## Short-guide to the AP-MS bioinformatics pipeline

1. Filtering out the MS output for protein groups that could not be uniquely matched to a set of peptide spectra in the search database
2. Computationally removing carry-over proteins observed between IPs that were run in consecutive order
3. Clustering of total protein spectral count (or # of unique peptides) correlation between all IPs to identify
   1. Clusters of replicates (good)
   2. Dispersed individual IPs that cluster more with the negative controls than with their replicates (bad)
4. Optional: Remove ‘bad’ IPs from the dataset
5. Scoring (See below)
   1. MIST with virus-host parameters
   2. MIST with PCA derived parameters
   3. COMPPASS
   4. (Optional: SAINT)
6. Optional: Adding additional background from AP-MS database to improve detection of commonly observed unspecific binders and rerun 5.
7. Optional: Searching for enriched complexes or pathways for a given bait

## Short-guides to ‘scored’ table format

* BAIT: custom bait name
* PREY: uniprot\_ac code for identified prey protein
* MIST\_hiv: MIST score with ‘HIV’ weights (See below)
* MIST\_self: MIST score with custom weights derived by PCA (See below)
* MIST\_R: MIST reproducibility feature
* MIST\_A: MIST abundance feature
* MIST\_S: MIST specificity feature
* TSC\_AVG: Average Total Spectral Count (or # unique peptides) over replicates
* COMPPASS\_Z: Z-score for this bait-prey pair’s TSC\_AVG with respect to observations of other baits interacting with this prey
* COMPPASS\_S: Empirical COMPPASS Specificity score
* COMPPASS\_D: Empirical COMPPASS Specificity & Reproducibility score
* COMPPASS\_WD: Empirical COMPPASS Specificity & Reproducibility score with background correction
* COMPPASS\_pZ: p-value indicating probability of finding this Z-score randomly
* COMPPASS\_pS p-value indicating probability of finding this S-score randomly
* COMPPASS\_pD p-value indicating probability of finding this D-score randomly
* COMPPASS\_pWD: p-value indicating probability of finding this WD-score randomly
* SAINT\_AVG\_P: Average SAINT score over all replicates for this bait-prey pair
* SAINT\_MAX\_P: Maximal SAINT score over all replicates for this bait-prey pair

Note on thresholds:

1. We suggest a MIST threshold of > 0.75 for significant interactions
2. COMPPASS score are not intuitive to interpret but we propose ranking any of the scores descending and applying a threshold on the score’s corresponding p-value of < 0.05
3. SAINT scores correspond to the probability of an interaction being true. The authors suggest a threshold of > 0.9

## Short-guides to score calculations

### MIST

The MIST algorithm first computes three features for every bait-prey interaction given the whole input set of observed interactions: *abundance*, *reproducibility* and *specificity*. The MIST total reported score is a weighted sum of these three features. The weights were determined by Principal Component Analysis (PCA) to maximize the feature space variance in one dimension.

We currently report a MIST score with the optimal parameters for the HIV-host interaction networks. We postulate that these parameters are suitable for most sparsely interconnected bait-prey datasets (eg. virus-host interaction networks).

We also report the MIST score with custom computed weights by performing PCA on the input set of interactions. For a more detailed description of the MIST algorithm we refer to the online published supplementary material of the ‘[Global landscape of HIV-human protein complexes paper’](http://www.ncbi.nlm.nih.gov/pmc/articles/PMC3310911/)

### COMPPASS

An excellent description of the COMPPASS score can be found online on the [Harper Lab website](http://falcon.hms.harvard.edu/ipmsmsdbs/cgi-bin/tutorial.cgi)

### SAINT

All the information regarding the SAINT scoring algorithm can be found online on the [SAINT website](http://saint-apms.sourceforge.net/Main.html)