# Community-wide hackathons establish foundations for emerging single cell data integration

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# **Draft outline**

Full outline described in 01.outline.md as comment if you need to go back to the big picture.

# Figures, Tables and online resources

### **Figures**

- Figure 1: Outline of manuscript (Provided below)
- Figure 2: scRNA-seq + seqFISH main results
- Figure 3: scTargeted proteomics main results
- Figure 4: scNMT-seq main results
- Figure 5: Illustration of partial overlap between studies (at the cells and / or features level)
- Figure 6: Within study cross-validation
- Figure 7: Infrastructure of Giotto package as illustration

### **Tables**

- Table 1: Summary of tasks and methods across all hackathons highlighting methods and common challenges (Provided below)
- Table 2: Glossary of terms, consensus terms will be used throughout the manuscript
- Table 3: Benchmarking single cell data sets
- Table 4: List of single cell analysis software

# **Online resources**

- Online resource 1: Three hackathon datasets (github)
  Online resource 2: R packages with open source reproducible vignettes (12 vignettes)

### **Abstract**

### Introduction

### Comprehensive characterization of biological systems with multiomics

- Single cell community has advanced technologies to enable concurrent processing of biological systems at multiple molecular resolutions
- The lack of prior knowledge and gold standard benchmark naturally leads to a data-driven approach

### New single cell multi omics initiatives:

- Human Cell Atlas (HCA): assess variation in normal tissues
- · Brain initiative and Allen Brain
- Human Tumor Atlas Network (HTAN): Single-cell, longitudinal, and clinical outcomes atlases of cancer transitions for diverse tumor types.

### What bulk multi-omics (e.g. TCGA, ENCODE) have taught us:

- Type of omics that can answer a specific biological question
- The value of open resources for methodological developments
- New hypotheses

# Using hackathons to illustrate analysis standards and challenges for capturing biological information from multi-omics technologies

- Brief overview of our three hackathon studies highlighting state of the art challenges (e.g., spatial transcriptomics, cross-study analysis, epigenetic regulation)
- Challenges include issues with noise and experimental design, Time lag between regulatory levels
  not addressed and many open questions remain (e.g methylation / gene expression), Direction of
  regulation not captured
- We present our findings from hackathon case studies that helped us obtain benchmarks and define a common language for multi-omics

### · Objectives of this paper

 Provide guidelines on tools / data / technologies / methods and needs to model the multi-scale regulatory processes in biological systems for a computational biologist audience

### Outline and messages

- Cellular and molecular regulation is fundamentally multi-scale and captured by distinct data modalities
- Traditional hypothesis-driven multi-omics/view studies only consider one facet of these technologies, but more can be learned through a holistic approach extending into atlases

• We present our findings from hackathon case studies that helped us obtain a broader picture and language

### Outline of the paper:

### Sc spatial

Challenge 1: overlay scRNA-seq onto seqFISH for resolution enhancement Challenge 2: determine signatures of cellular co-localization or spatial coordinates in non-spatial scRNA-seq

### Sc proteomics

**Challenge 1**: lack of overlap between proteins across studies

**Challenge 2**: integrate spatial protein expression analysis

### scNMT-seq

Challenge 1: Definine genomic features
Challenge 2: DNA methylation imputation

Challenge 3: Link epigenetic features / chromatin accessibility to gene expression

### Summary: Common challenges

- Data integration challenges
- From generic towards study specific methods

### Interpretation challenges

- Common language, no black box
- incorporation of contiguous information
- Over summaries and over interpretation

### Benchmarking

- Definition
- Strategy
- Are hackathons the solution?

#### Software infrastructure

- Data management strategies for multi-modal data
- Scalability strategies
- Interpretable visualizations

### **Future directions**

- Upcoming cell atlases technologies
- Unifying analysis goals for new computational methods
  - What do we need for the future of multi omics single cell

### Outputs

- Guidelines
- Glossary of terms
- Reproducible vignettes

Figure 1: Main challenges discussed during our brainstorming sessions from the hackathons.

# Overview of hackathon studies to illustrate multi-omics standards and challenges

Multi-omics technologies provide the unique capacity for full spatial and molecular characterization biological systems. In these analyses, each data modality can uniquely resolve specific biological scales from which complementary data integration techniques can resolve multi-scale interactions. In spite of this potential, computational techniques and benchmark strategies for even pairs of datasets across high-throughput measurement technologies remain an active area of research. The hackathon studies at the Mathematical Frameworks for Integrative Analysis of Emerging Biological Data Workshop and described here were tailored to highlight independent challenges of data integration. First, while spatial molecular profiling is rapidly emerging, these technologies often provide lower molecular resolution than non-spatial counterparts. In these cases, integration strategies that merge these datasets have the promise to enhance the molecular resolution of spatially resolved profiling. To address this challenge, We designed a hackathon using spatially resolved transcriptional data from seqFISH with corresponding non-spatial single cell profiling data for the mouse visual cortex from [1]. A second challenge for multi-omics technologies in cases for which tissue availability limits the ability to obtain multiple measurements in samples from identical conditions. This limitation raises the question as to whether information can be transferred from datasets between disparate sample cohorts. Therefore, we designed the second hackathon to contain two triple negative breast cancer cohorts profiled with single cell proteomics profiling from mass cytometry (CyTOF) [3] and spatial insitu proteomics from Multiplexed Ion Beam Imaging (MIBI) [4]. While each of these hackathons use data at the same molecular scale, genetic and epigenetic alterations to DNA further drive the transcriptional regulation that mediates intra- and inter-cellular signaling processes underlying cellular fate transitions and states. Our third hackathon uses scNMT-seq data to obtain concurrent DNA methylation, chromatin accessibility, and RNA expression from the same cells to delineate the regulatory networks that underlie mouse gastrulation [5]. Altogether, exploration of the analysis approaches employed to address these disparate hackathons across biological contexts provides an unique opportunity to identify technology-specific challenges and unifying themes that are essential to effectively employ multi-omics datasets into new biological knowledge.

### scRNA-seq + FISH as a case study for spatial transcriptomics

### Overview and biological question

The first hackathon is designed to leverage the complemenatary strengths of sequencing and imaging based single-cell transcriptomic profiling with comptuational techniques to integrate scRNA-seq and seqFISH data in the mouse visual cortext. While single cells are considered the smallest units and building blocks of each tissue, they still require proper spatial and structural three-dimensional organization in order to assemble into a functional tissue that can exert its physiological function. In the last decade single-cell RNA-seq (scRNA-seq) played a key role to capture single cell gene expression profiles, which allowed us to map virtually all the different cell types and states in whole organisms. Despite this remarkable achievement this technology is based on cellular dissociation and hence does not maintain spatial relationships between single cells. More recently technologies have begun to emerge that can profile the transcriptome of single cells within their original environment. These technological advancements offer the possibility to examine how gene expression is influenced by cell-to-cell interactions or organized in a spatially coherent manner. One such approach is sequential single-molecule fluorescence in situ hybridization (seqFISH), which can identify single molecules at (sub)cellular resolution with high sensitivity. Nevertheless, and in contrast with scRNAseq, seqFISH [6] and many other spatial transcriptomic technologies often pose significant technological challenges and hence the number of profiled genes per cell is usually restricted to a smaller number (10-100s). Although the newer generation of seqFISH technology (called seqFISH+)

has greatly enhanced its capacity which can now be used to profile 10,000 genes [8], the technology is significantly more complex and costly. As such, it is desirable to develop computational approaches to effectively integrate scRNAseq and seqFISH data analyses. Therefore, this hackathon provided with seqFISH [2] and scRNAseq [1] data corresponding to the mouse visual cortex and challenged to accurately identify cell-types by integrating both datasets. The scRNA-seq data in this hackathon provides transcriptional profiles at a high molecular resolution whereas the seqFISH data yield spatial characterization at a lower molecular resolution that were divided into two key challenges to enable high-resolution spatial molecular resolution. In this first challenge, this hackathon explored a number of strategies to identify the most likely cell types in the seqFISH dataset based on information obtained from the scRNAseq dataset. Likewise, the second challenge sought to go in the opposite direction to transfer spatial information obtained from the seqfish dataset to that of the scRNAseq dataset.

Elana - revised to streamline flow with other sections. Consider cutting the remaining text ~Single cells are considered the smallest units and building blocks of each tissue, but they still require proper spatial and structural three-dimensional organization in order to assemble into a functional tissue that can exert its physiological function. In addition, most tissues are composed of multiple cell types whose identity and function can be inferred through their unique transcriptomic profile. In the last decade single-cell RNA-seq (scRNA-seq) played a key role to capture single cell gene expression

profiles, which allowed us to map virtually all the different cell types and states in whole organisms.

Despite this remarkable achievement this technology is based on cellular dissociation and hence does not maintain spatial relationships between single cells.~

~More recently technologies have begun to emerge that can profile the transcriptome of single cells within their original environment. These technological advancements offer the possibility to examine how gene expression is influenced by cell-to-cell interactions or organized in a spatially coherent manner. One such approach is sequential single-molecule fluorescence in situ hybridization (seqFISH), which can identify single molecules at (sub)cellular resolution with high sensitivity. Nevertheless, and in contrast with scRNAseq, seqFISH and many other spatial transcriptomic technologies often pose significant technological challenges and hence the number of profiled genes per cell is usually restricted to a smaller number (10-100s).~

~The first hackathon is designed to explore comptuational techniques for integration of a matching scRNAseq and seqFISH dataset in the mouse visual cortex that can overcome the lack of spatial information with scRNAseq and the common limited coverage in spatial datasets. More specifically, this hackathon explored a number of strategies to identify the most likely cell types in the seqfish dataset based on information obtained from the scRNAseq dataset. And in the opposite direction, we sought out how to transfer spatial information obtained from the seqfish dataset to that of the scRNAseq dataset.~

![Overview of analysis. *note: letters order can be changed* **A** A spatial network was built from cells' positions using Voronoi tessellation, whilst cell types were inferred from an SVM trained on scRNAseq data. Left: The neighbors aggregation method described in [challenge 2] computes aggregation statistics such as mean or standard deviation on the seqFISH gene expression data for each node and its first order neighbors. Right: This approach identified spatially coherent areas that can contain one or several cell types, and can be used to detect genes whose expression is modulated by spatial factors rather than cell type. B\*\* Assessment of cell type prediction using different data normalizations and classifiers. Normalization strategies included none (raw), counts per million (cpm), ComBat batch correction applied to cpm (cpm\_combat), scRNAseq and seqFISH scaled using the first eigenvalue (cpm\_eigen), latent variables retained for both datasets after applying partial least squares to cpm\_eigen normalized data (cpm\_pls). A supervised multinomial classifier with an elastic net penalty (enet) was trained using the scRNAseq data and cell-type labels (provided). Each model was then applied to the seqFISH data to determine the predicted cell-type labels. A semi-supervised

multinomial classifier with an elastic net penalty (ssenet) was trained using the scRNAseq and seqFISH data and cell-type labels (provided) to impute the "missing" cell-types labels of the seqFISH dataset. The gower distance between each method-normalization pair was computed (including SVM-based predictions from [2]). The multidimensional scaling plot depicts the similarity between different methods and normalizations based on predicted cell-type labels. The first dimension (Dim1) separates methods that normalize the scRNAseq and seqFISH data together (dashed) and separately (solid). The normalization method used had a stronger impact on cell-type predictions than the classification method used.

**C** Challenge 1: LIGER was applied to combine spatial and single cell transcriptomic datasets. From the separate and integrative analyses, plots of identified and known clusters were generated and metrics of integration performance were compared in context showing some loss of information as a result of the integration. **D** Challenge 2: what is the minimal number of genes required for data integration? SVM classification models (C=1e-6, C=1e-3, C=1) were trained and evaluated with different number of genes in scRNAseq data using Recursive Feature Elimination (RFE). The results show that a small (*do you mean smaller that the original study?*) gene list to identify cell types in both data types, but the lack of goad standard hinders our ability to evaluate the relevance of such genes. **Hang** .] (images/seqFish\_mockup.png){#fig:spatial}

### **Computational challenges**

### Challenge 1: overlay of scRNA-seq onto seqFISH for resolution enhancement

### [suggestion 1]

Sequencing and imaging based single-cell transcriptomic profiling have complementary strengths. Whereas single-cell RNAseq generates transcriptome-wide information, it does not have spatial information. On the other hand, seqFISH (Lubeck 2014; Shah 2016) provides single-cell resolution spatial information, but typically profiles the expression level of only 100-300 genes. Although the newer generation of seqFISH technology (called seqFISH+) has greatly enhanced its capacity which can now be used to profile 10,000 genes (Eng 2019), the technology is significantly more complex and costly. As such, it is desirable to develop computational approaches to effectively integrate scRNAseq and seqFISH data analyses.

In this hackathon, the participants were provided with seqFISH (Zhu 2018) and scRNAseq (Tasic 2016) data corresponding to the mouse visual cortex and challenged to accurately identify cell-types by integrating both datasets. Cell type labels, derived from scRNAseq analysis (Tasic 2016) and previous seqFISH/scRNAseq integration (Zhu 2018) were also provided as reference. A variety of computational approaches were applied to achieve this goal, including: supervised classification with support vector machines (Coullomb, Xu), supervised and semi-supervised (self-training) elastic net classifiers (Singh), and unsupervised matrix factorization methods (Sodicoff) (Figure ??). While the methodologies are different, a number of themes recur, such as the importance of gene selection and batch effect correction. As expected, the ability to identify refined cell-type structure relies on the selection of cell-type specific marker genes in seqFISH data, suggesting a potential benefit of using single-cell RNAseq data to guide seqFISH experimental design. Batch effect is another important factor affecting the accuracy of data integration. While a number of batch effect correction methods have been developed (COMBAT, Seurat, Scanorama, etc), it remains challenging to distinguish technical from biological variations if the biological samples do not match exactly.

### [suggestion 2]

The mouse visual cortex consists of multiple complex cell types, however the number of profiled genes in the seqFISH dataset is limited to 125 genes. Moreover, these genes were not prioritized based on their ability to discriminate between cell types and thus assigning the correct cell identity is challenging. In contrast the scRNAseq dataset is transcriptome wide, thus including the 125

aforementioned genes. As such we first used all genes to identify the cell type labels for each cell in the scRNAseq data with high certainty. Next, we leveraged that information to build a classifier based on (a subset of) the 125 common genes only. This classifier could subsequently be applied to the seqFISH dataset in order to assign cell types to each cell with high probability.

During the hackathon we tested various machine learning or data integration models, but also noted that initial dataset normalization strategies might have a significant impact on the final results (see (Figure ??B). Although unique molecular identifier (UMI) based scRNAseq and seqFISH can both be considered as count data, we observed dataset specific biases that could be attributed to either platform (imaging vs sequencing) or sample specific sources of variation. Here we opted to apply a quantile normalization approach which forces a similar expression distribution for each shared gene.

The hackathon participants selected two machine learning classifiers, a supervised support vector machine (SVM) and, a supervised and semi-supervised generalized linear model regularized with an elastic net penalty (enet and ssenet). To further improve the SVM model the participants tried multiple kernels and searched for optimal hyperparameters using a combined randomized and zoomed search. In addition, different flavors of recursive feature elimination was used to find the optimal or minimum number of genes needed to correctly classify the majority of the cells. Importantly, participants tested different classification accuracy metrics to alleviate the major class imbalance in the dataset, since more than 90% of cells were excitatory or inhibitory neurons. The ssenet approach used a iterative model building approach, which combines both datasets and initially only retains the highest confidence labels and then gradually adds more cell type labels until all cells have been classified. This type of self training approach might in fact be more generalizable to other datasets.

Finally, a data integration approach based on integrative non-negative matrix factorization (NMF) was used by applying the previously published LIGER approach. By integrating both datasets in a similar subspace based on shared factors, cell type labels can be transferred using a nearest neighbor approach.

# Challenge 2: Identifying spatial expression patterns at the tissue level through integration of gene expression and spatial cellular coordinates

### [suggestion 1]

How could one identify spatial patterns in the seqFISH data? This broad question can be divided into a number of specific tasks, such as detecting genes whose expression is spatially coherent, cell types whose spatial distribution is confined to distinct regions, recurrent multi-cell-type interaction clusters, etc, and the spatial scale may vary from subcellular all the way to tissue-wide organizations. While there have been abundant studies in the geo-spatial analysis domain, computational tools targeting specific spatial transcriptomic questions are still lacking. In this hackathon, one group tackled this challenge by aggregating gene expression data from neighboring cells followed by spatial clustering (Coullomb). Much more work in the future is need to further explore such information.

### [suggestion 2]

Most analyses that were originally developed for scRNAseq data can be immediately applied to spatial transcriptomic datasets, however methods to extract sources of variation that originate from spatial factors are still sparse. To incorporate spatial information the cells from the seqFISH dataset were first connected through a spatial network based on Voronoi tessellation and then the expression of each individual cell was spatially smoothed by calculating the average gene expression levels over all the neighboring cells. This smoothened and aggregated data matrix was subsequently used to create a 2 dimensional UMAP from which clusters were identified through a density based clustering approach. The obtained cluster labels can then be mapped back to the original spatial locations for further visual inspection and analysis.

### [suggestion 2 detailed]

Most analyses that were originally developed for scRNAseq data can be adapted for spatial transcriptomic datasets, however methods to extract sources of variation that originate from spatial factors are still sparse. The latter requires the development of novel methods which can integrate the information obtained from gene expression with that of the spatial coordinates from each individual cell or transcript (for sub-cellular resolution) within a tissue of interest. For this hackaton we aimed to incorporate the spatial information by connecting the cells from the seqFISH dataset through a spatial network based on Voronoi tessellation. Then, for each node, its RNA count data was aggregated with its first order neighbors' count data. The mean and standard deviation were computed for each gene in the gathered data in order to capture the global tendency as well as the variability in the area arround each node. Thus, each node has nb\_genes x nb\_statistics (here 2) variables. These "aggregation statistics" can be visualized on a 2D UMAP projection. These data were clustered in UMAP reduced spaces of dimensionality between 2 and 9, higher dimensions allowing to define more fine-grained clusters. The clustering was performed with HDBSCAN, a noise-aware density-based algorithm that can define arbitrary-shaped clusters. These clusters can then be visualized on the 2D UMAP projection and on the 2D spatial map of seqFISH data. The clusters are spatially coherent, some of them contain several cell types, and a given cell type is not necessarily limited to one specific cluster. During the exploratory phase consisting in varying the number of dimensions and the minium cluster size, a specific spot area was found clustered for several parameters combinations, suggesting it wasn't an artifact of the choice of parameters. "Differential expression" analysis was performed between this spot and the other areas, although we don't look at differences in gene data but in aggregation metrics. This area seems to correspond to a "regeneration hub", but this analysis has to be considered carefully and further analyzes is required to confirm this hypothesis. This "neighbors aggregation" method has been extended to aggregate RNA counts (or other node attributes) to higher orders of neighbors in order to define aggregation metrics on wider areas, which could be useful for analyzes of bigger tissues. One interesting extension would be to substract phenotypes contributions to RNA counts for each cell before performing the neighbors aggregation analysis in order to highlight genes that are modulated by spatial factors. But if we want to retrieve the mean expression of a cluster for cells belonging to it, we should first check cluster's convexity and be sure that no other cluster lies within it.

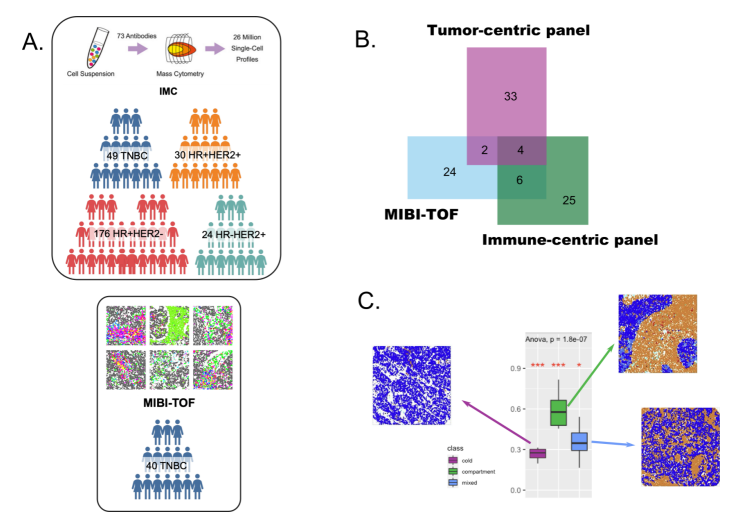
# Spatial proteomics and cross-study analysis

### Overview and biological question

The single-cell targeted proteomics hackathon investigated the tumor-immune microenvironment of primary breast cancer tissue. The challenge was the joint cross-study, cross-platform integrative analysis of single cell proteomics data that had low feature overlap, were generated on different antibody-based targeted proteomics technological platforms and in different laboratories [1, 2]. Mass cytometry (CyTOF) from Wagner et al. [3] measured 73 proteins in two panels (immune, tumor) in 194 tissue samples from 143 subjects, of which 6 patients had triple-negative negative breast cancer. The second dataset applied Multiplexed Ion Beam Imaging (MIBI) to quantify spatial in-situ expression of 36 proteins in 41 triple-negative breast cancer patients [4]. 2 Whereas the first hackathon had matched seqFISH and and scRNA-seq data of samples from the same biological conditions, the data in this hackathon is obtained from different cohorts of patients posing a further challenge for analysis.

Whilst this is a formidable data integration challenge, it reflects the bioinformatics analysis of clinical teams who wish to compare and investigate data collected on different measurement platforms and cohorts of cancer patients. The first questions simply asked if partially-overlapping proteomic data collected on different patients with similar phenotypes could be integrated, and asked participants to test if measurements in one technology could be transferred and used to predict information in the second. The MIBI data provided spatial location and expression of proteins, therefore the challenge

asked if the spatial expression patterns of proteins measured on mass-tag could be predicted. Secondly it raised questions unique to spatial 'omics technologies. Does spatial capture unique information, beyond cell compositions. Could information about the spatial location of immune cell populations in breast cancer be discovered in integrated analyses of these datasets. Finally the datasets had no overlap in patients, so how could heterogeneous phenotype information be used to integrate patient 'omics data with low feature and no tumor biological sample overlap.



**Figure 2:** A Overview of sc\_targeted proteins hackathon challenge. **B** Challenge 1: Partial to no overlap between protein features across studies. **C** Challenge 2: Spatial analysis with Moran's index, a computed on Gabriel graph (using both dummy variables and protein expression measurements) that differs significantly between groups. Boxplot of Moran's index values on tumor/immune status with examples from Keren et al. [4] corresponding to each tumor category. Red asterisks indicate significance of an ANOVA of each group with all others, and the p-value from an overall ANOVA across the three groups is reported above.

### **Computational challenges**

### Challenge 1: Lack of overlap between protein features across studies

The low number of features precluded integration of features at the level of gene set or pathways. There were only 20 proteins that were assayed in both studies (Figure 2A-B). The majority of features were cell-type markers or biomarkers targets of breast cancer therapeutic intervention. The limited overlap in these studies necessitated the use of surrogate measures of cross- study association.

Although the overlap in proteins was low, many proteins were cell-type markers, providing the opportunity to perform cross-study integration of cell type proportions in tumor tissue samples. Hackathon participants applied several semi-supervised and supervised algorithms to transfer cell labels and cell compositions from one dataset to the second. To capture the hierarchical structure of

cell lineage, Lauren Hsu (Harvard) applied a simple random forest (RF) approach to perform feature transfer learning of cell type labels. An adaptation of the prediction strength approach described in Tibshirani [9] demonstrated model robustness: first, a model was trained on the labeled dataset and used to predict labels in the unlabeled dataset; next, a second model was trained based on the second dataset with the newly predicted labels; finally, she assessed the ability of the second model to recover the correct original labels when making predictions on the labeled dataset.

Yingxin Lin (Univ of Sydney) mapped the cells from CyTOF to imaging with spatial information by solving an entropic regularization optimal transport problem [4, 11], utilising the cosine distance of the common proteins between the two datasets as transport cost. The constructed optimal transport plan can be considered as likelihood of cells from one modality mapped to cells from the other modality, which allows the prediction of protein expression measured only in CyTOF on imaging data. By clustering on the imputed expression matrix, she was able to identify a sub tumour cell type that is not revealed in the original matrix.

However the scales of protein expression was a possible limiting factors When integrating cell compositions using the correlation expression of protein markers, some cell markers were expected on a range of cell types (e.g., CD45), whereas others are more specialized and represent a subset of those cells (e.g., CD4). Others challenges associated with cell compositions analysis of proteomics analysis included uncertainty over antibody specificity and consistency between studies, the sensitivity and specificity of protein markers for cell types, tissue and disease heterogeneity

The assignment of cell type relied on manually curated protein annotation, and was dependent on domain-specific knowledge; for example that CD4 is expressed by T-cells. To date, methods for cell type assignment, classification or extraction of differentially expressed proteins cannot easily be applied to targeted proteomics. Participants expressed a need for a unifying map between cells present in different datasets, and for annotation resources to provide quality metric or priors of protein cell type markers. There is a need for protein expression atlases to support cell type classification and potentially if this could be developed from large scale consortiums IHC of proteins (Human Protein Atlas [5, 6]), although the antibodies used and their performances might vary between labs.

Standards/QC/Normalization

### Challenge 2: spatial protein expression analysis

Whereas the CyToF mass spectrometry provided protein expression and counts/composition of cells in breast tumor-immune environment, the MIBI-TOF data provided spatial information that quantified cell attributes (shape/size/spatial coordinates) in addition to expression levels, thus providing the opportunity to examine protein expression, cell microenvironment, and predict cell-cell interactions and the cellular community ecosystem.

Spatial information can be encoded as a set of XY coordinates (cell centroid), a line (eg tumor-immune boundary) or a polygon, which is a closed plane defined by a number of lines. A polygon can define complex shapes such as a cell or a community of cells. Spatial protein expression can be summarized using spatial descriptive statistics, such as the autocorrelation of the expression of a protein within a neighborhood of polygons. The neighborhood of polygons can be defined with a Euclidean distance or sphere, by a number of bounded cells or other measures, many of which were developed in geographical information science or ecology and assess if a spatially measured variable has a random, dispersed or clustered pattern [7].

Kris Sankaran examined the extent to which expression data could be used to predict spatial properties of tissue samples. To build predictors, cells were first clustered (K = 20) on the basis of

protein expression. Sample-level expression summaries were defined as the proportions of cells belonging to each cluster. To build the spatial response variables, a K-nearest neighbor graph was obtained from cell centroids (K = 5). For each cell, the average distance to its 5 nearest neighbors was computed, reflecting its local density. Further, the entropy of the cluster memberships across nearest neighbors was found, reflecting local heterogeneity. To summarize samples' cell ecosystems, cell-level statistics were averaged across each sample's cells. A random forest model trained from expression to spatial predictors achieved an average cross validation RMSEs of (tk) for neighborhood size and entropy, respectively, relative to baselines of () obtainable by predicting the mean.

Dr. Pratheepa Jeganathan applied topic modelling and defined five topic trained on protein expression and cell compositions in the CyToF data were sufficient to predict cell co-locations, in 10% MIBI-ToF Test data. Pratheepa Jeganathan (Stanford) applied a Bayesian modelling approach based on latent Dirichlet allocation (Blei, Latent dirichlet allocation Journal of machine Learning research 3.Jan (2003): 993-1022). Topic modeling was used to identify the dominated topics and assign spatial location of MIBI-TOF cells to the CyTOF data or vice-versa, based on the topic distribution in each cell (**Ref topic modelling?**). Among the five topics identified, the first topic was dominated in most of the immune cells from CyTOF data and the other four dominated in all other cells. Cells from MIBI-TOF were depicted in five clusters (link to vignette) and were consistently based on the observed and predicted marker expression, but these clusters were not identified with only observed marker expressions. [further details from Pratheepa available in pdf file in debrief folder]

Yingxin Lin (Sydney) examined the prognostic performance of different higher level spatial metrics. She measured protein autocorrelation using Moran's Index (I) with a sphere distance, cell type localisation using nearest neighbour correlation, or cell type interaction composition, Ripley's L-function. High-dimensional Cox models with fused lasso penalty and random forest survival models were fitted utilising different features, including clinical features such as tumour stage, tumour grade, age and tumour size, as well as features like cell type composition. Evaluating by the c-index via cross-validation, the spatial metrics are found to be predictive, especially in triple negative breast cancer where clinical features such as grade are poorly prognostic.

Lauren Hsu (Harvard) also considered Moran's I but used a graph-based neighborhood measure (Gabriel graph, based on Delaunay triangulation) instead of a sphere euclidean distance, and found that Moran's I differed significantly between the three prognostic tumor scores described by Keren, et al. [4](Figure 2C).

Need for development of spatial measure - different in dimensions of RNA v proteins

### **Challenge 3: Fourth corner Integration of data (at the level of phenotype)**

Another question often faced in cross-study integration is integration of biological samples that are non-overlapping but have similar phenotypes. The aim is to identify biomarkers from the different omics data to predict the same phenotype, and, more importantly, to explore how the markers selected from multiple datasets are in agreement with or distinct from each other. The integration of the markers from each dataset should enable to extend biological knowledge that is not available by single omics data. To solve the challenge, phenotypical data (such as the cell attributes) are the critical factors that should be used to link the two datasets (Figure 2D).

The participants were successful at data integration using patient phenotype measures such as grade, stage and overall survival. Breast cancer is highly heterogeneous, and multiple breast cancer molecular subtypes have been described [8, 9]. Both MIBI and Jackson data used different approaches to cell type annotation and had 13 proteins in common.

Borrowing from ecology and french school of ordination, Chen Meng (Munich) described this problem as a case of the fourth corner problem (or RLQ). Briefly, given two omics data where both rows (features) and columns (samples) are non overlapping, and phenotypical data available for each omics data, multiplying the two phenotypical factors will derive a bridging matrix that links the features of two omics data. We should note that the two phenotypical matrices need to be multipliable, i.e. the phenotypical data should describe the same phenotypical factors over the samples in the corresponding dataset. The Chessel fourth corner RLQ is a matrix decomposition method to solve the problem [10; doi:10.1111/ecog.02302]. It decomposes the bridging matrix (phenotypical matrix) into components, each of which often represents a specific phenotypical pattern in the data. The loading matrix of each of the omics data indicate how a feature is correlated with phenotypical factors.

### scNMT-seq as a case-study for epigenetic regulation

### Overview and biological question

The first two hackathons leveraged paired datasets from complementary technologies that measure the same molecular modality at distinct resolutions. The aim was to develop integration strategies to gain a holistic view into and enable high molecular and spatial resolution of biological systems. Datasets spanning disparate molecular scales, such as DNA and RNA level measurements, are critical to further resolve the regulatory networks that mediate cell fate decisions. For example, the maturation of single-cell sequencing technologies has enabled the identification of transcriptional profiles associated with lineage diversification and cell fate commitment[11]. Yet, the accompanying epigenetic changes and the role of epigenetic layers in driving cell fate decisions still remains poorly understood[12].

scNMT-seq is one of the first experimental protocols that enable simultaneous quantification of RNA expression and epigenetic information from individual cells[13]. Briefly, in scNMT-seq cells are incubated with a GpC methyltransferase enzyme that labels accessible GpC sites via DNA methylation. Thus, after bisulfite sequencing, GpC methylation marks can be interpreted as direct read-outs for chromatin accessibility. This stands in contrast to CpG methylation marks, which can be interpreted as endogenous DNA methylation. In addition, by physically separating the genomic DNA from the mRNA, scNMT-seq can profile RNA expression, DNA methylation and chromatin accessibility read-outs from the same cell. This hackathon leverages scNMT-seq data to assess the performance of data integration strategies to infer complex gene regulatory mechansisms driving cell fate decisions.

### **Data set description**

Gastrulation marks a major lineage specification event in mammalian embryos, accompanied by profound transcriptional rewiring and epigenetic remodelling [5].

In this hackaton we used a data set where scNMT-seq was applied to study mouse gastrulation [14] by profiling cells across four developmental stages (E4.5 to E7.5). For simplicity, in the workshop we focused on the integration of RNA expression and DNA methylation, quantified over the following genomic contexts: gene bodies, promoters, CpG islands, and DHS open sites. A total of 799 cells passed quality control for the two modalities. Overview of data is shown in Figure ??A.

Applying standard dimensionality reduction algorithms confirms that all three embryonic stages can be separated on the basis of RNA expression. The task of the workshop was to leverage the multifaceted nature of measurements to better resolve the single-cell subpopulations from distinct embyonic stages.

# **Computational integration**

We considered three computational strategies:

- **block sPLS** [15]:\*\* block sparse Projection to Latent Structures model implemented in mixOmics[16]. A sparse generalization of canonical correlation analysis that maximizes covariation between multiple sets of data modalities.
- **LIGER** [17]: unsupervised non-negative matrix factorization model for manifold alignment. It assumes a common feature space by aggregating DNA methylation over gene-centric elements (promoters or gene bodies) but allows cells to vary between data modalities.
- MOSAIC: a multi-omics supervised integrative clustering algorithm inspired by survClust [18].



hackathon analyses. **A** A summary of a subset of data types used. Challenges 1 and 2: defining genomic contexts, varying dimensions and different amount of missing values across various modalities. **B** Uniform Manifold Approximation and Projection of RNA measurements using 671 highly variable genes resolves the various embryonic stages shown by different colors. **C** Supervised analysis using view-specific and integrative distance measures (MOSAIC). The integration finds 5 clusters of cell populations based on Adjusted Mutual Information (AMI) and Standardized Pooled Within Sum of Squares (SPWSS) and outperforms individual analyses. **D** Joint clustering of cells finds common and

view-specifc patterns of variation. LIGER uses non-negative matrix factorization to integrate transcriptome and genebody methylome and align cells in a common subspace. The poor alignment suggests decoupling of the gene expression and genebody methylation during gastrulation and highlights the importance of linking relevant epigenetic features to transcriptome (challenge 3). **E** The shared variation among views is largely driven by embryonic stage. Unsupervised integration with variable selection using Projection to Latent Structures finds covarying components from RNA and methylation data which separate the cells from each stage in all views (mean balanced accuracy of kmeans using components 1 and 2 = 88%). Each point represents a cell and the axes represent the derived components for each view. The method either omits the missing values (challenge 2) while integration or incorporates the inferred values in the integration (shown). Balanced accuracy measures for a k-means clustering analysis using the extracted components from the two scenarios shows that genebody methylation components have considerably improved in predicting the embryonic stage after imputation (bottom right).

### **Computational challenges**

### **Challenge 1: defining genomic features**

The first challenge concerns the definition of the input data. The output of single-cell bisulfite sequencing are binary DNA methylation measurements for individual CpG sites. Integrative analysis at the CpG level is extremely challenging due to (1) the sparsity levels, (2) the binary nature of the readouts, and (3) the intricacy in interpretation of individual dinucleotides. To address these problems, DNA methylation measurements are typically aggregated over pre-defined sets of genomic elements (i.e. promoters, enhancers, etc.). This preprocessing step reduces sparsity, permits the calculation of binomial rates that are approximately continuous and can also improve interpretability of the input features.

There are two common strategies to define genomic elements. The first one is to use a running window approach across the entire genome. This strategy has been successful to distinguish heterogeneous cell types, but it does not improve interpretability and it typically leads to an enormously large feature set. The alternative strategy is to adopt a biologically-informed approach where genic annotations (e.g. genebody, exon, or promoter regions) as well as ChIP-seq data or chromatin accessibility information is employed to restrict the feature space to genomic regions of regulatory potential.

### [DESCRIBE RESULTS]

Our results confirm that the appropriate selection of the feature space is critical for a successful integration with RNA expression.

## **Challenge 2: Missing values in DNA methylation**

Because of the low amounts of starting material, single-cell bisulfite sequencing protocols are limited by incomplete CpG coverage. Nonetheless, in contrast to scRNA-seq, missing data can be discriminated from dropouts.

Two strategies were put forward in the workshop to handle missing values. The first is to define an inference framework that omits missing values, as done in PLS and MOSAIC. The second approach, for methods that do not handle missing information, including LIGER, is to perform *a priori* imputation of DNA methylation values.

Here we compared the integration performance for PLS ~and MOSAIC~ with and without imputation. We observe that after inferring the missing values in methylation data using a nearest neighbors approach, the axes of concordnat variation with gene expression better resolve the population structure in some epigentic regions. To quantify any improvement in biological relevance of the extracted components of shared variation, we used a supervised analysis which used the number of embryonic stages as the number of clusters. Notably, the balanced accuracy of prediction based on genebody methylation components improved from 70% for the original data to 86% after inferring the missing values.

### Challenge 3: Linking epigenetic features to gene expression

One of the main advantages of single-cell multi-modal assays is the ability to unbiasedly link epigenetic variation with gene expression. Transcriptional activation is associated with specific chromatin states near the gene of interest. This includes deposition of activatory histone marks such as H3K27ac (in promoters and enhancers), H3K4me3 (in promoters) and H3K36me3 (in gene bodies), binding of transcription factors, promoter and/or enhancer demethylation and chromatin remodelling. All these events are closely interconnected and leave a footprint across multiple molecular layers that can be (partialy) recovered by performing an association analysis between a specific chromatin read-out and mRNA expression. However, given the large amount of genes and regulatory regions, this task can become prohibitively large and it is mandatory to restrict the feature space to avoid a complex multiple testing problem.

A simple and practical approach from a computational perspective involves considering only putative regulatory elements within each gene's genomic neighbourhood. Nonetheless, this might miss important links with regulatory elements located far away from the neighbourhood.

In recent years, chromosome conformation capture experiments, have uncovered a complex network of chromatin interactions inside the nucleus connecting regions separated by multiple megabases along the genome and potentially involved in gene regulation. Early genome-wide contact maps generated by HiC uncovered domains spanning on the order of 1 Mb (in humans) within which genes would be coordinately regulated. Thus, a second strategy to associate putative regulatory elements to genes is to build on existing promoter-centered chromatin contact networks to restrict the association analysis to putative regulatory elements that are in 3D contact with genes. Although this is a promising strategy to reduce the complexity of the association analysis, most of our 3D interaction datasets are produced in bulk samples and it is so far unclear how much of these structures are preserved across individual cells. Single-cell conformation capture experiments are still limited by data sparsity and high levels of technical noise, but we envision that technological advances in this area will deepen our understanding on the regulatory roles of chromatin states.

# Commonalities between analytical multi-omics approaches for hackathons

Each hackathon study highlighted disparate challenges to multi-omics from different measurement technologies. Still, these studies were unified by the underlying problem of data integration. We summarize the common main challenges faced across all hackathons and the common approaches participants adopted to highlight the critical computational challenges faced in multi-omics single cell data analysis.

The choice of methods mostly relied on the challenge or biological question to address: data integration was conducted using projection approaches across all studies, whilst cell prediction required machine or statistical learning methods (SVM, Enet) and spatial analysis Hidden Markov random field or Moran's Index. As the computational methodologies span technologies, so do the

central challenges highlighted in each hackathon. For example the accuracy of the analysis depends critically on data pre-processing, (normalization, upstream feature selection), differences in scale across data sets and overlap (or lack thereof) features and cells (Figure 3). In many cases, this preprocessing can yield data mapping to common molecular features, such as genes, that can be the focus of the integration. However, as we highlighted particularly in the example of the single cell proteomics challenge, many multi-omics questions may have limited shared genes of features between studies. In this case, analysis methods investigating hierarchical structure among the 'omics, cell and phenotype layers and applied a number of measures of higher order concordance are crticial for successful cross-study intergation. Even in cases with matching molecular features, analyses focusing on hierarchical structure can reveal novel aspects of the biology.

Table ?? summarizes the main methods that were applied across all hackathons. A large number of computational analysis methods that were applied derive from bulk RNA-seq literature, with the exception of projection methods such as tSNE, UMAP and LIGER (the latter two are based on the common techniques NMF and PCA that were further developed for single cell data). In this section, we briefly highlight three common challenges faced across all hackathons.

# Common challenge 1: Dependence on pre-processing method and/or variable selection

Pre-processing steps strongly affect downstream analyses. Our participants thoroughly assessed the effect of normalization and data transformation (e.g. spatial transcriptomics, Figure ??Letter), as well as preliminary feature selection (mostly on based on highly variable genes) or feature summarization (scNMT-seq study). Ease of comparisons between analyses was facilitated by providing processed input data ([???]{sec:software}), but even such step did not avoid facing reproducibility issues between the original published study and the new analyses. For example in the spatial transcriptomics study, Coullomb selected 19 genes (*in scRNA-seq? or seqFISH?*) whereas the original paper selected 47 genes based on the same feature selection process [2]. No consensus was reached across participants analyses regarding the best way to process such emerging data, as no extensive benchmark, ground truth nor established biological results are yet available ([???]{sec:benchmark}).

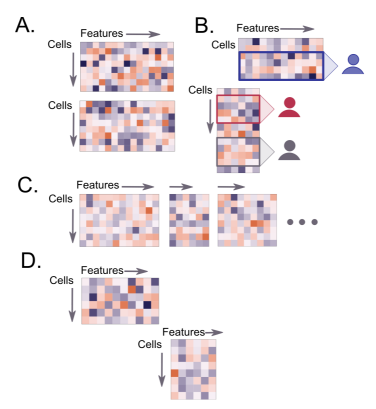
# Common challenge 2: Managing differences in scale and size across datasets

Different types of techniques were used to address the differences in scale or resolution across data sets. For spatial transcriptomics and proteomics, participants focused on a common set of genes (via feature selection in spatial transcriptomics) or proteins. The scNMT-seq study that included overlap between cells raised the issue of differences in data set size with a varying number of features per dataset ranging from 6673 to 18345, Figure ??). Some projection-based methods, such as MOFA [19], require a similar number of features in each data set, whilst others such as PLS / sGCCA [15] are not limited by such data setting and enable flexible analysis (Abadi). Difference in data scale may result in one data set contributing to either too much variation or noise during data integration. Techniques such as re-scaling (Jenagan), batch effect removal approaches, such as Combat [20] (Singh), or weighting specific data sets (Arora, Abadi) were considered and all offer further improvement in the analyses.

# Common challenge 3: Addressing partial overlap of information across cells or features

Degree of feature or cell overlap between datasets varied dramatically within each study. Intuitively, one requires at least one type of overlap (whether on the features, or on cells, Figure 3) in order to

integrate information across modalities. The field has made progress in developing methods to integrate data sets across the same (bulk) samples of single cells, mostly based on dimension reduction techniques. Amongst them, NMF (LIGER), Projection to Latent Structures (sGCCA [15]) were used for the scNMT-seq study. When there was no cell overlap (spatial studies), imputation methods were used to predict gene, protein or spatial expression values based on nearest neighbors, latent variables or optimal transport, or to predict cell types (Hsu). The most challenging study was the spatial proteomics, which raised the issue of no overlap between cells or features - the so called fourth corner that relies on phenotypes ( [???]{sec:proteomics}). We anticipate that this scenario will be avoided once technological progress and increase in data sets availability is achieved [??? 10.1186/s13059-020-1926-6].



**Figure 3: A.** Overlap of features (genes) but not cells (e.g. spatial transcriptomics where cell type prediction for seqFISH data was performed based on scRNAseq where cell types are known. **B.** Partial overlap of features (proteins) but no overlap of cells (e.g. spatial proteomics that required data imputation or cell type prediction). **C.** Overlap of cells across assays, but no overlap of features (e.g. scNMT-seq where data integration was performed). **D.** Lack of overlap between cells and features (the so-called fourth corner problem [????]{sec:proteomics}).

Table: Different methods were used in the hackathon. \* indicates the method was not applied on the hackathon data. For some common challenges, 'bulk' indicates the method was originally developed for bulk omics, 'sc' indicates the method was specifically developed specifically for single cell data **table will include links to vignettes** {#tbl:common}

challenges proteomics
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Tasks	sc Spatial	sc targeted proteomics	sc NMT-seq
Normalization & data transformation	Data distribution checks (Coullomb, Singh) High Variable Genes selection (Xu)	Variance Stabilization Normalisation [21] (Meng) Arcsinh transformation (Jeganathan). Inverse transformation (Jenagan) Selection of patients (Jenagan)	Summaries of DNA measurements (input data provided in hackathon)
Data integration	LIGER [22] (Sodicoff) (sc) ComBat (Singh) Projection methods MFA, sGCCA [15] (Singh*) (bulk) UMAP/tSNE (Sodicoff) (sc)	Multi-block PCA [23] Weighting matrices based on their similarities: STATIS, MFA (Chen*)(bulk) Scale MIBI-TOF to the range of CyTOF values (Jenagan)	LIGER [22] (Welch) (sc) Projection method sGCCA [15] (Abadi) (bulk) Multi Omics Supervised Integrative Clustering with weights (Arora) (bulk)
Cell overlap (features not matching)			Dimension reduction and projection methods: LIGER [22] (Welch) (sc) sGCCA [15] (Abadi) (bulk)
Partial feature overlap (cells not matching)		Imputation: Direct inversion with latent variables (Sankaran) Optimal transport to predict protein expression (Lin) K Nearest Neighbor averaging (Jenathan)  No imputation: Biological Network	
Partial cell overlap (features not matching)		Interaction (Foster)  Multi block PCA [23] (Meng*)	
No cell overlap (complete feature overlap)	Averaging nearest neighbors in latent space to impute unmeasured expression values (Coullomb?)	Transfer cell type label with Random Forest (Hsu)	LIGER [22] (Welch)
No cell overlap (partial feature overlap)		Topic modeling to predict cell spatial co- location or spatial expression (Jenathan, partial feature overlap)	
	Normalization & data transformation  Data integration  Cell overlap (features not matching)  Partial feature overlap (cells not matching)  Partial cell overlap (features not matching)  No cell overlap (complete feature overlap)	Normalization & data transformation Checks (Coullomb, Singh) High Variable Genes selection (Xu)  Data integration LIGER [22] (Sodicoff) (sc) ComBat (Singh) Projection methods MFA, sGCCA [15] (Singh*) (bulk) UMAP/tSNE (Sodicoff) (sc)  Cell overlap (features not matching)  Partial feature overlap (cells not matching)  No cell overlap (complete feature overlap) complete feature overlap (complete feature overlap) who cell overlap (coullomb?)  No cell overlap (coullomb?)	Normalization & data transformation Singh) High Variable Geness selection (Xu)  Data (IGER [22] (Sodicoff) (Igenagan)  LIGER [22] (Sodicoff) (Soc) ComBat (Singh) Projection methods MFA, SGCCA [15] (Singh*) (bulk) UMAP/tSNE (Sodicoff) (Soc) UMAP/tSNE (Sodicoff) (So

Common challenges	Tasks	sc Spatial	sc targeted proteomics	sc NMT-seq
	No overlap		RLQ [ <u>24</u> ] (Chen*)	
Generic approaches	Classification & feature selection	Backward selection with SVM (Coullomb) self training ENet (Singh) Balanced error rate (Coullomb, Singh) Recursive Feature Elimination (Xu) (all bulk)		Multi Omics Supervised Integrative Clustering (Arora) (bulk) Lasso penalization in regression- type models (bulk)
	Cell type prediction	Projection with LIGER [22] (Sodicoff) SVM (Coullomb, Xu) ssEnet (Singh) (all bulk)		
	Spatial analysis	Hidden Markov random field Voronoi tesselation (Coullomb) (bulk)	Spatial autocorrelation with Moran's Index (Hsu, Lin)  Selection of spatial discriminative features: Moran's Index, NN correlation, Cell type, interaction composition, L function (Lin)  (all bulk?)	
	Inclusion of additional information		Survival prediction: Cox regression based on spatial features (Lin)	Include annotated hypersensitive sites index to anchor new/unseen data from DNase-seq, (sc)ATAC-seq, scNMT-seq, for <i>de novo</i> peak calling (Meuleman*) (bulk)

# **Challenges for interpretation**

While the three hackathon analyses emphasized that regardless of the common challenges faced by our participants, there is no one method fits all multi-omics challenges. An equally important complement to the diverse computational methods used to solve multi-omics analysis problems rests in the biological interpretation of their solutions. While better resolving biological systems, the integrated data resulting from these approaches are often even higher dimensional than each of the input datasets. For example, abstract lower dimensional representations of spatial coordinates successfully captured higher level cellular structure and were more prognostics that individual feature information, suggesting that new measures of the tumor or cell ecosystems of interacting cells are needed because these interactions are fundamental to disease progression. Both this high-dimensionality and biologicaly complexity introduce further challenges in the understanding and communication of results from complex data sets and analyses.

These efforts will require standardized vocabulary, benchmarked methods, and common abstracted variables that can be compared between studies.

### Supervised versus unsupervised

Interpretation hinges on the analysis method selected for a given dataset. One simple delineation between methods used throughout the hackathons and summarized in Table {#tbl:common} is that some aim to predict a clearly defined outcome at the start of the project, such as recognizing the environment of tumor cells versus that of healthy cells [???](sec:scProteomics). The supervised setting often provides easier interpretations, one can easily rank the covariates and contiguous data in terms of their predictive potential.

On the other hand when data are collected using multiple different technologies the data integration needs to provide organizing patterns that enable interpretation. Clustering is often used as one unsupervised method and is a good example of the use of a latent variable, in this case a factor or categorical variable which was not directly measured on the data but is often used to enable simple interpretations.

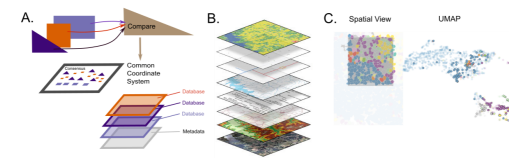
In cellular biology, a favorite such division into clusters is that involved in the definition of cell type [25].

Sometimes people get carried away in "clustering data" and manipulate the data, in cytometry one often sees cell gating done. The goal there is to eliminate cells in intermediary states to give clearly delineated inventories of cell types or cells in discrete states, this is a static description and will not enable researchers down the road to predict or understand transtitions between types.

Although a latent factor can be a useful first approximation, keep in mind that development of cells and their fate is a dynamic process and it can often be beneficial to keep data that enable interpretation of the cell trajectories: in that case, locally the underlying latent variable of interest is continuous along a gradient of development.

So far, we have seen two types of latent variables: clusters and a one dimensional continuous "gradient", (pseduo-time, disease progression are two examples of such latent gradients). However the idea of latent variables is a rich anchor for many multimodal methods and can often be useful in highlighting what the modalities have in "common" and how they differ. The commonalities are well understood in the case of classical multivariate factor analyses where the data are decomposed into "commonalities" and uniqueness components [26].

A schematic summary of the different stages in interpretation is provided here:



**Figure 4:** A Schematic diagram of stages of interpretaion and integration of data sources. **B** Standards in Geographic Information Systems enable the integration of multiple layers of data. **C** Brushing an UMAP with a covariate can illustrate the dynamics of cell changes (**Kris: I have this feeling this is not the type of figure Susan intended to show**).

Multiple domains of knowledge can be combined easily if there is a common coordinate system, as in geospatial analyses. This is often a goal in multimodal or conjoint analyses, when the first step is to find a common compromise or consensus on which to project each of the individual modalities. Conjoint analyses also known as STATIS [27] was a very early multimodal method designed as PCA of PCAs where the first step in the analyses was to find what the different modalities had in common and define a consensus [28] onto which the individual tables were projected. This method can be seen as an extension of the class of matrix decomposition methods to data cubes. Many extensions to matrix decompositions have been designed for multimodal data, [29] offers an overview of the relations between many of them.

### Reasonning by analogy with geospatial problems

In both the proteomics example [???](sec:scProteomics) and the [???](sec:scSpatial) examplary data, a spatial dimension is already naturally available. As in previous studies one can leverage extensive methods developed in spatial statistics to quantify spatial effects [30]. Contiguity and clustering can be tested and easily understood in the spatial context.

In these cases, layers of information can be mapped to the natural coordinate system in the same way a GIS system incorporates them (Figure  $\frac{4}{9}$ B).

The spatial coordinate system analogy can be pursued further by finding a "consensus space" that provides a common coordinate system.

There are however pitfalls in using very sophisticated dimension reduction techniques which lead to over-interpretation or misinterpretation (size of clusters in tSNE related to sampling baselines rather than density, ...)

# Disparate sources of evidence are more compelling than more of the same.

Following <u>Cardinal Newman's principle</u> disparate sources of evidence, or in this case data from different technologies, are more compelling than many replicates of the same technology. Thus, if different technologies allow a consensus on underlying latent variables, this information is worth retaining.

# Explaining results to biologists through generative models and simulations (ex: Factor Analysis, Hierarchical models).

Several difficulties arise when explaining summaries and conclusions, problems encountered include non-identifiability of models or non-sufficiency of summaries, simulations can often provide effective communication tools.

One can often generate data from different probabilistic models and show that the methods cannot differentiate between the generation processes, this is the identifiability problems that most overparametrized models lead to. Added constraints on the parameters can often be integrated into the analyses to make them more realistic and reduce if not eliminate the identifiability issues.

### Meaningful Interpretation by linking in databases

In the right side of Figure 4A we show how connections to layers of information from outside databases can be incorporated into the final output. Real biological understanding is often

subordinated to the integration of this contiguous information. Either from the metadata already available in the multiassay containers as for instance in the <u>MultiAssayExperiment package</u> or from exterior sources such as Gene Ontologies, Biomart [31], Kegg, Human Cell Atlas (HCA) or other sources often available through links provided within systems like bioconductor ().

Redundant biological knowledge is often enlightening, as many methods suffer from identifiability issues (ie in a gradient, the direction of the direction is unknown). By providing information on the extreme points in a map or brushing a map with known gene expression features one can delineate orientations and clusters.

For instance coloring by CD56 across time shows the dynamics of immune response [32] (Figure 4C).

### Visualization tools for interpretation and communication to biologists

An example of effective visual interpretation tools is interactive brushing of UMAP plot, see Figure 4C by Kris Sankaran.

# Interpretation for data scientists reading the methods sections requires a good understanding of the building blocks

Spanning all of these interpretation challenges is a further central communication barriers within the community of data scientists, computer scientists and computational biologists ie communicating about methods within a community of practitionners who do not have the same vocabulary or background.

Many tools are used as black boxes and users don't know or agree on what exactly the methods are doing (MOFA and tSNE are examples). The first step in unblinding these black boxes used as methodology shortcuts is to have a clear glossary of terms and how we are using them. Many synonyms for multimodal data exist and some have nuances, see the table we have compiled (ref: Table1). Understanding the relation between methods developed by different teams is essential and we often try to organize the methods first, thus it is useful to create a dichotomy of methods and their underlying properties.

A very useful tool for making methodological black boxes more transparent are simulated data. These can follow benchmark methods such as those presented in [???](sec:sec-benchmark) and use well defined generative processes to clarify what some complex methods do.

Visualization of the data, following the step by step transformations and optimizations of data representations also help clarify how certain methods fit models or compress and reduce data dimensionality. These visualizations are often very specialized (think for instance, correspondence analyses, goodness of fit plots like qqplots or rootograms or mean-variance fitting). These intermediary plots don't usually end up in the main text of final biological publications and serve as intermediary checks to unpack the black boxes.

### Missing

- Validation through complementary data and sequential experimental design.
- Examples from other parts, references and commentary here missing until documents become availabe ([???])

### References

Cell type definition: [25]

Factor Analysis: [26]

Statis, conjoint analysis: [27]

The French way: [28]

Overview and connections of methods: KS [29]

Kevin Murphy: Probabilistic Machine Learning, MIT Press [???] [33] [33]

GIS: reference https://www.usgs.gov/faqs/what-a-geographic-information-system-gis

Original: https://prd-wret.s3.us-west-2.amazonaws.com/assets/palladium/production/s3fs-public/styles/full\_width/public/thumbnails/image/8BaseLayersofTheNationalMap.JPG

Biomart: [31]

UMAP: [34] https://arxiv.org/abs/1802.03426v2

Spatial tumor and immune cells: [30]

CD56 Immune cell coloring, paper with C. Blish: [32]

Footnote: Cardinal Newman wrote **The Grammar of Assent.** and cited in [Bruno de Finetti, Volume 1, 1974 Theory of Probability]:

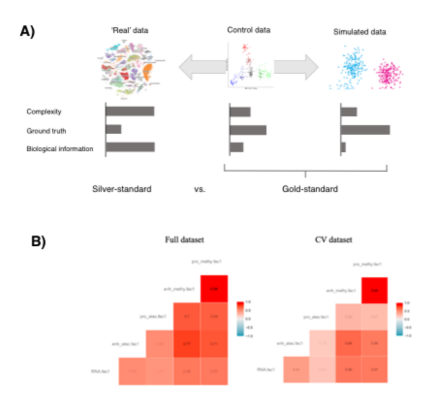
Supposes a thesis (e.g. the guilt of an accused man) is supported by a great deal of circumstantial evidence of different forms, but in agreement with each other; then even if each piece of evidence is in itself insufficient to produce any strong belief, the thesis is decisively strengthened by their joint effect.

### Techniques and challenges for benchmarking methods

### **Definition of benchmarking**

Visualizations and biological assessment of marker gene lists resulting from multi-omics analyses provide critical interpretation of integrative analysis of high-throughput data. Additional quantitative metrics are essential to establish to delineate biologically-relevant features from features resulting from either comptuational and techical artifacts. Beyond interpretation, such quantitative benchmarks are also essential to enable unbiased comparison between the numerous analysis methods used for analysis. For example, the goal of multi-platform single cell data analysis is often the recovery of known cell types through computational methods. In this case, The adjusted Rand Index (ARI) or other metrics for partitions can directly compare known cell types with the results from processing of raw data, quantification, and clustering. In cases where cell types or biological features are unknown a priori, one may also attempt to benchmark methods for their ability to discover known relationships between data modalities, e.g. gene regulatory relationships observed between chromatin accessibility and gene expression. However, this is made difficult by the fact that these

relationships are not fully known at the single cell level which provides the very motivation for the data generation and analysis. Cross-validation within a study and cross-study validation can be used to assess whether solutions found by multi-modal methods generalize to held-out observations or held-out studies.



**Figure 5:** A Silver standard: Svensson et al. (2017) Nat Methods; Wang et al. (2019) bioRxiv; Cole et al. (2019) Cell Systems; Zhang et al. (2017) bioRxiv; Soneson et al. (2018) Nat Methods; Saelens et al. (2019) Nat Biotechnol; Gold standard control data: Tian et al. (2019) Nat Meth; Freytag et al. (2018) F1000Res; Gold standard simulated data with the splatter R package appia et al. (2017) Genome Biol [credit: Matt Ritchie] (**Refs to be added proper**). **B** scNMT-seq study: correlations with linear projections (MOFA+) evaluated with cross-validation [credit: Wancen Mu and Michael Love].

### Strategies for benchmarking

Benchmarking multi-modal methods on typical multi-modal datasets is inherently difficult, as we rarely know the ground truth [35]. Simulation methods that generate high-throughput data in silico are useful for having known truth, but it is difficult to simulate realistic covariance structure across features and across data modalities (Figure 5A). Moreover, these simulations can inadvertently embed the same underlying assumptions as the computational methods employed for analysis, introducing further biases into benchmark studies. Therefore, high-throughput datasets with a known ground truth are also critical for multi-omics studies and robust testgrounds for future hackathon studies that were widely discussed throughout the workshop.

### **Cross-validation within study**

One strategy for quantitative assessment of multi-model methods is cross-validation (CV). While CV cannot determine the accuracy of a method, it can be used to assess a method's self-consistency. For example a <u>cross-validation analysis of the scNMT-seq dataset</u> was performed as part of the hackathon for this meeting using MOFA+ (Figure 5B). Such community-based benchmarking efforts in the area of multi-modal data analysis could follow the paradigm of the <u>DREAM Challenges</u>, with multi-modal training data provided and test samples held out in order to evaluate the method submissions from participating groups.

Evaluation of methods using permutation or cross-validation has been performed previously, typically to optimize a tuning parameter or other aspects of model selection. Permutation has been used to create null datasets, either as demonstration that a method is not overfitting, or for tuning parameter selection, where the optimal parameter setting should produce an objective that is far from the null distribution [36,37,38]. Cross-validation using folds or leave-one-out has likewise been used in many multi-modal method development papers [16,39,40,41,42,43,44,45,46,47,48,49].

A challenge with within study cross-validation is how to match dimensions of latent space across folds. Previous evaluations of multi-modal methods using cross-validation have focused on the top factor, swapping the sign of the project as needed to align the top latent factor across folds [50].

Finally, we note that for assessing clustering, a number of papers have suggested to resampling or data-splitting strategies to determine prediction strength [9,51,52,53]. For clustering of cells into putative cell types or cell states, such previously developed techniques could be applied in a multi-modal setting.

Cross-study validation would assess if relationships discovered in one dataset present in other datasets, potentially looking across single cell and bulk.

### **Creating benchmarking datasets**

### (add examples from Google Doc)

Benchmark datasets for single cell studies have largely centered around measuring sequencing depth and diversity of cell types derived from a single assay of interest (e.g. scRNAseq). A benchmark dataset serves a few purposes:

- Provides ground truth for the intended effect of exposure in a proposed study design.
- Provides validation for a data integration task for which a new computational method may be proposed.

For multi-modal assays, while the intended effects can vary based on the leading biological questions, one may abstract out common data integration tasks such as co-embedding, mapping or correlation, and inferring causal relationships. We distinguish data integration from further downstream analyses that may occur on integrated samples such as differential analysis of both assays with regard to to a certain exposure.

Both the intended effects and data integration task rely on study design that takes into account:

- Biological and technical variability via replicates, block design, and randomization.
- Power analysis for the intended effect or data integration task.
- Dependencies between modalities, for e.g. gene expression depending on gene regulatory element activity, requires that experiment design must also account for spatial and temporal elements in sampling for a given observation.

As such, no universal benchmark data scheme may suit every combination of modality, and benchmark datasets may be established for commonly used combinations of modalities or technologies, towards specific data integration tasks.

# Software strategies to enable analyses of multimodal single cell experiments

Community-developed, open source software and data sharing platforms enable the application of novel computational frameworks to multimodal genome-scale experiments. Our hackathons were designed to leverage and build on these open frameworks. We provided easy to use data packages to distribute the multi-omics data, and we developed vignettes that reproducibly demonstrate each of the analysis approaches employed for these datasets

(https://github.com/BIRSBiointegration/Hackathon). These open-source hackathon components provide a critical resource to enable the use and evaluation of robust benchmark strategies, and to support the enhancement and extension of the components themselves. Data structures and vignettes were largely developed on the R/Bioconductor ecosystem for reproducible, open-source, open development genomic analysis. In this section, we reflect on the software infrastructure that enabled the analytic frameworks for inference and visualization of the multimodal single-cell experiments in the hackathons. Additionally, we discuss unmet challenges identified through the hackathons, that require future development. A wide array of open-source genomics frameworks for multi-platform single cell data are also developing in R, Python, and in other software frameworks that enable graphical user interfaces for point-and-click exploration. Exploration and engagement with all these tools is richly enabled through code sharing resources like GitHub.

### **Basic aims**

We take it for granted that **openness** is a *sine qua non* for computational tooling in this area. All components need to be accessible for full vetting by the community, so licensing in Creative Commons, Artistic, or GNU frameworks is expected. We also aim for a **coordinated approach**, so that duplication of effort between groups working on similar problems can be avoided. Finally, we seek solutions that are **efficient**, avoid **lock-in**, and lead to reproducible analyses. Real-time improvements to the tool-set should be feasible, respecting needs for stability, reliability, and continuity of access to evolving components.

These objectives are fluid and open to interpretation. Community engagement and communication are important to achieving desired goals in this domain.

## **Key questions**

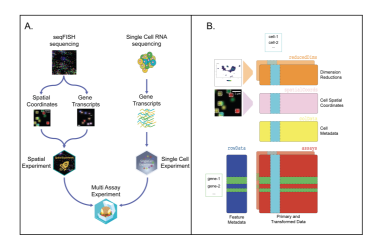
- How should multimodal single cell data be managed for interactive and batch analyses?
- What methods will help software developers create scalable solutions for multimodal single cell analysis?
- How can we ensure that visualization methods that are central to multimodal single cell analysis are usable by researchers with visual impairments?

These questions will ultimately be answered through the creation of a **software ecosystem** [54,55,56]. As an example of an ecosystem of broad scope, we cite bioconductor.org [57]. This project produces code in R for **data representation and data services** for many data modalities used in genome-scale experimentation. Bioconductor's resources for achieving **scalability** [58] include tools for analyzing massive data resources with tunable RAM footprints [???,??], and tooling for supporting fault-tolerant parallel distributed computing in various cluster and cloud contexts [59]. Finally, Bioconductor supports *developers* who seek to build broad user bases by providing multiplatform/multistream **continuous integration/continuous delivery** of contributed packages [60], and *users* with different skill sets by articulating standards for documentation, and testing, and by hosting community forums and workshops [61].

### **Data management strategies**

A ready-to-use integrative data class with multiAssayExperiment. The Bioconductor S4 class implementing an abstract data type called multiAssayExperiment is highly relevant for multimodal single-cell experiments as each mode is characterized by a different collection of features on possibly non-overlapping collections of samples [62].

The Metadata on features is bound directly into the class instance. For example, genes and transcripts can be enumerated using Ensembl [63] catalog identifiers, represented as GRanges instances [64]; regions of accessibility from, e.g., ATAC-seq experiments, may be defined using genomic coordinates in a clearly specified reference build. Metadata on samples includes all relevant information on experimental conditions such as treatment, protocol, and date of technical processing. Figure 6 shows how this class was used to amalgamate and annotate results of a multimodal dataset consisting of seqFISH and scRNA-seq experimental data. To combine these two different experiments, the seqFISH data were stored into the SpatialExperiment S4 class object, while the scRNA-seq data were stored into a SingleCellExperiment Bioconductor class object [65]. Then, these objects were easily stored into a MultiAssayExperiment class object and released with the SingleCellMultiModal Bioconductor package [???].



**Figure 6:** Panel A: Combination of seqFISH-based SpatialExperiment and SingleCellExperiment instances into a MultiAssayExperiment. Panel B: details of the SpatialExperiment class design.

The multiAssayExperiment class includes 1) Assay slots containing variables or features from multiple modalities (e.g. gene expression units from scRNA-seq and protein units in sc-proteomics), either from the same cells or distinct cells from the same or distinct starting samples or biological specimen of origin. In some cases, the feature may be multidimensional (e.g. spatial coordinates, locations of eQTLs). 2) Metadata for sample of origin for the individual cells, e.g. study, center, phenotype, perturbation. 3) A map between the different assays to enable analysis.

Some of our contributors (Al Abadi, Patheepa Jeganathan) used the multiAssayExperiment class to integrate the multi-modal single cell data (scNMT-seq, sc Proteomics, **pointers to vignettes**), allowing for easier preprocessing, transformation, extraction of spatial information from raster objects, addition of cell information and visualization **Al, details please if needed**.

While RNA-seq has well-defined units and IDs (e.g., transcript names), other assays may be summarized at different genomic scales, e.g., gene promoters, exons, introns, or gene bodies. The GenomicRanges package [66] may be used to compute summaries at different scales and overlap between signal (e.g., ATAC-seq peaks) and genomic annotation. Further, the observations of different modalities may not be directly comparable: for instance, gene expression may be measured from individual cells in single-cell RNA-seq but spatial transcriptomics may have a finer (sub-cellular) or

coarser (multi-cellular) resolution. Methods such as SPOTlight [67] may be used to deconvolute multi-cellular spots signal. Finally, in the absence of universal standards, the metadata available may vary from analysis to analysis. A potential solution is to define the minimum set of metadata variables necessary for each assay, and for pairs of assays to be comparable for common analyses.

# **Scalability strategies**

In addition to standardize data infrastructures that allows for scalability of *storing* and *access* of large datasets [68,69], new strategies are emerging that allow for scalable *algorithms* that allow for data to be stored in memory or on disk such as unsupervised clustering of cell types in either R [70,71] or Python [72]. While these strategies were originally developed in application of single-cell RNA-sequencing analyses, these scalable algorithms are applicable to multimodal single-cell experiments.

# Data access for the hackathons and long term software strategy for multimodal single-cell experiments

Reproducibility and transparency are crucial aspect in hackathons, starting with data availability. Providing data to the community is a long standing issue (e.g. SAGEBionetwork **complete Elana**). In our context, input data across all hackathons were pre-processed with steps documented, and the data were stored in MultiAssayExperiment [73] objects which allows collating of standard data formats as well as powerful data access and data processing methods. The datasets were centrally managed and hosted on ExperimentHub [74] and the SingleCellMultiModal package [doi:10.18129/B9.bioc.SingleCellMultiModal] was used to query the relevant datasets for each analysis. At the moment, it includes the hackathons scNMT-seq and the seqFISH+scRNA-seq (Fig. 6) datasets, easily stored because of the samples overlapping between multiple modalities, while we are working on the integration of the scProteomics dataset which has no overlap between samples.

### The contributed analyses reports

[https://github.com/BIRSBiointegration/Hackathon/blob/master/analysis-vignettes.md] were compiled into reproducible workflows [using GitHub Actions] enabling continuous integration of changes to source codes and continuous delivery of vignettes as well as containerized snapshots of the analyses environments. For challenge studies conducted in R, the source code and vignettes were assembled into R packages which took the loaded libraries as dependencies. For those carried out in Python, the requierments were included in the source files for automatic installation and deployment[using https://github.com/fastai/fastpages].

### Reducing barriers to interpretable visualizations

Color is a powerful data visualization tool that helps representing the different dimensions of our increasingly complex and rich scientific data. Color vision deficiencies affect a substantial portion of the population[75] and leads to difficulties in perceiving patterns (the basis for the Ishihara's color vision tests) in multi-colored figures. In some cases, the perceived patterns such as heat maps and reduced dimension plots can differ between individuals with normal and color deficient vision.

One strategy to present scientific information accessible information to all readers is to include colorblind friendly visualizations [76,77] as a default setting, using palettes such as form the R packages viridis [78] and dittoSeq [79], whilst limiting to a number of 10 colors. Additional visual cues to differentiate regions (hatched areas) or cells (point shapes) can also reduce the dependence on colors. The inclusion an "accessibility caption" accompanying figures which "guide" the reader's perception of the images would greatly benefit broader data accessibility.

# **Details of working components - trimmed**

you can interact with underlying data at google sheet

Туре	Brief name (link)	Description
Matlab package	<u>CytoMAP</u>	CytoMAP: A Spatial Analysis Toolbox Reveals Features of Myeloid Cell Organization in Lymphoid Tissues
Matlab package	histoCAT	histoCAT: analysis of cell phenotypes and interactions in multiplex image cytometry data
Python library	<u>PyTorch</u>	General framework for deep learning
Python package	<u>SpaCell</u>	SpaCell: integrating tissue morphology and spatial gene expression to predict disease cells
Python package	<u>Scanpy</u>	Python package for single cell analysis
R data class	MultiAssayExperiment	unify multiple experiments
R data class	<u>SpatialExperiment</u>	SpatialExperiment: a collection of S4 classes for Spatial Data
R package	Giotto	Spatial transcriptomics
R package	<u>cytomapper</u>	cytomapper: Visualization of highly multiplexed imaging cytometry data in R
R package	<u>Spaniel</u>	Spaniel: analysis and interactive sharing of Spatial Transcriptomics data
R package	Seurat	R toolkit for single cell genomics
R package	<u>SpatialLIBD</u>	Transcriptome-scale spatial gene expression in the human dorsolateral prefrontal cortex
R package	<u>Cardinal</u>	Cardinal: an R package for statistical analysis of mass spectrometry-based imaging experiments
R package	CoGAPS	scCoGAPS learns biologically meaningful latent spaces from sparse scRNA-Seq data
R package	<u>projectR</u>	ProjectR is a transfer learning framework to rapidly explore latent spaces across independent datasets
R package	<u>SingleCellMultiModal</u>	Serves multiple datasets obtained from GEO and other sources and represents them as MultiAssayExperiment objects
R scripts	<u>SpatialAnalysis</u>	Scripts for SpatialExperiment usage
Self- contained GUI	ST viewer	ST viewer: a tool for analysis and visualization of spatial transcriptomics datasets

Туре	Brief name (link)	Description
Shiny app	<u>Dynverse</u>	A comparison of single-cell trajectory inference methods: towards more accurate and robust tools
R package	<u>mixOmics</u>	R toolkit for multivariate analysis of multi-modal data
Python package	totalVI	A variational autoencoder (deep learning model) to integrate RNA and protein data from CITE-seq experiments
Python web application		<u>lmJoy</u>
Python package	<u>napari</u>	Interactive big multi-dimensional 3D image viewer
Software	QuPath	Multiplex whole slide image analysis
Python package	<u>Cytokit</u>	Multiplex whole slide image analysis
Python package	<u>cmIF</u>	Multiplex whole slide image analysis
Software	<u>Facetto</u>	Multiplex whole slide image analysis, not available yet
Software, Python based	<u>CellProfiler</u>	Image analysis

### **Discussion**

The rapid pace of technological development can be seen as only six years after single-cell sequencing was name method of the year, single-cell multimodal omics claimed the title [80]. Beyond the title, multimodal omics has inherited not just the explosion in datasets and the formation of Atlas based initiatives, but also the wave of statistical and computational techniques they launched. Thus, a central question is what can be carried forward from both current initiatives and current methods. In the case of consortia, the ability to impose uniform processing and community standards such as those for ENCODE and IHEC will remain important. The Human Cell Atlas [81] showed how community driven initiatives have the ability to influence and inspire across larger proportions of the scientific community. Hackathons are also excellent resources for bringing the community together and providing formal benchmarks. Together with large consortia studies, hackathons have the potential both to inform and require multi-omics data analysis methods. In regard to what is needed computationally for the future of multi-omics much will be driven by the data itself. While on the one hand the inherently coupling of measurements simplifies certain challenges, on the other hand the layering of an entirely new set of molecular measurement intrinsically increases complexity. Thus, considerations must be made for cross-modal integration and modeling with both matched and unmatched multimodal measurements. In both cases methods for dealing with data quality, data loss by summarization, timing variances both between and within omics layers, and batch effects will be necessary challenges to overcome. Conversely, the ability to infer regulatory relationships between different omics and expand to dynamic models across time and space presence new opportunities.

### **Emerging analytical methods and technologies**

# Community needs for data structures, analysis methods, etc

# **Glossary**

**Table 1:** Glossary of interchangeable terms in the field of single-cell and bulk multi-omics (multi-source) data analysis.

Consensus Term	Related Terms	Description	Citation
network	graph, adjacency matrix	A set of <i>nodes</i> , representing objects of interest, linked by <i>edges</i> , representing specific relationships between nodes.	?
node	vertex	Element of interest in a network and linked to other nodes. For example: people, cells, proteins or genes. Nodes can have several properties called attributes like cell type or position.	?
edge	link	The relationship between 2 nodes in a network. For example: friendship in social networks, cells in contact in a spatial network, or gene-gene interactions in a gene regulatory network.	?
concordant	concordant, coherent, consistent	?	<u>50</u>
contributions	variable weights, loadings, eigenvector, axis, direction, dimension, coefficients, slopes	Contributions of the original variables in constructing the components.	<u>16, 82</u>
latent factors	variates, scores, projections, components, latent/hidden/unobserved variables/factors	Weighted linear combinations of the original variables.	<u>16, 82</u>
multimodal	Multiview, multiway arrays, multimodal, multidomain, multiblock, multitable, multi-omics, multi-source data analysis methods, N- integration	Methods pertaining to the analysis of multiple data matrices for the same set of observations.	<u>16, 29, 83</u>
conjoint analysis	conjoint analysis, P- integration, meta-analysis, multigroup data analysis	Methods pertaining to the analysis of multiple data matrices for the same set of variables.	<u>16, 82, 84</u>

Consensus Term	Related Terms	Description	Citation
variable	feature, variable	A measurable quantity that describes an observation's attributes. Variables from different modalities include age, sex, gene or protein abundance, single nucleotide variants, operational taxonomic units, pixel intensity etc.	?
biomarker	marker, biomarker	A variable that is associated with normal or disease processes, or responses to exposures, or interventions. Any change in this variable is also associated with a change in the associated clinical outcome. These variables may be used for diagnostic, monitoring, Pharmacodynamic responses. Examples include LDL cholesterol, CD4 counts, hemoglobin A1C.	<u>85</u>
panel	biomarker panel, biomarker signature	A subset of the originally measured variables that are determined to be associated with the outcome or response variable. This may be determined using statistical inference, feature selection methods, or machine/statistical learning.	<u>86</u> , <u>87</u>
observation	sample, observation, array	A single entity belonging to a larger grouping. Examples include patients, subjects, participants, cells, biological sample, usually the unit of observation on which the variables are measured etc.	?

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1. Supposes a thesis (e.g. the guilt of an accused man) is supported by a great deal of circumstantial evidence of different forms, but in agreement with each other; then even if each piece of evidence is in itself insufficient to produce any strong belief, the thesis is decisively strengthened by their joint effect. ←