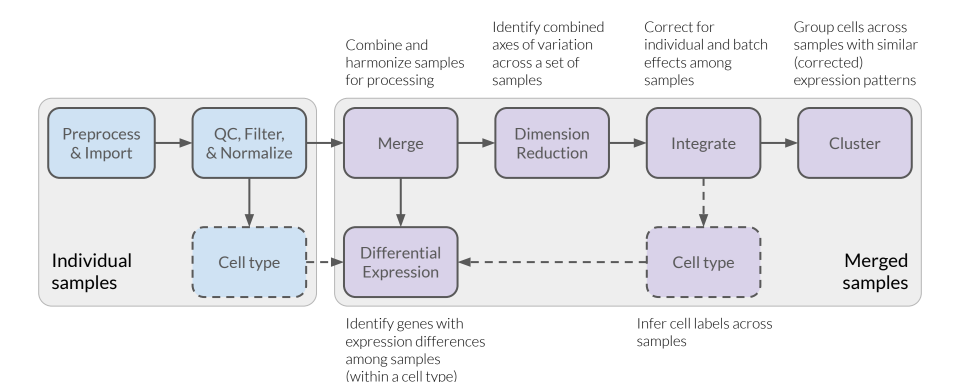
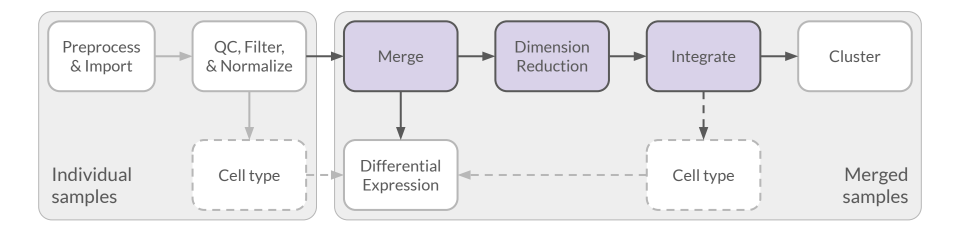
Integrating Different Samples in Single-cell RNA-seq

The Data Lab

Working with multiple samples in scRNA-seq



Integration in scRNA-seq overview



Why integrate samples?

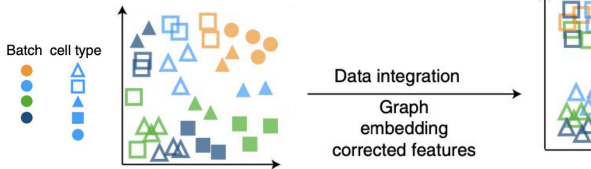
The goal of integration is to mitigate the *batch effects* caused by technical variation across samples, while still preserving biological information.

- Let's call each sample a "batch" of cells
- Cells in a given sample will share some technical variation
- This becomes a problem when we want to jointly consider several samples
 - Cells within a given sample appear more similar than they are, simply because they're from the same sample.
- To compare cells across samples, we need to remove this batch-level technical variation. Then, we can hopefully hone in on the more interesting biological variation 🕵

What can('t) integration do for you?

- Integration is performed on reduced dimension representations (often principal components)
 - Integration also *returns* reduced dimension representations for downstream use
 - Some integration methods will "back-calculate" corrected gene expression values, but these aren't as important as you think!
 - For example, we do *not* use these for differential expression (stay tuned for more!)
 - Recommended reading on when to use, and not to use, corrected expression values: <u>http://bioconductor.org/books/3.16/OSCA.multisample/using-corrected-values.html</u>
- Integration allows us to...
 - Jointly visualize **cells** from multiple datasets
 - Jointly cluster **cells** from multiple datasets
 - Annotate or identify similar **cell types** across datasets

What does successful integration look like?



Before integration, the primary "clustering" is by batch

• Orange tends to group with orange, green with green, etc.

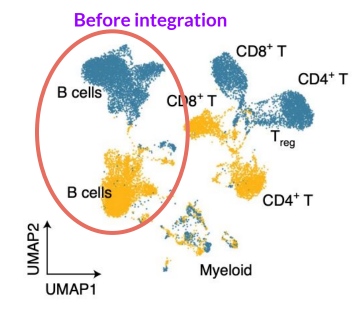
After successful integration:

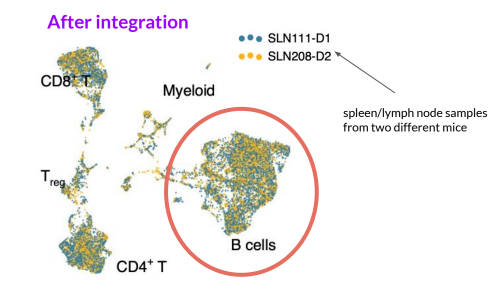
- Batches show lots of mixing
- Cell types ("biology") cluster together, and do *not* show lots of mixing

Successful integration depends on *shared information* across batches.

Figure adapted from Lueken et al. (2022) https://doi.org/10.1038/s41592-021-01336-8

Example of (what looks like!) successful integration





How to evaluate integration

• Compare before and after UMAP vibes



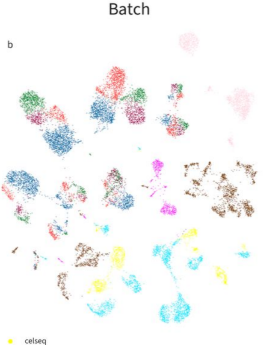
- Before integration, batches (datasets) will mostly cluster together
- After integration...
 - Batches should not group together but should be highly mixed across the UMAP
 - Biologically similar cells (tissue, cell type, disease vs healthy) should group together
- Usually, when it fails, *it fails*.

- There are several metrics for evaluating batch correction
 - Luecken *et al.* (2022) is an excellent reference <u>https://doi.org/10.1038/s41592-019-0619-0</u>
 - Caution: Metrics do not measure "was integration successful," but other proxies which *sometimes can help us tell* if integration was successful (or at least not unsuccessful)

How I stopped worrying and learned to love (the) UMAPs

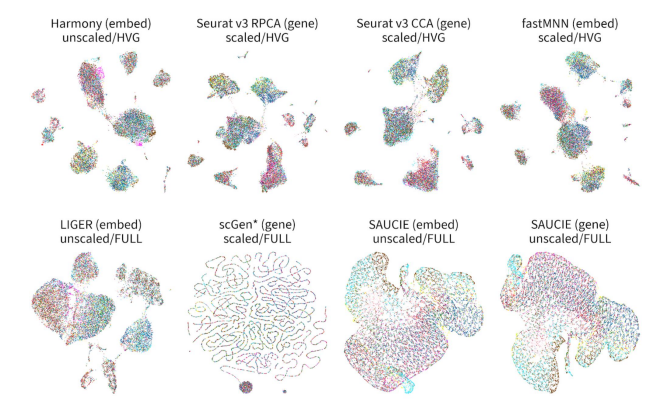
- Some examples from Luecken et al. (2022)
 - Luecken, M.D., Büttner, M., Chaichoompu, K. et al. Benchmarking atlas-level data integration in single-cell genomics. (2022). <u>https://doi.org/10.1038/s41592-021-01336-8</u>

- Panels from Figure S13 are shown on the next two slides
 - https://static-content.springer.com/esm/art%3A10.1038%2Fs41592-021-01336-8/Medi aObjects/41592 2021 1336 MOESM1 ESM.pdf

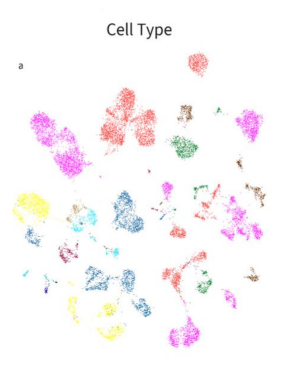


- celseq2
- fluidigmc1
- inDrop1
- inDrop2
- inDrop3
- inDrop4
- smarter
- smartseq2

Top 4 "best" integration methods



Bottom 4 "worst" integration methods

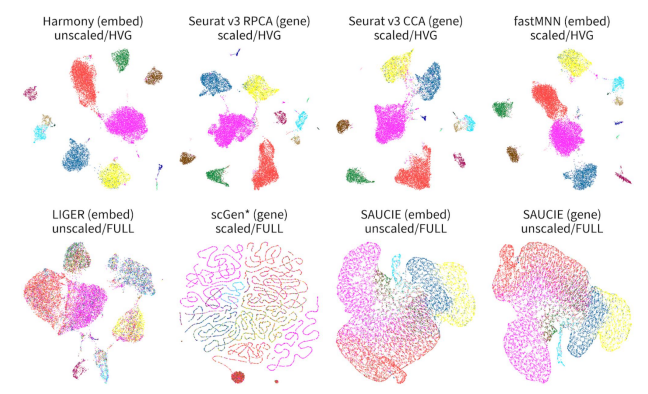


t_cell



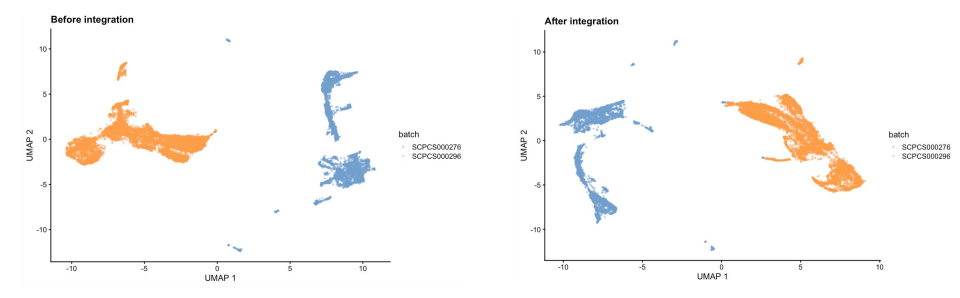
endothelial

Top 4 "best" integration methods



Bottom 4 "worst" integration methods

An example of failed integration



Will it integrate?



- Datasets that don't have shared cell types or states will be hard to integrate
 - Patient and xenograft
 - Healthy and normal
 - Data from different tissue types
 - Data from different organisms

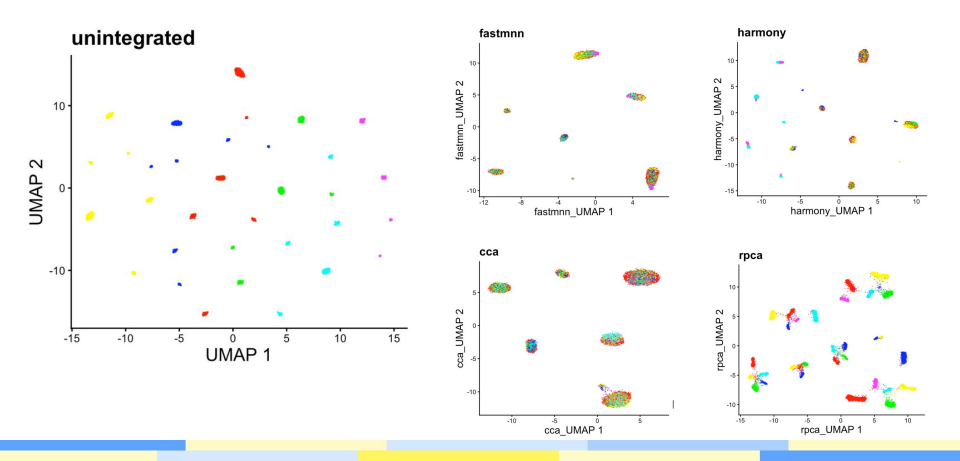
• The extent of "overlap" among datasets may also influence which integration method you should use, along with the results themselves

We performed some benchmarking on simulated data from Luecken *et al*.

- We evaluated several methods, four of which we'll show here:
 - FastMNN
 - Harmony
 - Seurat using CCA (canonical correlation analysis)
 - Seurat using RPCA (reciprocal PCA)
 - (We'll note that we also looked at scVI, which seemed to work well but is slow on CPU and it's in Python which is beyond the scope of our workshop!)
- We chose these methods based on performance in Luecken *et al.* and their usability
- See <u>https://github.com/AlexsLemonade/sc-data-integration</u> for our benchmarking code

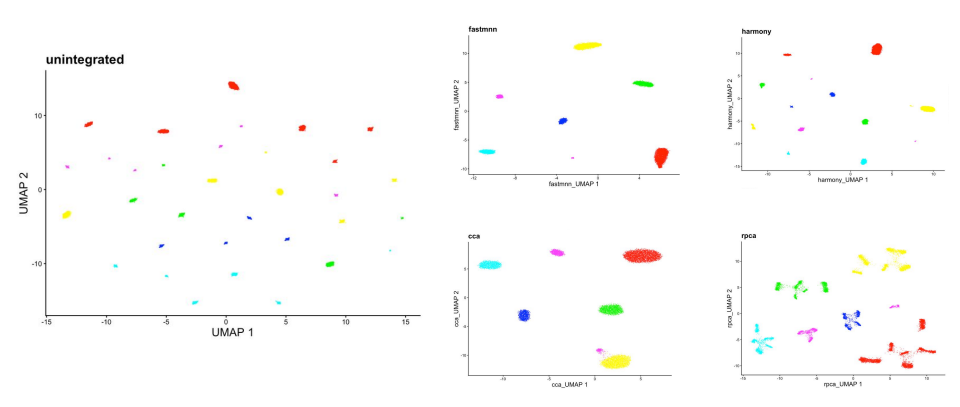
Scenario 1: All cell types are present in all batches

UMAPs colored by Batch



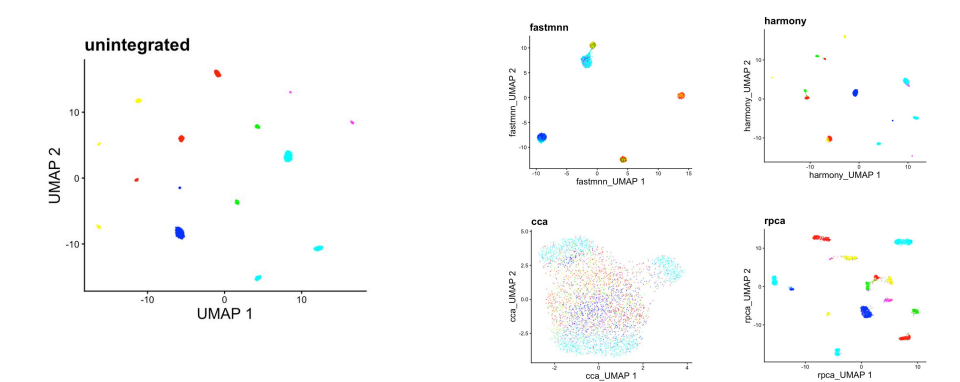
Scenario 1: All cell types are present in all batches

UMAPs colored by Cell Type



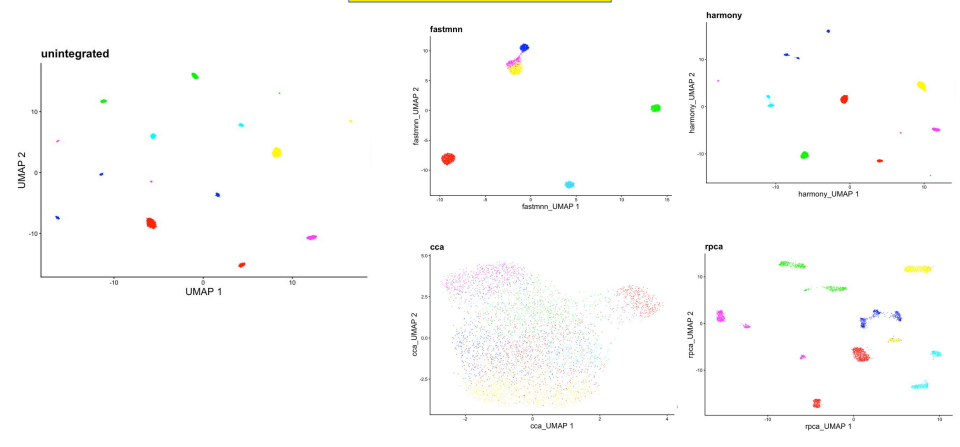
Scenario 2: Cell types are not present in all batches, and not all batches have cells in common

UMAPs colored by Batch

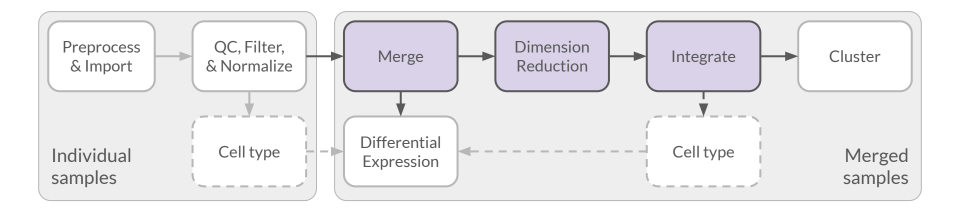


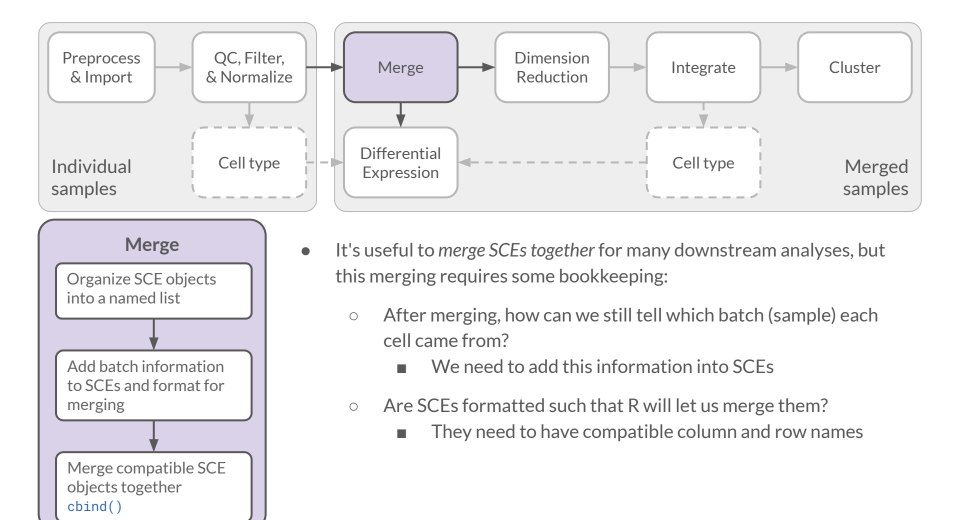
Scenario 2: Cell types are not present in all batches, and not all batches have cells in common

UMAPs colored by Cell Type



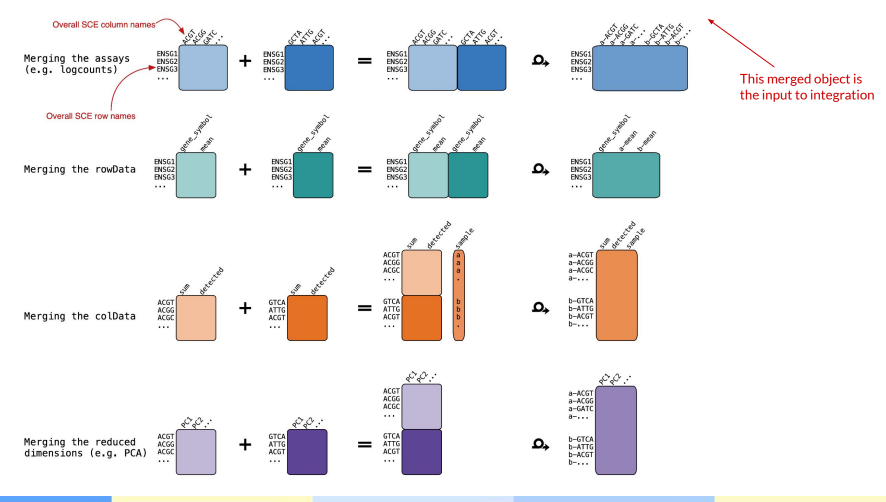
Integration in scRNA-seq overview



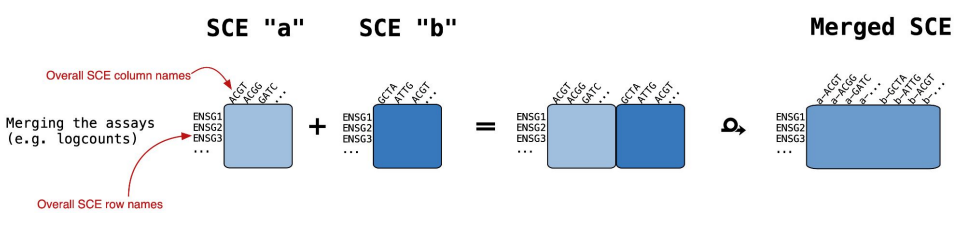


SCE "a" SCE "b"

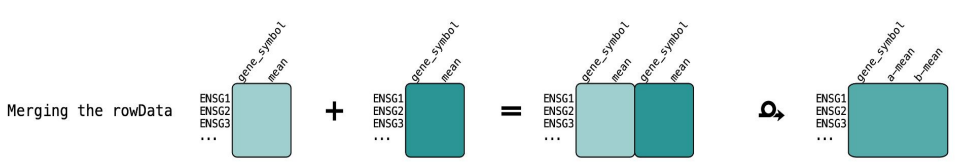
Merged SCE



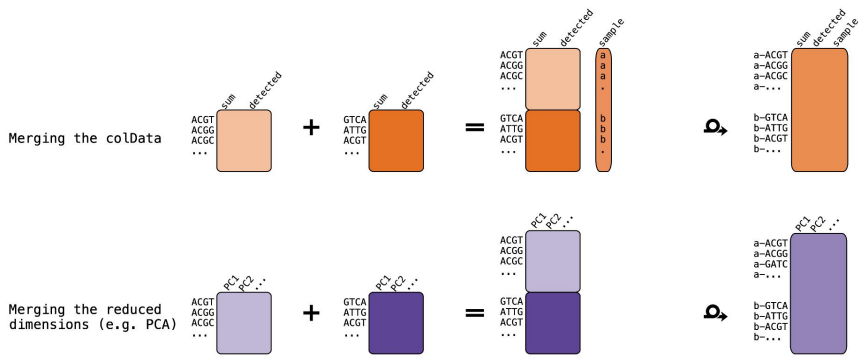
Merging SCE assays

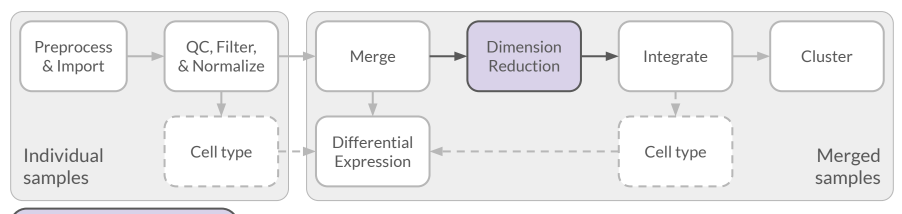


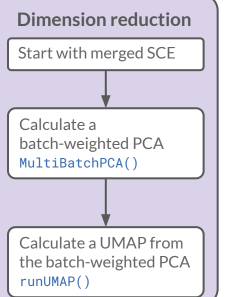
Per-gene (feature) data: Each row is a gene



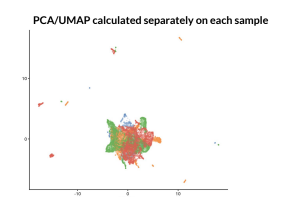
Per-cell data: Each row is a cell



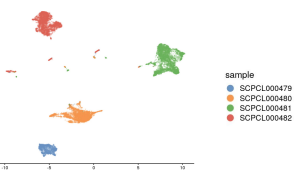


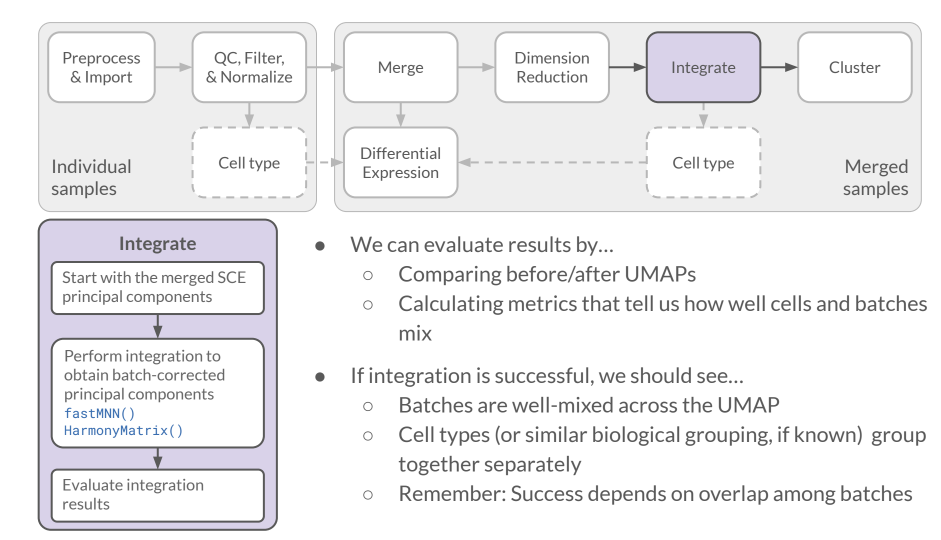


- Dimension reduction techniques like PCA and UMAP start by scaling data to be centered at 0.
- To use PCA/UMAP across samples, we need to calculate the variation jointly









Let's have a closer look at methods we'll be using

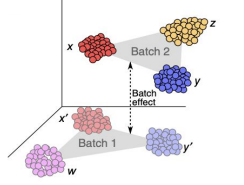
- MNN: Mutual nearest neighbors
 - Specifically, we'll use FastMNN 🚀
 - Haghverdi, L, Lun, A, Morgan, M, et al. Batch effects in single-cell RNA-sequencing data are corrected by matching mutual nearest neighbors. (2018) <u>https://doi.org/10.1038/nbt.4091</u>

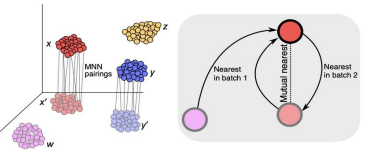
• Harmony

 Korsunsky, I, Millard, N, Fan, J, et al. Fast, sensitive and accurate integration of single-cell data with Harmony. (2019) <u>https://doi.org/10.1038/s41592-019-0619-0</u>

Mutual nearest neighbors batch correction

- Imagine we have 2 batches, each with 3 cell types
 - Red (x) and blue (y) are shared but pink (w) and yellow (z) are not!
 - Before beginning integration, cosine distances are first calculated among pairs of cells *within each sample*
 - This enables expression profile comparisons and sets up the data for integration
- First, we identify pairs of cells with mutually similar expression profiles
 - These are our "mutual nearest neighbors"

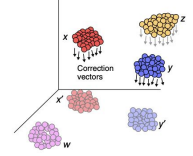


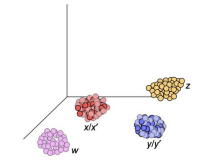


Mutual nearest neighbors batch correction

• Next, compute a batch correction vector for each MNN pair

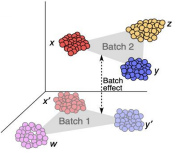
- Finally, calculate the weighted average of these vectors to get cell-specific batch corrections to perform the final integration
 - Note that w and z don't "look" as "integrated"! Why?





Some assumptions that MNN makes

- At least one cell population is present in both batches
- The batch effect is almost orthogonal to the biological effects
 - Roughly means, batches and biology are expected to have separate variation



• The batch-effect variation is much smaller than the biological-effect variation across cell types

Harmony batch correction

• "Soft k-means clustering algorithm"

