER)\(^38\), and co-detection by indexing (CODEX)\(^22,23\), allow one-step immunostaining and fast sequential barcode readout through rapid binding/unbinding of fluorescent oligonucleotides. Although these methods can detect a high number of target molecules in a wide range of tissues, their utility is limited in samples in which harsh fixation has reduced epitope retention or tissues with high autofluorescence. Amplification methods such as Immuno-SABER\(^38\), immunosignal hybridization chain reaction\(^36,39\), and enzymatic (for example horseradish peroxidase\(^36\)) or tyramide signal amplification (TSA)\(^4\) approaches can be used to improve signal-to-noise while increasing the detection of low-abundance epitopes. Until recently, TSA methods such as Opal IHC\(^8\) have been restricted to the simultaneous detection of 6–8 markers\(^8\) as tyramide-linked fluorophores remain bound to the tissue despite rounds of antibody removal and amplification. Nonetheless, it has recently been shown that the fluorophore limitations of Opal IHC could be extended using lithium borohydride to eliminate signal from several Opal dyes, providing a means for the capture of highly multiplexed images in heavily fixed tissues\(^34\).

An alternative to fluorescence-based approaches is mass spectrometry (MS)-based methods that incorporate ionizable metal mass tags (Fig. 1b). These technologies enable visualization of >40 biomarkers in a tissue section using a single master mix of metal-conjugated primary antibodies and do not require antibody staining/removal cycles. The two most common technologies are multiplex ion beam imaging (MIBI)\(^25\) and imaging mass cytometry (IMC)\(^8\), which differ by the use of an ion beam or laser, respectively, for tag ionization. The major benefit of these MS-based technologies is the ability to detect and resolve dozens of metal-isotope-labeled antibodies simultaneously, aptly named ‘all in one.’ The metal ion barcodes possess low background signal by circumventing auto-fluorescence and incorporating high instrumental mass-resolving power. Unlike other multiplexed approaches, samples must be vacuum stable, and antibody labels cannot be amplified. However, the limit of detection of MS systems is on the order of attomoles. The availability of bovine serum albumin (BSA)-free antibody formulations is a limitation shared with other imaging workflows requiring custom reagent creation, for example chelation or fluorophore conjugation. An additional potential barrier associated with expanding metal-tagged antibody panels is access to sufficient amounts of isotopically pure lanthanide metals. While image acquisition occurs in one cycle, it can be slow when considering large fields of view (~200 px/s or 1 mm²/2 h)\(^3\). Finally, to date, the instruments are less operationally stable than conventional fluorescence microscopes and require more specialized staff and environments for effective use.

Other spectroscopy methods that also employ all-in-one data collection include multiplexed vibrational imaging\(^36\) using techniques such as stimulated Raman scattering (SRS) or surface enhanced Raman spectroscopy (SERS). Instead of mass barcodes, these methods utilize the enhanced vibrational signatures of fluorophores and specialized modifiers at different wavelengths for multiplexing over 20 targets. Sample preparation for spectroscopic multiplexed imaging is similar to that for fluorescence microscopy and can be performed on various tissue preparations without a vacuum system.

Beyond investigating tissues in two dimensions (2D), recent advances in sample preparation and imaging have enabled exploration of entire tissue volumes (three dimensions, 3D) to allow a deeper understanding of total organ architecture while permitting detailed characterizations of rare cells that are frequently undersampled in thin (~5–10 µm) tissue sections\(^41\). For instance, chemical clearing methods can be applied to transform intact tissues and organs into robust, transparent samples for future antibody staining and imaging. Importantly, clearing methods must preserve the spatial distribution of proteins and nucleic acids while remaining permeable to fluorescent probes. Several tissue-clearing options can be used, such as: clear lipid-exchanged anatomically rigid imaging/immunostaining-compatible tissue hydrogel (CLARITY), clearing-enhanced 3D (Ce3D), system-wide control of interaction time and kinetics of chemicals (SWITCH), magnified analysis of proteome (MAP), stabilization under harsh conditions via intramolecular epoxide linkages to prevent degradation (SHIELD), entangled link-augmented stretchable tissue-hydrogel (ELAST), and protein retention expansion microscopy (pro-ExM)\(^42–46\). After clearing, tissues have been reported to be amenable to antibody staining and subsequent probe removal using either heat, detergents\(^42,44–46\), oligo-probe dehybridization, or fluorochrome reduction. This allows the potential for higher multiplexed 3D imaging. Owing to the volume of tissue being analyzed, staining and imaging steps are significantly longer than in conventional histology, although the use of electrophoresis-accelerated antibody transport\(^46\) and light-sheet microscopy enables whole murine organs to be stained in a day and imaged within hours. Together, these approaches enable rapid and multiplexed 3D interrogation of intact tissues, providing system-scale structural and molecular information.

Initial experimental considerations

When selecting a multiplexed antibody-based workflow to adopt or implement, we suggest following the mantra: ‘begin with the end in mind.’ The number of samples, whether multiplexed imaging will be a core focus of the outlined work, available budget, and analytical support are key considerations when choosing an approach for spatial profiling of tissues (Fig. 2a). As discussed above, each
multiplexed imaging method has distinct advantages and disadvantages. Therefore, questions about final data requirements, available samples and format, and existing infrastructure must guide the choice of multiplexed imaging method (Fig. 2b and Supplementary Table 1). Similarly, the first and most essential step of creating a multiplexed imaging panel is to determine the scientific questions to be addressed. As all multiplexed imaging modalities share the process of creating and validating multiplexed antibody panels, we focus on the majority of our recommendations on creating a validated panel of 10–60 antibodies (Fig. 3).

Tissue handling. The multiplexed antibody-based imaging methods described here are compatible with a wide range of tissue and sample preparations including fresh frozen (FF), formalin-fixed, or formalin-fixed paraffin embedded (FFPE, Fig. 3). When deciding on a sample preparation method, one should consider the preservation of tissue architecture, long-term storage conditions, epitope accessibility, and ease of multiplex tissue imaging. To this end, FFPE workflows preserve overall tissue architecture and cellular morphology better than do most FF methods.54,55. Additionally, most clinical or archival tissues are stored as FFPE blocks due to enhanced preservation and compatibility with room-temperature storage. However, FFPE preservation renders many target epitopes inaccessible, increases autofluorescence due to formalin fixation56–59, and frequently requires antigen retrieval52,53. In contrast to single-marker IF/IHC, antigen retrieval conditions suitable for one epitope may be incompatible with another epitope, requiring different antigen-retrieval methods for distinct molecules, making multiplex panel development increasingly challenging. In contrast, FF tissues overcome the need for antigen retrieval but instead require immediate snap freezing and ultralow temperature storage (≤−80 °C) for best tissue preservation60–62. Lastly, a tissue-preservation method employing 1% paraformaldehyde and a detergent has been shown to preserve tissue architecture, minimize tissue autofluorescence by lysing red blood cells, and enable immunolabeling with a large range of antibodies without the need for antigen retrieval in a variety of human and mouse tissues63–65.

Costs. An overlooked consideration when designing highly multiplexed imaging experiments is the overall cost and time investment, including that for reagents, antibodies, training, panel optimization, image acquisition, and data analysis (Fig. 3). Ultimately, the cost per unique data point and power of contextual multiplexed data can be useful metrics when rationalizing the increased costs described below. For example, whereas a single-plex experiment can cost a few hundred US dollars, a 50-plex panel could cost many thousands of US dollars for an identical number of tissue samples. Several factors contribute to the substantial increase in time and cost of the panel, including the complexity of the panel (for example number of antibodies, specimen type, sample preparation steps), stability of target epitopes over multiple cycles (if applicable), antibody compatibility and performance under unified immunolabeling conditions, and iterative panel design and adjustment for optimal results. Moreover, identifying and obtaining candidate antibodies can take days to weeks and may additionally require modification of primary antibodies with direct tags. In particular, in situ validation can consume weeks or even months for complex panels applied to a new tissue type where each antibody needs to be validated first in single-plex experiments and then assessed for performance within the entire panel. Not all antibody clones are likely to work across all multiplexed technologies, and each multiplexing technology requires sample-specific antibody titration. MS-based platforms are even more expensive, requiring specialized imaging instrumentation and expensive labeling reagents. Beyond raw materials, training requirements vary widely across different assays and methodologies and can dramatically increase costs. More time is invested in panel development in fluorescence-based approaches than in MS-based approaches because of the cyclic nature of these methods. However, as multiplexed methodologies are commercialized and adopted by core facilities, the time, training, and optimization costs will ultimately decrease.

Target selection. Additionally, it is critical to consider downstream analytical requirements: image registration (for example nuclear labels such as DAPI or Hoechst 33342)66–69, cellular segmentation (for example robust nuclear and membrane stains)70, and unsupervised clustering (for example phenotyping discrete populations using a combination of markers)71,72. By considering these requirements, putative markers are suggested to be ranked and categorized as ‘essential’ or ‘desired,’ greatly expediting overall panel development. This information, along with the budget and approximate timeline, will determine the number of markers to be included in a multiplexed imaging panel.

Marker selection can be guided by expert knowledge, existing literature, orthogonal datasets, and/or online resources (Supplementary Table 2). After establishing a target list of molecules, evaluating and purchasing appropriate antibodies is the next step. In recent years, multiple antibody search engines have been generated, offering intuitive user interfaces that curate antibody clones by citations or artificial-intelligence-based approaches (Supplementary Table 3). In addition to identifying highly cited antibody clones, these platforms allow investigators to query many relevant search fields, such as tissue and cell type, application, company, and host species, while often providing reference images. Notably, these databases do not provide details on performance related to multiplexed imaging, such as epitope stability, steric hindrance, and compatibility with other antibodies. For these reasons, we provide a list of antibody clones previously validated for FF, formalin-fixed, and/or FFPE prepared mouse and human tissues using high-content imaging methods (Supplementary Dataset). Additionally, a recent report outlining tissue-specific panels for highly multiplexed imaging of diverse human tissues is available73. These resources serve as a starting point for creating custom panels and can be tailored to the multiplexed imaging method and question(s) of interest.

For laboratories interested in routine multiplexed imaging, we recommend creating base panels with well-established antibodies that can be expanded as needed. Such an approach reduces the time and capital associated with antibody validation. Moreover, recombinant monoclonal or multiclonal antibodies with appropriate quality-control measures offer the least lot-to-lot variation, making them preferable for highly multiplexed panel development. Beyond that, preference is given to rigorously quality controlled hybridoma-derived monoclonal antibodies and, lastly, to polyclonal antibodies. Applications and sample-specific testing of each vendor lot is recommended to ensure reproducibility across experiments regardless of antibody type.

Creating custom reagents

Many low-plex IHC and IF methods utilize primary antibody detection via a secondary antibody (indirect detection) to enable signal amplification. While convenient, these indirect labeling approaches often cannot be employed to obtain highly multiplexed datasets owing to significant overlap between the primary antibody host species and isotype. Consequently, it is often necessary to modify primary antibodies with functional groups that enable detection (Fig. 4 and Supplementary Table 1). Ultimately, reagent preparation strategies are driven by the cost, availability of materials, technical capabilities, scale, and intended experimental design.

Numerous conjugation methods are used for antibody labeling, each having distinct advantages and disadvantages as well as several tunable variables including, but not limited, to: crosslinker chemistry, crosslinker concentration, pH, modifier concentration,
reaction buffer, and purification strategy. Factors like degree of labeling and conjugate purity/homogeneity might vary depending on the method and adjustable variables. Frequently, conjugation chemistries target lysine, cysteine, or glycan residues of the antibody (Fig. 4), though there are other site-directed approaches that target the amino terminus, defined lysine groups, or the fragment

---

**Fig. 2** | Considerations for the choice and implementation of multiplexed antibody-based imaging technologies into existing workflows. a, Open-source, commercial, and core facility options can be separated by advantages and disadvantages related to ease of implementation, initial cost investment, cost per experiment, flexibility and customization, and the required expertise. b, Several factors govern which method to implement: imaging data requirements (area of tissue needed to be imaged per sample, resolution of final images, time required for imaging each sample, number of markers), sample requirements (number and format of samples, preservation method used for samples, tissue autofluorescence, and whether 2D or 3D volume data are needed), and infrastructure requirements (where existing equipment can be leveraged, level of bioinformatics needed for analysis, technical support, and whether reagents can be purchased or must be customized). Comparisons between different multiplexed imaging techniques are summarized in Supplementary Table 1 and have been described in greater detail in other reviews.
Conjugation chemistries. Lysine-directed conjugations are typically preferred for conventional dyes and oligonucleotide barcodes because they are quick, cost effective, controllable, and scalable. A typical IgG has over 40 solvent-exposed lysines, and the extent of conjugation on these residues enables adjustment of the antibody to modifier ratio. While optimized, the labeling is not site-specific and can result in heterogeneous conjugation and crossreactivity with non-lysine amino acid residues. If critical lysines are in the antibody active site, the modification can reduce epitope binding activity. These factors can make probe standardization and validation more difficult, although recent publications have described site-specific lysine modification, which may address some of these issues.

Alternatively, cysteine-directed conjugations are also fast, cost effective, scalable, and preferred for the addition of metal tags and oligonucleotide barcodes. While cysteine-based conjugation has been shown to generate more homogeneous and reproducible outcomes, there are only 12 available cysteine sites on a typical IgG. In addition to a lower degree of labeling, the proximity of cysteine residues may result in dyes conjugating near one another, which tends to reduce fluorescence due to quenching. Furthermore, the antibody must be mildly reduced with dithiothreitol or 2-carboxyethyl phosphine (tris) prior to conjugation with maleimide-fluorophore/chelator or crosslinkers, which can impact the antibody structure and affinity.

Often, glycan-directed approaches are preferred for small-scale conjugations, when other protocols fail, or when conjugating oligonucleotide barcodes. On average, an IgG contains two glycan sites on the heavy chain. Because of this, glycan-directed conjugations are site-specific, require less optimization of the antibody to modifier ratio, are more reproducible, and avoid binding site inactivation. Unfortunately, the protocol is typically longer and requires enzymatic reactions, which are more costly and difficult to scale.

Standard commercial antibody formulations may not be directly compatible with conjugation chemistries since they often contain sodium azide, bovine serum albumin (BSA), glycerol, and/or cryoprotectants. To facilitate conjugation, we recommend purchasing antibodies in simple buffers such as PBS or performing affinity purification. Further, stocks should be maintained in PBS or borate buffered saline. As each conjugation results in antibody loss, starting with 50–100 µg of the purified antibody is recommended to yield an appreciable amount of product. Although conjugation procedures and methods significantly vary, we visually summarize common approaches (Fig. 4).

Conjugation considerations. Ultimately, the generation and validation of custom antibody reagents is a major bottleneck for developing multiplexed antibody-based imaging panels. Extended experimental planning and validation of conjugated antibodies associated with multiplexed experiments increases the cost and effort compared to their single-plex counterparts. Custom conjugation also introduces additional batch-to-batch variations, especially for the small-scale preparations done in academic research groups. While commercial custom conjugation services and ready-made conjugation kits make it easier to modify antibodies for custom panels, a wider selection of off-the-shelf primary antibodies conjugated with standard sets of dyes, metals, and oligonucleotide barcode libraries will facilitate their greater adoption. Additionally, antibody performance is frequently affected by the choice of metal tag, fluorophore, or oligonucleotide barcode to which it is conjugated (Supplementary Fig. 1). For this reason, antibodies that target low-abundance epitopes should be conjugated to bright fluorophores that do not overlap
with native autofluorescence. Similarly, metal groups and oligonucleotide sequence combinations that have been previously validated for the tissue type are preferred. After custom conjugations, the functionality of the final antibody–conjugate should be confirmed by comparison to the unconjugated version and reassessed with the target assay (Fig. 4). Modified antibodies may also require alterations in tissue blocking and antibody incubation conditions for optimal performance.

Multiplexed antibody panel design and development

Antibody validation. Antibodies must be validated for each workflow as antibody performance depends on tissue type, preservation method, antigen retrieval conditions, final antibody concentration, incubation buffers, and detection method (Fig. 3). While best practices for antibody validation have been described by Uhlen et al., and have been extensively covered before, for multiplexed antibody-based tissue imaging, we recommend considering: (1) tissue and subcellular location of an antibody target, (2) positive and negative tissue controls, (3) native tissue autofluorescence and other types of background, (4) marker compatibility (for example no crossreactivity or steric hindrance between antibodies), (5) confirming antibody specificity through single-plex IF or IHC experiments, and (6) signal-to-noise, particularly for experiments that cannot be amplified (Supplementary Fig. 1). In addition to the antibody-specific search engines discussed here (Supplementary Table 3), regularly updated databases (Supplementary Table 2) can
aid in establishing the location and relative abundance of a marker. Using this information, one can validate antibodies using appropriate tissues with well-documented expression of targeted molecules. Additionally, knockout or knockdown cell lines\(^{24,27}\) or tissues with distinct spatial expression patterns of targets, can be used to further validate the specificity of an antibody\(^{31,36}\). Finally, pathologists can provide invaluable input into the specificity of antibody staining, particularly in the case of rare clinical samples or atypical markers, where artifactual staining may be more difficult to discern. While we provide an overview of antibody validation for multiplexing here, earlier resources provide a more in-depth and focused discussion of the antibody validation process\(^{26,29}\). To increase reproducibility and confidence in the results, we recommend including metadata on antibody validation with published work (Supplementary Table 4).

An additional consideration affecting antibody performance is native tissue autofluorescence, which can vary widely with tissue type, disease state, sample preparation, and fixation method\(^{30,31}\). Autofluorescence can significantly impact the visualization of markers, particularly if these markers are not paired with fluorophores or spectral channels that have higher signal yields. Thus, even if an antibody is validated for its target specificity, its effective use will depend on the anticipated signal in each tissue as compared with tissue autofluorescence, particularly if the label cannot be amplified. Although there is no standard protocol for controlling autofluorescence, we provide strategies for minimizing signal from commonly encountered endogenous fluorophores as a resource to the field (Supplementary Table 5).

**Full-panel validation.** Many of the multiplexed imaging methods discussed here employ high numbers of antibodies incubated simultaneously or iteratively\(^{31,21,25,30}\). Therefore, once the desired marker list is established and antibody specificity validation is complete, it is necessary to validate the full panel, as there may be cross-reactivity between antibodies or spectral overlap from reporters or barcodes within the panel that need to be adjusted (Fig. 3). When antigen retrieval is required for any of the antibodies in a multiplex panel, the same method of antigen retrieval must work for all antibodies in the panel. Thus it is crucial to compare specificity of an antibody within a multipplexed panel to its performance within a single-plex experiment. If an antibody does not perform similarly to this single-plex experiment, further adjustments may be needed (for example optimization of the staining or conjugation conditions, or alternative clones).

Finally, we recommend titrating antibodies over their dynamic range to determine the best signal-to-noise ratio while limiting spectral overlap\(^{31}\). Initial antibody validation and multiplex panel design can be performed in control tissues that are prepared in the same manner as the experimental tissues. The use of control tissue allows precious samples to be reserved for final image collection using the fully validated panel of antibodies. Cell or tissue microarrays (TMAs) are particularly useful for a quantitative assessment of the optimal antibody titrations on the same slide\(^{22,78}\). TMAs, especially multi-organ TMAs, are also recommended for validation of the antibody specificity across multiple tissues in a single staining round. Recently, TMAs were used to validate IMC antibodies and predict biomarkers relevant to disease outcome using AQUA\(^{26,29}\), a software that uses molecularly defined compartments to automatically measure signal intensity with subcellular resolution. For cyclic methods, it is necessary to verify that tissue morphology and target epitopes remain intact during each stain/bleach cycle or the antibody-removal steps. Verification methods have been detailed elsewhere and include (1) comparing antibody performance in single-plex (serial) and cyclic (iterative) imaging experiments\(^1\), (2) varying the order of antibody addition (first cycle versus the fourth cycle), and (3) reimaging the same target across multiple cycles\(^{24}\). If loss of antibody immunogenicity is observed, we recommend placing the affected antibody earlier within the cycle order, substituting with a different clone, or optimizing the cycling conditions such that low-intensity antibodies proceed high-intensity antibodies.

**Reproducibility and data analysis**

One of the greatest challenges facing any scientific method involves reproducibility and rigor of the published conclusions\(^{30}\), particularly for antibody-based approaches that are prone to variable results in the absence of tightly controlled experiments\(^{31}\). Owing to the complexity surrounding multiplexed antibody-based experiments, we encourage the inclusion of detailed validation data in published work or for these data to be deposited as part of publicly available datasets. By mandating and standardizing these processes, we can facilitate appropriate data publishing and use practices, which will undoubtedly improve the experimental rigor and reproducibility. Sharing data on effective antibodies collectively saves an enormous amount of time and resources, but requires extensive, multipronged validation and stable antibody production. We believe that commercial products play a major role in the future adoption and high-fidelity use of multiplexed methods. Access to application-specific vendor validation and quality-control data with detailed information (for example optimal concentration, tested antigen-retrieval protocols, buffer conditions, stability/storage conditions, RRID/clone information), production of monoclonal antibodies with known sequences/epitopes, broader antibody tag options, and flexible formulations (for example conjugation compatible buffers, lyophilized products) will greatly support the field. Publicly available data repositories, such as the HubMAP data portal\(^2\), HTAN data portal\(^3\), The Human Protein Atlas\(^4\), The Cancer Imaging Archive\(^5\), and The Image Data Resource\(^6\), can be leveraged as early domains for sharing and assessing data.

Another large source of variability comes from how and to what extent the raw imaging data are processed. To facilitate this discussion, we propose using predefined data states that can describe any multiplexed imaging set (Supplementary Table 6). Many of these processing and segmentation analyses can be performed using a combination of commercial, freeware, and lab-built software packages. Some of the most widely used software packages include Zen (commercial and freeware, Zeiss), ImageJ (open source)\(^7\), ASHILAR (open source)\(^8\), cytofiks (open source)\(^9\), CellProfiler (open source)\(^10\), CellPose (open source)\(^11\), NAPARI (open source), CODEX Processing and inForm/MAV (commercial, Akoya Biosciences), HALO (commercial, Indica Labs), QUPath (open source)\(^12\), DeepCell (open source)\(^13\), ilastik (open source)\(^14\), Visiopharm TissueAlign/TMA (commercial, Visiopharm), his-tocAT (open source)\(^15\), MCMICRO (open source)\(^16\), Squidpy (open source)\(^17\), CytoMAP (open source)\(^18\), and many others\(^19-100\). Alternative, segmentation-free approaches have also been developed for quantitative analysis of imaging data using pixel-based assignment of subcellular features\(^92,97,99\) as well as classification of irregularly shaped myoepithelial cells\(^101\).

Multiplexed tissue-imaging experiments typically result in sizable files owing to multiple imaging cycles for large regions of interest scanned with tiling and z-stacks. To account for this, workstations or computer servers with extensive memory, high processing speeds, efficient graphic cards, integrated storage, and multicore processors are required for handling these large datasets. Generally, processing pipelines work down the data state levels and are, principally, an accumulation of different software packages that transform the raw data into processed and segmented data. Of note, cell type or functional unit annotations typically require incorporating the opinion of a field expert (for example a pathologist or tissue biologist) which limits the speed, and sometimes the accuracy, of the assignment, as such tasks cannot be easily automated.
Future directions and challenges

Despite the efforts required for more complicated experimental design and thorough validation, multiplexed imaging experiments are important, and often the only way to visualize diverse cell types and physiological states within complex biological systems in situ. For instance, the most recent advances in multiplexed imaging technologies allow over 60 antigens to be stained within a single tissue section; an order of magnitude higher than single-pass experiments performed on the best microscopy systems. Most mammalian systems contain hundreds of different cell types that serve essential functions and are subject to regulation by their spatial position and neighbors. Importantly, disease formation, progression, and treatment response may be governed by unique cellular associations within tissues. All of these factors necessitate highly multiplexed microscopy approaches to understand the complexity of healthy and diseased tissues. In a recent example, >50 proteins were visualized to reveal how distinct cellular neighborhoods orchestrate the spatial organization of the immune tumor microenvironment and contribute to colorectal cancer outcome.

Extending capabilities and reach. Antibodies have long formed the backbone of in situ protein detection, yet they are imperfect tools that require careful characterization and empirical optimization, which makes multiplexed implementation tedious. They are large molecules whose diffusion, target access and binding are highly sensitive to sample preparation conditions, and critical biochemical characterization information, such as antibody protein sequence, stability, exact epitopes, or binding kinetics, is rarely available. Widespread availability of recombinant antibodies, a bigger selection of nanobodies, and alternative probes like aptamers and engineered binders have the potential to make multiplexed antibody panel creation more efficient and lower cost in the future. Also adoption of multiplexable signal amplification methods (as described above) and brighter nano-emitters as alternatives to commonly used organic dyes would improve the detection sensitivity of multiplex experiments.

Further improvements are expected to include improved reproducibility, automated end-to-end workflows, a higher number of markers per experiment, larger imageable areas, and inspiration from outside the field of multiplexed imaging. Of note, a recent study incorporated principles from astronomy to aid in the collection of high-quality datasets from a six-plex TSA-based imaging assay of human melanoma samples. The resulting workflow, AstroPath, offers guidelines for operator-independent field selection and solutions for commonly encountered problems in multispectral imaging including image processing artifacts, improper image segmentation and classification, and batch-effects due to variation in marker intensities. Importantly, these principles are scalable to the high-content imaging assays described here. Another exciting area of development involves computational/automated segmentation of multiplexed images for specific cell types and subtypes to enable tissue-wide assessment of cellular populations. To this end, the HuBMAP consortium has invested significant resources to compare existing cell-segmentation algorithms for their ability to segment membrane and nuclei across multiplexed imaging platforms and defined data-processing pipelines. In contrast to the many methods described for cellular segmentation, our ability to segment bulk functional units, such as glomeruli within the kidney or ventricles in the heart, are not as advanced. While individual cells are functionally important, their active coordination within anatomical structures is critical for overall organ function. Multiplexed microscopy approaches are key for advancing this field, as several markers are often needed to define a structure completely, and automation enables bulk tissues to be analyzed quickly. Finally, effective implementation of these analytical methods in thicker volumes will greatly advance organ-level studies.

Dealing with data: standardization, reproducibility, and storage challenges. While the increased parameter depth is crucial for understanding complex mammalian systems, each cycle or protein biomarker provides additional data, often leading to terabytes of raw microscopy images. As such, there are considerable challenges associated with transferring, storing, and sharing raw data. These datasets present challenges, as multiplexed experiments often require proprietary software, and scale with the number of cycles, markers, and samples captured. The sheer size of datasets alone presents unique challenges for storage and handling. For instance, the raw data for a 10-cycle CODEX experiment (200 tiles, 11 z-stacks, ×20 objective) is >1.5 TB (150 GB/cycle of 3 probes and DAPI). Many databases or data banks will not store data of this size without considerable cost to the authors or journal, discouraging open data sharing. Processing steps such as compression, extended depth of field, and overlap cropping will reduce the file size considerably (~290 GB using the above example), but this involves altering the image, preventing others from reprocessing the original data with improved algorithms and software. Moreover, data analysis and processing steps should be transparent, such that analysis-specific metadata or processing standards are reported to enhance the reproducibility and reliability of the final dataset in line with FAIR data reporting standards. The HTAN consortium has recently provided detailed guidelines and references for components and a structured database for recording of multiplex tissue-imaging metadata. Datasets are most valuable if available in an open or nonproprietary format, such as OME-tiff or BDV, which enables free access to datasets in standardized formats. As an alternative, long-term storage or archiving solutions (cheap but slow) can be utilized for raw data (state 0), whereas higher-level data states are maintained on more accessible storage solutions (more expensive but easier to access) (Supplementary Table 6).

Challenges in managing these large datasets are further compounded by the different approaches and tools used to acquire and process the data. Remedying this requires standardized, nonproprietary formats and common infrastructures for sharing and depositing data. We see value in adopting the open science culture and nationally or internationally funded and managed large data repository structures, such as the HuBMAP data portal and the Cancer Imaging Archive. This endeavor surely requires a concerted effort from all stakeholders including funding bodies, publishers, method developers, biotechnology companies that commercialize these methods into hardware, software, and chemical products, and the scientists who utilize these resources.

From multiplexed imaging to sequencing. As multiplexed imaging approaches are integrated with orthogonal modalities such as flow cytometry, CITE-seq, mass spectrometry, transcriptomics, and spectroscopy, it is crucial to bring the resulting data closer to established ‘omics’ practices. Incorporating the multiplexed antibody recommendations discussed here will improve the characterization of cell types and potentially lead to the discovery of cell (sub)types. In addition to independent validation of new discoveries, such multimodal analysis will enable simultaneous measurement of gene expression and protein products for individual cell types and allow a better understanding of the physiological functions and interacting partners of cells in complex tissues. In addition to the multiplexed imaging methods covered here, new techniques such as DBit-Sequencing and Digital Spatial Profiling offer a spatially defined collection of nucleotide barcodes and enable multiplexed protein detection using sequencing platforms. The best practices described here are also largely applicable to these hybrid methods, although distinct considerations may apply for subcellular sampling and sequencing-based detection of targets.
Conclusion
Here, we have provided an overview for multiplexed antibody-based imaging, collated key resources, and outlined considerations for planning and executing multiplexed protein detection experiments which can provide the highly resolved spatial context that is much needed to create true cell and tissue atlases. Furthermore, these approaches offer direct insights for cell–cell interactions, signaling events, subcellular localizations or translational modifications of proteins, and can complement other bulk assays or single-cell omics analyses of dissociated tissues. By following the suggestions presented here and in cited resources, we aim to empower other investigators to obtain high-quality imaging data using state-of-the-art multiplexed methods. We believe that this primer will facilitate important contributions to biomedical research, and that the emerging findings will be of great impact to the larger scientific community.

Ethics declaration. Human tissues in Fig. 1 and Supplementary Fig. 1 were obtained by following a NIH Institutional Review Board (IRB)-approved protocol (13-C-0076) at the time of risk-reducing surgery performed as a consequence of germline genetic mutation(s). All tissue procured was grossly normal, as determined by the operative surgeon, and histopathologically normal, as determined by a board-certified pathologist.

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

Received: 2 April 2021; Accepted: 15 September 2021; Published online: 22 November 2021

References
1. Stack, C. E., Wang, C., Roman, K. A. & Hoyt, C. C. Multiplexed immunohistochemistry, imaging, and quantitation: A review, with an assessment of Tyramide signal amplification, multiplexed imaging and multiplex analysis. Methods 70, 46–58 (2014).
2. Thul, P. J. et al. A subcellular map of the human proteome. Science 356, eaal3321 (2017).
3. Gawad, C., Koh, W. & Quake, S. R. Single-cell genome sequencing: current state of the science. Nat. Rev. Genet. 17, 175–188 (2016).
4. Massonnet, P. & Heeren, R. M. A. A concise tutorial review of TOF-SIMS based molecular and cellular imaging. J. Anal. Spectrom. 34, 2217–2228 (2019).
5. Neumann, E. K., Do, T. D., Comi, T. J. & Sweetler, J. V. Exploring the fundamental structures of life: non-targeted, chemical analysis of single cells and subcellular structures. Angew. Chem. Int. Ed. 58, 9348–9364 (2019).
6. Zhang, L. & Vertes, A. Single-cell mass spectrometry approaches to explore cellular heterogeneity. Angew. Chem. Int. Ed. 57, 4466–4477 (2018).
7. Porwit, A. & René, M. C. Multiparameter flow cytometry applications in the diagnosis of mixed phenotype acute leukemia. Cytom. Part B: Clin. Cytom. 96, 183–194 (2019).
8. Bandura, D. R. et al. Mass cytometry: technique for real time single cell multistarget immunomassay based on inductively coupled plasma time-of-flight mass spectrometry. Anal. Chem. 81, 6813–6822 (2009).
9. Kolodziejczyk, A. A., Kim, J. K., Svensson, V., Marioni, J. C. & Teichmann, S. A. The technology and biology of single-cell RNA sequencing. Mol. cell 58, 610–620 (2015).
10. Stoeckius, M. et al. Simultaneous epitope and transcriptome measurement in single cells. Nat. Methods 14, 865–866 (2017).
11. Taube, J. M. et al. The Society for Immunotherapy of Cancer statement on best practices for multiplex immunohistochemistry (IHC) and immunofluorescence (IF) staining and validation. J. Immunother. Cancer 8, e001155 (2020).
12. Coltovei, Y. et al. Deep profiling of mouse splenic architecture with CODEX multiplexed imaging. Cell 174, 968–981.e1915 (2018).
13. van den Brink, S. C. et al. Single-cell sequencing reveals dissociation-induced gene expression in tissue subpopulations. Nat. Methods 14, 935–936 (2017).
14. Gernert, M. Y., Kastenmuller, W., Iirim, J., Kabat, J. & Germain, R. N. Histo-cytometry: a method for highly multiplex quantitative tissue imaging application applied to dendritic cell subset microanatomy in lymph nodes. Immunity 37, 364–376 (2012).
15. Asp, M., Bergenstråle, J. & Lundeberg, J. Spatially resolved transcriptomes—next generation tools for tissue exploration. BioEssays 42, 1900221 (2020).
16. Marx, V. Method of the year: spatially resolved transcriptomics. Nat. Methods 18, 9–14 (2021).
17. Uhlen, M. et al. Tissue-based map of the human proteome. Science 347, 1260419 (2015).
18. Uhlen, M. et al. A pathology atlas of the human cancer transcriptome. Science 357, eaan2507 (2017).
19. Tan, W. C. C. et al. Overview of multiplex immunohistochemistry/ immunofluorescence techniques in the era of cancer immunotherapy. Cancer Commun. 40, 135–153 (2020).
20. Bodenmiller, B. Multiplexed epitope-based tissue imaging for discovery and healthcare applications. Cell Syst. 2, 225–238 (2016).
21. Dijkstra, A. J. et al. IBEX: a versatile multiplex optical imaging approach for deep phenotyping and spatial analysis of cells in complex tissues. Proc. Natl. Acad. Sci. 117, 33455–33460 (2020).
22. Gut, G., Herrmann, M. D. & Peklmans, L. Multiplexed protein maps link subcellular organization to cellular states. Science 361, eaar7042 (2018).
23. Lin, J.-R., Fallahi-Sohani, M. & Sörger, P. K. Highly multiplexed imaging of single cells using a high-throughput cyclic immunofluorescence method. Nat. Commun. 6, 8390 (2015).
24. Lin, J.-R. et al. Highly multiplexed immunofluorescence imaging of human tissues and tumors using t-CyCIF and conventional optical microscopes. elife 7, e31657 (2018).
25. Gérardes, M. et al. Highly multiplexed single-cell analysis of formalin-fixed, paraffin-embedded cancer tissue. Proc. Natl Acad. Sci. USA 110, 11982–11987 (2013).
26. Schubert, W. et al. Analyzing proteome topology and function by automated multiparameter fluorescence microscopy. Nat. Biotechnol. 24, 1270–1278 (2006).
27. Wälchly, C., Erlandsson, F., Bengtsson, E. & Zetterberg, A. Sequential immunofluorescence staining and image analysis for detection of large numbers of antigens in individual cell nuclei. Cytometry 47, 32–41 (2002).
28. Zrazhevskiy, P. & Gao, Q. X. Quantum dot imaging platform for single-cell molecular profiling. Nat. Commun. 4, 1–12 (2013).
29. Du, Z. et al. Qualifying antibodies for image-based immune profiling and multiplexed tissue imaging. Nat. Protoc. 14, 2900–2930 (2019).
30. Wang, Y. et al. Rapid sequential in situ multiplexing with DNA exchange imaging in neuronal cells and tissues. Nano Lett. 17, 6131–6139 (2017).
31. Saka, S. K. et al. Immuno-SABER enables highly multiplexed and amplified protein imaging in tissues. Nat. Biotechnol. 37, 1080–1090 (2019).
32. Schürch, C. M. et al. Coordinated cellular neighborhoods orchestrate antitumoral immunity at the colorectal cancer invasive front. Cell 182, 1341–1359.e1319 (2020).
33. Neumann, E. K. et al. Highly multiplexed immunofluorescence of the human kidney using co-detection by indexing. Kidney Int. https://doi.org/10.1016/j.kint.2021.08.033 (2021).
34. Lin, R. et al. A hybridization chain–reaction-based method for amplifying immunosignals. Nat. Methods 15, 275–278 (2018).
35. Wang, Y. et al. Multiplexed in situ protein imaging using DNA-barcoded antibodies with extended hybridization chain reactions. Preprint at bioRxiv https://doi.org/10.1101/274456 (2020).
36. Tsuchikawa, T. et al. Quantitative multiplex immunohistochemistry reveals myeloid-inflamed tumor-immune complexity associated with poor prognosis. Cell Rep. 19, 203–217 (2017).
37. Angelo, M. et al. Multiplexed ion beam imaging of human breast tumors. Nat. Med. 20, 436–442 (2014).
38. Giesen, C. et al. Highly multiplexed imaging of tumor tissues with subcellular resolution by mass cytometry. Nat. Methods 11, 417–422 (2014).
39. Chiang, Q. et al. Imaging mass cytometry. Cytom. Part A 91, 160–169 (2017).
40. Wei, L. et al. Super-multiplex vibrational imaging. Nature 544, 465–470 (2017).
41. Wong, H. S. et al. A local regulatory T cell feedback circuit maintains immune homeostasis by pruning self-activated T cells. Cell, 3981–3997 (2021).
42. Chung, K. et al. Structural and molecular interrogation of intact biological systems. Nature 497, 332–337 (2013).
43. Li, W., Germain, R. N. & Gernert, M. Y. Multiplex, quantitative cellular analysis in large tissue volumes with clearing-enhanced 3D microscopy (Ce3D). Proc. Natl Acad. Sci. USA 114, E7321–E7330 (2017).
44. Murray, E. et al. Simple, scalable proteomic imaging for high-dimensional profiling of intact systems. Cell 163, 1500–1514 (2015).
45. Ku, T. et al. Multiplexed and scalable super-resolution imaging of three-dimensional protein localization in size-adjustable tissues. Nat. Biotechnol. 34, 973–981 (2016).
46. Park, Y.-G. et al. Protection of tissue physicochemical properties using polyfunctional crosslinkers. Nat. Biotechnol. 37, 73–83 (2019).
107. Moore, J. et al. OME-NGFF: scalable format strategies for interoperable bioimaging data. Preprint at bioRxiv https://doi.org/10.1101/2021.03.31.437929 (2021).

108. Nirmal, A. J. et al. The spatial landscape of progression and immunoediting in primary melanoma at single cell resolution. Preprint at bioRxiv https://doi.org/10.1101/2021.05.23.445310 (2021).

109. Liu, Y. et al. High-spatial-resolution multi-omics sequencing via deterministic barcoding in tissue. Cell 183, 1665–1681 (2020).

Acknowledgements

We are grateful for engaging and thoughtful discussions from the Affinity Reagent Imaging and Validation Working Group, HuBMAP Consortium. The authors would like to acknowledge funding from the following sources: NIH U54 DK120058, NIH U54 EY032442, NIH R01 AI145992, NIH R01 AI138581 (R. M. C. and J. M. S.), NIH T32ES007028 (E. K. N.), NIH U54 HG010426-01 (M. P. S. and G. P. N.), NIH UG3 HL145600-01, NIH UH3 CA246633-01 (R. M. A.), NIH UH3 CA246635-01 (N. L. K.), Swedish Research Council 2018-06461 (E. L.), Wallenberg Foundation (E. L.), NIH UH3 CA246594-01 (A. S. and E. M.), NIH T32CA196585 and ACS PF-20-032-01-CSM (J. W. H.), and NIH UH3 CA255133-03 (S. K. S.) and European Molecular Biology Laboratory (S. K. S.). This work was supported, in part, by the Intramural Research Program of the NIH, NIAID and NCI. We thank J. Hernandez and J. Davis (National Cancer Institute, NIH) for providing de-identified human tissues featured in this work, and K. Prummel (EMBL) for comments on the manuscript. Figures were made using the tools on Biorender.com.

Author contributions

J. W. H., E. K. N., A. J. R, J. M. C., and S. K. S. conceived the plan, designed figures, and wrote the manuscript. R. T. B. designed display items and helped write the manuscript. A. A., K. C., E. M., J. H., A. E. W., J. E., J. C., A. S., R. M. C., R. M. A., G. P. N., K. C., S. M. H., R. N. G., J. M. S., E. L., M. P. S., and N. L. K. provided domain expertise and assisted with the conception and writing of the manuscript.

Competing interests

A. E. W. is an employee and shareholder of Abcam plc. J. F. is an employee of Cell Signaling Technologies. J. C. is an employee and shareholder of BioLegend. J. H. is an employee of Bio-Technne. E. S. is an employee of Thermo Scientific. E. M. and A. S. are current or past employees of GE Research. K. C. is an inventor for patent applications covering some technologies described in this paper and a cofounder of LifeCanvas Technologies. G. P. N. is inventor on a US patent, covering some technologies described in this paper, has equity in and/or is a member of the scientific advisory board of Akoya Biosciences. S. K. S. is an inventor for patent applications related to some of the methods described here.

Additional information

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41592-021-01316-y.

Correspondence should be addressed to Andrea J. Radtke or Sinem K. Saka.

Peer review information Nature Methods thanks Trevor McKee, David Rimm and Takahiro Tsujikawa for their contribution to the peer review of this work. Rita Strack was the primary editor on this article and managed its editorial process and peer review in collaboration with the rest of the editorial team.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.
Reporting Summary

Nature Portfolio wishes to improve the reproducibility of the work that we publish. This form provides structure for consistency and transparency in reporting. For further information on Nature Portfolio policies, see our Editorial Policies and the Editorial Policy Checklist.

Statistics

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

- n/a
- Confirmed

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

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

| Data collection | Provide a description of all commercial, open source and custom code used to collect the data in this study, specifying the version used OR state that no software was used. |
|-----------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Data analysis   | Provide a description of all commercial, open source and custom code used to analyse the data in this study, specifying the version used OR state that no software was used. |

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

Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our policy

Provide your data availability statement here:
Field-specific reporting

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

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

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

Life sciences study design

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

- **Sample size**
  - Describe how sample size was determined, detailing any statistical methods used to predetermine sample size OR if no sample-size calculation was performed, describe how sample sizes were chosen and provide a rationale for why these sample sizes are sufficient.

- **Data exclusions**
  - Describe any data exclusions. If no data were excluded from the analyses, state so OR if data were excluded, describe the exclusions and the rationale behind them, indicating whether exclusion criteria were pre-established.

- **Replication**
  - Describe the measures taken to verify the reproducibility of the experimental findings. If all attempts at replication were successful, confirm this OR if there are any findings that were not replicated or cannot be reproduced, note this and describe why.

- **Randomization**
  - Describe how samples/organisms/participants were allocated into experimental groups. If allocation was not random, describe how covariates were controlled OR if this is not relevant to your study, explain why.

- **Blinding**
  - Describe whether the investigators were blinded to group allocation during data collection and/or analysis. If blinding was not possible, describe why OR explain why blinding was not relevant to your study.

Reporting for specific materials, systems and methods

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

| Materials & experimental systems | Methods |
|---------------------------------|---------|
| n/a                             | n/a     |
| ☑ Antibodies                    | ☑ ChiP seq |
| ☐ Eukaryotic cell lines         | ☐ Flow cytometry |
| ☐ Palaeontology and archaeology | ☐ MRI-based neuroimaging |
| ☐ Animals and other organisms   |         |
| ☐ Human research participants   |         |
| ☐ Clinical data                 |         |
| ☐ Dual use research of concern  |         |

**Antibodies**

- Antibodies used
  - The review paper includes a reference antibody list where the clone numbers are included and reference publications or vendors are cited.

- Validation
  - Describe the validation of each primary antibody for the species and application, noting any validation statements on the manufacturer’s website, relevant citations, antibody profiles in online databases, or data provided in the manuscript.

**Eukaryotic cell lines**

Policy information about [cell lines](http://celllines.org)

- **Cell line source(s)**
  - State the source of each cell line used.

- **Authentication**
  - Describe the authentication procedures for each cell line used OR declare that none of the cell lines used were authenticated.

- **Mycoplasma contamination**
  - Confirm that all cell lines tested negative for mycoplasma contamination OR describe the results of the testing for mycoplasma contamination OR declare that the cell lines were not tested for mycoplasma contamination.

- **Commonly misidentified lines**
  - Name any commonly misidentified cell lines used in the study and provide a rationale for their use.

(See [CTAT register](http://ctat.org))
Palaeontology and Archaeology

Specimen provenance
Provide provenance information for specimens and describe permits that were obtained for the work (including the name of the issuing authority, the date of issue, and any identifying information). Permits should encompass collection and, where applicable, export.

Specimen deposition
Indicate where the specimens have been deposited to permit free access by other researchers.

Dating methods
If new dates are provided, describe how they were obtained (e.g. collection, storage, sample pretreatment and measurement), where they were obtained (i.e. lab name), the calibration program and the protocol for quality assurance OR state that no new dates are provided.

Ethics oversight
Tick this box to confirm that the raw and calibrated dates are available in the paper or in Supplementary Information.

Ethics oversight
Identify the organization(s) that approved or provided guidance on the study protocol, OR state that no ethical approval or guidance was required and explain why not.

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

Animals and other organisms

Policy information about studies involving animals; ARRIVE guidelines recommended for reporting animal research

Laboratory animals
For laboratory animals, report species, strain, sex and age OR state that the study did not involve laboratory animals.

Wild animals
Provide details on animals observed in or captured in the field; report species, sex and age where possible. Describe how animals were caught and transported and what happened to captive animals after the study (if killed, explain why and describe method; if released, say where and when) OR state that the study did not involve wild animals.

Field-collected samples
For laboratory work with field-collected samples, describe all relevant parameters such as housing, maintenance, temperature, photoperiod and end-of-experiment protocol OR state that the study did not involve samples collected from the field.

Ethics oversight
Identify the organization(s) that approved or provided guidance on the study protocol, OR state that no ethical approval or guidance was required and explain why not.

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

Human research participants

Policy information about studies involving human research participants

Population characteristics
Describe the covariate relevant population characteristics of the human research participants (e.g. age, gender, genotypic information, past and current diagnosis and treatment categories). If you filled out the behavioural & social sciences study design questions and have nothing to add here, write “See above.”

Recruitment
Describe how participants were recruited. Outline any potential self-selection bias or other biases that may be present and how these are likely to impact results.

Ethics oversight
Identify the organization(s) that approved the study protocol.

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

Clinical data

Policy information about clinical studies
All manuscripts should comply with the ICMJE guidelines for publication of clinical research and a completed CONSORT checklist must be included with all submissions.

Clinical trial registration
Provide the trial registration number from ClinicalTrials.gov or an equivalent agency.

Study protocol
Note where the full trial protocol can be accessed OR if not available, explain why.

Data collection
Describe the settings and locales of data collection, noting the time periods of recruitment and data collection.

Outcomes
Describe how you pre-defined primary and secondary outcome measures and how you assessed these measures.

Dual use research of concern

Policy information about dual use research of concern

Hazards
Could the accidental, deliberate or reckless misuse of agents or technologies generated in the work, or the application of information presented in the manuscript, pose a threat to:

| No | Yes |
|----|-----|
| ☒ Public health |     |
| ☒ National security |     |
| ☒ Crops and/or livestock |     |
| ☒ Ecosystems |     |
| ☒ Any other significant area |     |

Experiments of concern

Does the work involve any of these experiments of concern:

| No | Yes |
|----|-----|
| ☒ Demonstrate how to render a vaccine ineffective |     |
| ☒ Confer resistance to therapeutically useful antibiotics or antiviral agents |     |
| ☒ Enhance the virulence of a pathogen or render a nonpathogen virulent |     |
| ☒ Increase transmissibility of a pathogen |     |
| ☒ Alter the host range of a pathogen |     |
| ☒ Enable evasion of diagnostic/detection modalities |     |
| ☒ Enable the weaponization of a biological agent or toxin |     |
| ☒ Any other potentially harmful combination of experiments and agents |     |

ChIP-seq

Data deposition

☒ Confirm that both raw and final processed data have been deposited in a public database such as GEO.

☒ Confirm that you have deposited or provided access to graph files (e.g. BED files) for the called peaks.

Data access links
May remain private before publication.
For "Initial submission" or "Revised version" documents, provide reviewer access links. For your "Final submission" document, provide a link to the deposited data.

Files in database submission
Provide a list of all files available in the database submission.

Genome browser session
(e.g. UCSC)
Provide a link to an anonymized genome browser session for "Initial submission" and "Revised version" documents only, to enable peer review. Write "no longer applicable" for "Final submission" documents.

Methodology

Replicates
Describe the experimental replicates, specifying number, type and replicate agreement.

Sequencing depth
Describe the sequencing depth for each experiment, providing the total number of reads, uniquely mapped reads, length of reads and whether they were paired- or single-end.

Antibodies
Describe the antibodies used for the ChIP-seq experiments; as applicable, provide supplier name, catalog number, clone name, and lot number.

Peak calling parameters
Specify the command line program and parameters used for read mapping and peak calling, including the ChIP, control and index files used.

Data quality
Describe the methods used to ensure data quality in full detail, including how many peaks are at FDR 5% and above 5-fold enrichment.

Software
Describe the software used to collect and analyze the ChIP-seq data. For custom code that has been deposited into a community repository, provide accession details.
Flow Cytometry

Plots

- Confirm that:
  - The axis labels state the marker and fluorochrome used (e.g., CD4-FITC).
  - The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a ‘group’ is an analysis of identical markers).
  - All plots are contour plots with outliers or pseudocolor plots.
  - A numerical value for number of cells or percentage (with statistics) is provided.

Methodology

| Sample preparation | Describe the sample preparation, detailing the biological source of the cells and any tissue processing steps used. |
|--------------------|--------------------------------------------------------------------------------------------------|
| Instrument         | Identify the instrument used for data collection, specifying make and model number. |
| Software           | Describe the software used to collect and analyze the flow cytometry data. For custom code that has been deposited into a community repository, provide accession details. |
| Cell population abundance | Describe the abundance of the relevant cell populations within post-sort fractions, providing details on the purity of the samples and how it was determined. |
| Gating strategy    | Describe the gating strategy used for all relevant experiments, specifying the preliminary FSC/SSC gates of the starting cell population, indicating where boundaries between “positive” and “negative” staining cell populations are defined. |

- Tick this box to confirm that a figure exemplifying the gating strategy is provided in the Supplementary Information.

Magnetic resonance imaging

Experimental design

| Design type         | Indicate task or resting state; event-related or block design. |
|---------------------|-----------------------------------------------------------------|
| Design specifications | Specify the number of blocks, trials or experimental units per session and/or subject, and specify the length of each trial or block (if trials are blocked) and interval between trials. |
| Behavioral performance measures | State number and/or type of variables recorded (e.g., correct button press, response time) and what statistics were used to establish that the subjects were performing the task as expected (e.g., mean, range, and/or standard deviation across subjects). |

Acquisition

| Imaging type(s) | Specify: functional, structural, diffusion, perfusion. |
|-----------------|--------------------------------------------------------|
| Field strength  | Specify in Tesla                                       |
| Sequence & imaging parameters | Specify the pulse sequence type (gradient echo, spin echo, etc.), imaging type (EPI, spiral, etc.), field of view, matrix size, slice thickness, orientation and TE/TR/flip angle. |
| Area of acquisition | State whether a whole brain scan was used OR define the area of acquisition, describing how the region was determined. |
| Diffusion MRI   | □ Used □ Not used                                      |

Preprocessing

| Preprocessing software | Provide detail on software version and revision number and on specific parameters (model/functions, brain extraction, segmentation, smoothing kernel size, etc.). |
|------------------------|--------------------------------------------------------------------------------------------------|
| Normalization          | If data were normalized/standardized, describe the approach(es): specify linear or non-linear and define image types used for transformation OR indicate that data were not normalized and explain rationale for lack of normalization. |
| Normalization template | Describe the template used for normalization/transformation, specifying subject space or group standardized space (e.g. original Talairach, MN1305, ICBM152) OR indicate that the data were not normalized. |
| Noise and artifact removal | Describe your procedure(s) for artifact and structured noise removal, specifying motion parameters, tissue signals and physiological signals (heart rate, respiration). |
### Statistical modeling & inference

**Model type and settings**
Specify type (mass univariate, multivariate, RSA, predictive, etc.) and describe essential details of the model at the first and second levels (e.g. fixed, random or mixed effects; drift or auto-correlation).

**Effect(s) tested**
Define precise effect in terms of the task or stimulus conditions instead of psychological concepts and indicate whether ANOVA or factorial designs were used.

Specify type of analysis:  
- [ ] Whole brain  
- [ ] ROI-based  
- [ ] Both

**Statistic type for inference**
(See [Ikund et al. 2016](#))
Specify voxel-wise or cluster-wise and report all relevant parameters for cluster-wise methods.

**Correction**
Describe the type of correction and how it is obtained for multiple comparisons (e.g. FWE, FDR, permutation or Monte Carlo).

### Models & analysis

| n/a | Involved in the study |
|-----|-----------------------|
|     | Functional and/or effective connectivity |
|     | Graph analysis |
|     | Multivariate modeling or predictive analysis |

**Functional and/or effective connectivity**
Report the measures of dependence used and the model details (e.g. Pearson correlation, partial correlation, mutual information).

**Graph analysis**
Report the dependent variable and connectivity measure, specifying weighted graph or binarized graph, subject- or group-level, and the global and/or node summaries used (e.g. clustering coefficient, efficiency, etc.).

**Multivariate modeling and predictive analysis**
Specify independent variables, features extraction and dimension reduction, model, training and evaluation metrics.