



DIAmeter: Matching peptides to dataindependent acquisition mass spectrometry data

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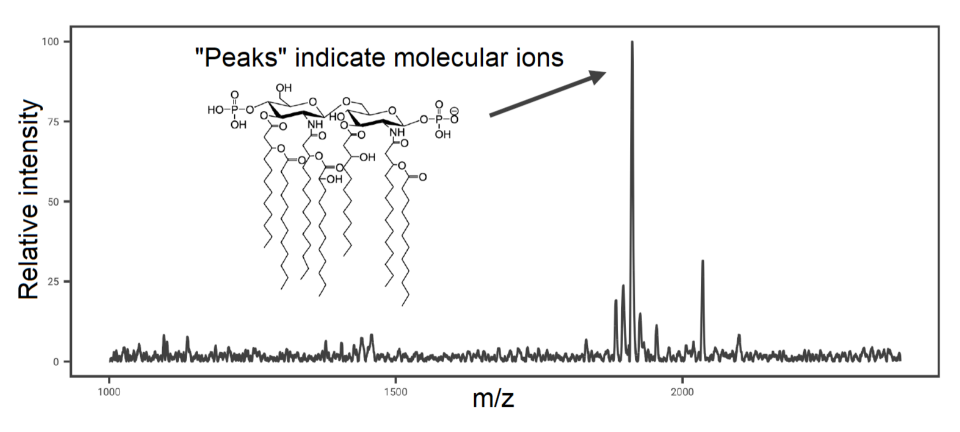




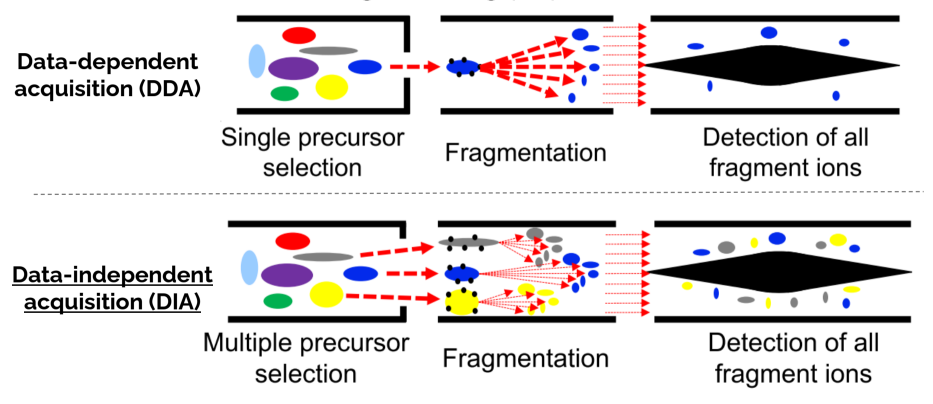




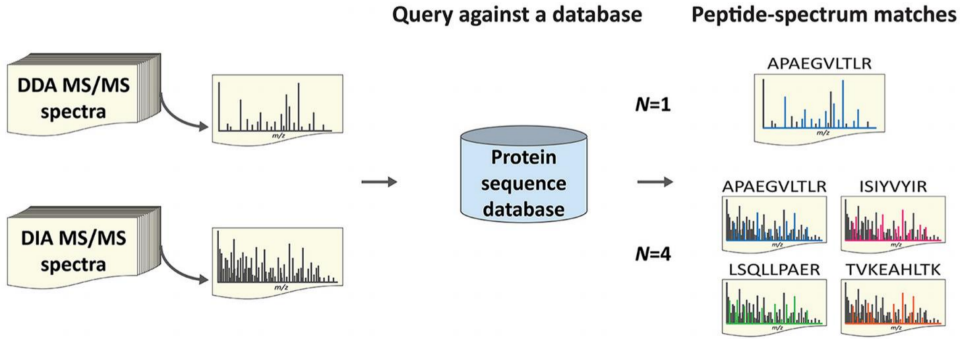
The mass spectrum is a record of the detected ions



The mass spectrum is obtained by isolating and fragmenting peptides



Standard proteomics database search process is typically designed for DDA data



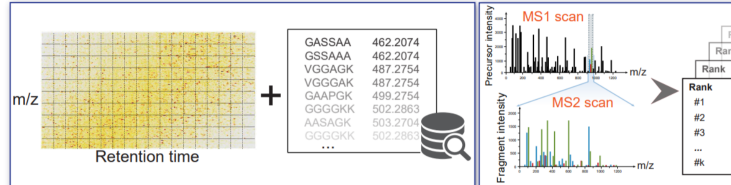
There is demand for new methods that overcome limitations of existing methods

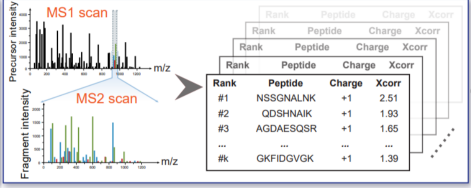
	DIAmeter	DIA-Umpire	PECAN	Prosit + EncyclopeDIA
Support wide isolation windows	\checkmark	√	X	X
Detects PTMs			1	X
Instrument independent	\checkmark	1	X	X
Detects peptides with undetectable precursor	\	X	\checkmark	

DIAmeter uses a standard DDA search engine with an extra PSM filtering step

1. Input: DIA data and a peptide database

2. Tide search identifies multiple matches per spectrum

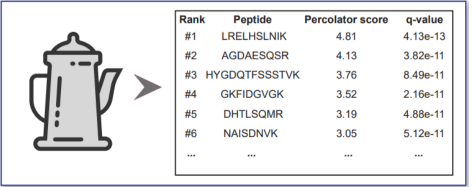




3. Matches are reduced by using PSM features

Retain/Discard Rank Peptide Additional features Composite score Rank Peptide Retain/Discard Additional features Composite score Rank Peptide Additional features Composite score Retain/Discard Retain/Discard Rank Peptide Additional features Composite score #1 NSSGNALNK 10.35 #2 **ODSHNAIK** 12.51 **AGDAESQSR** 8.96 **GKFIDGVGK** 16.38

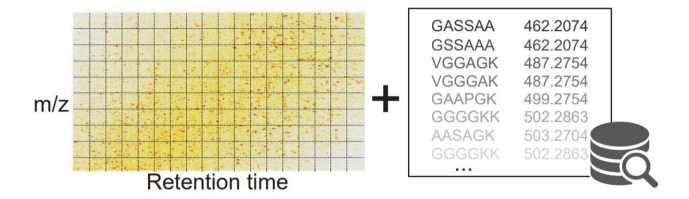
4. DIA-tailored Percolator ranks peptides by q-value

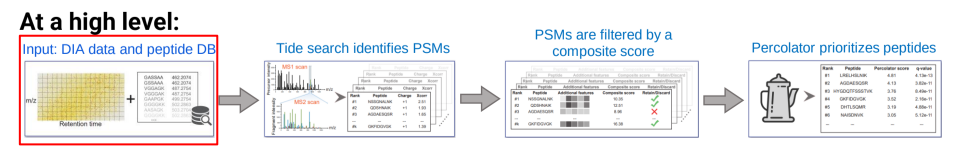


DIAmeter searches spectra by using a peptide sequence database

Input:

- □ DIA data
- A peptide database

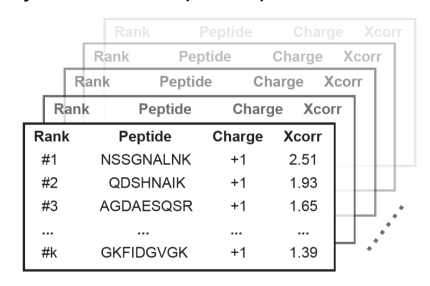


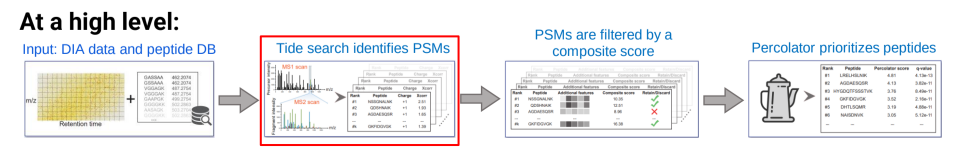


DIAmeter uses Tide to identify matches per spectrum

Tide search for PSMs:

- ☐ Use XCorr as the primary score
- ☐ Identify top-k (k=5) matches for each spectrum and each charge state

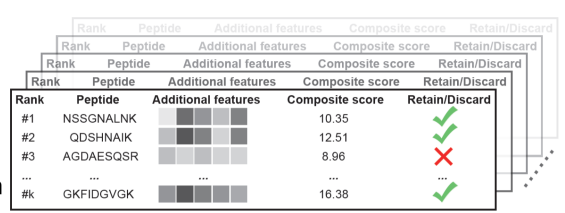




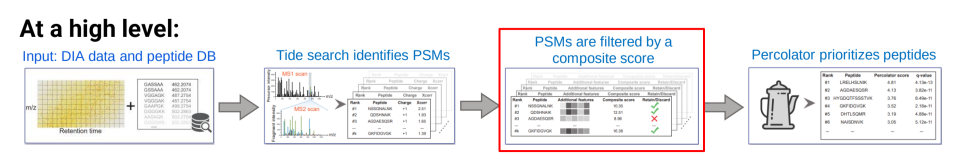
DIAmeter filters the PSMs by a composite score

The composite score is defined as the weighted sum of PSM features:

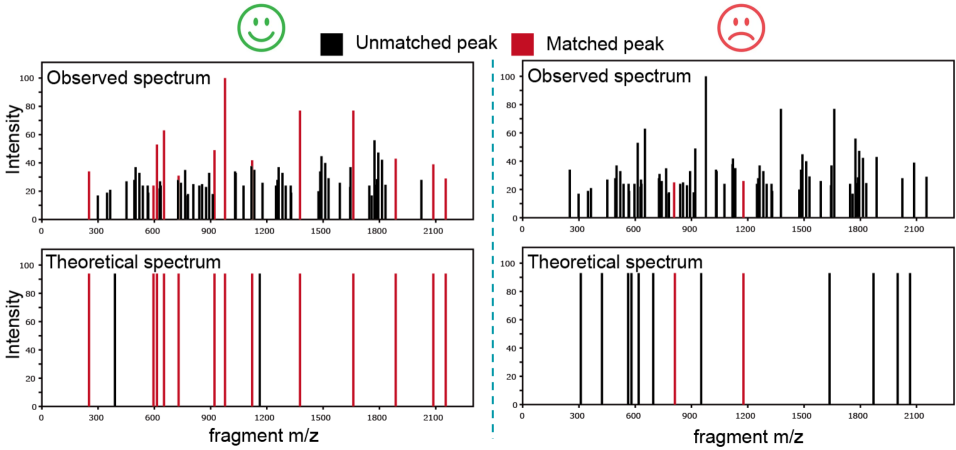
- ☐ XCorr (with Tailor calibration)
- □ Precursor intensity rank
- ☐ Fragment matching p-value
- □ Retention time score
- □ Precursor and fragment coelution



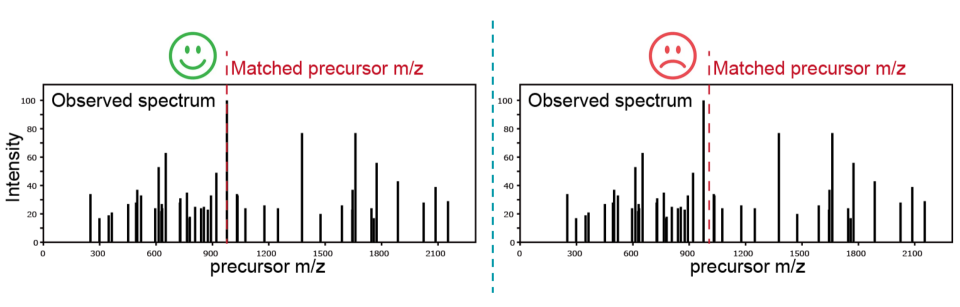
Sulimov et al. Journal of Proteome Research (2020)



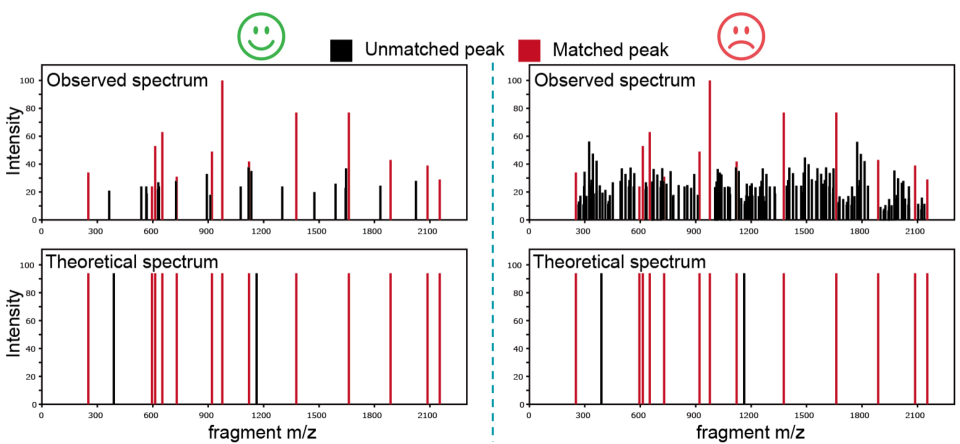
XCorr score measures the fragment-level matching evidence



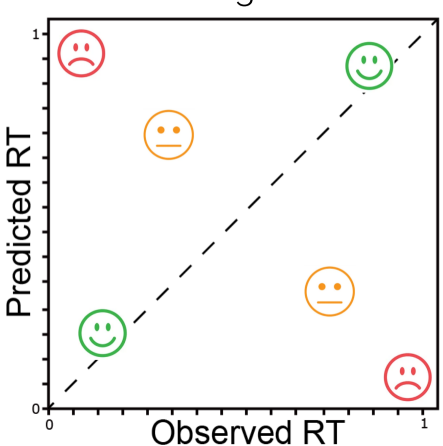
Precursor intensity measures the precursor-level matching evidence



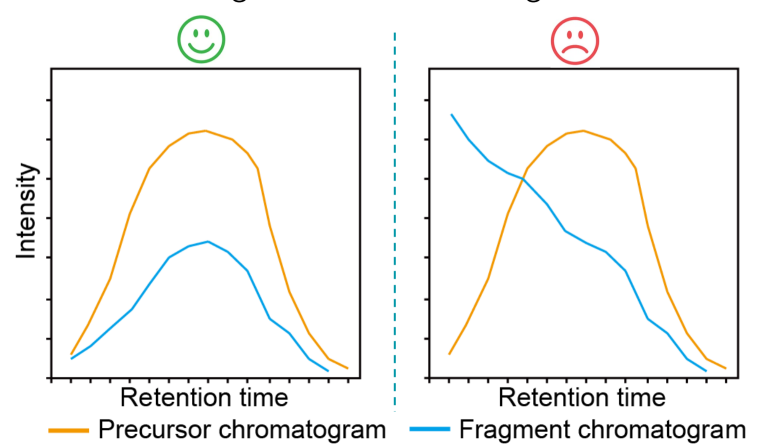
Fragment matching p-value measures how likely the theoretical fragments are to match by random chance.



Retention time score measures the temporal matching evidence



Precursor and fragment coelution measures the chromatogram-level matching evidence



DIA-tailored Percolator ranks peptides by q-value

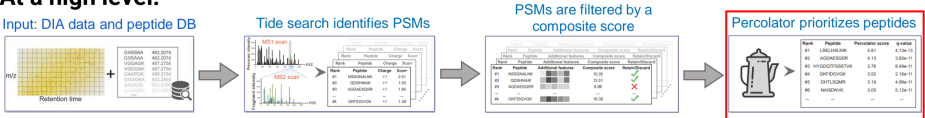
Percolator learns to re-rank these PSMs:

- ☐ The modified Percolator assigns all PSMs related to the same target-decoy pair to the same cross-validation split
- □ DIAmeter selects the top-scoring PSM per target-decoy pair
- □ DIAmeter calculates its own peptidelevel FDR estimates.

Kall et al. Nature Methods (2007)

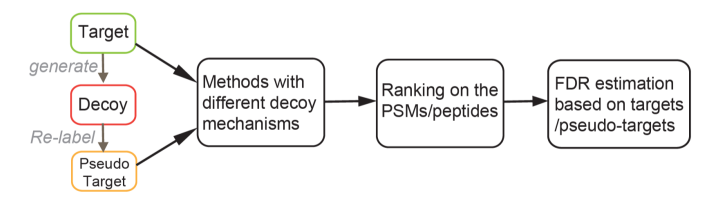
Rank	Peptide	Percolator score	q-value
#1	LRELHSLNIK	4.81	4.13e-13
#2	AGDAESQSR	4.13	3.82e-11
#3	HYGDQTFSSSTVK	3.76	8.49e-11
#4	GKFIDGVGK	3.52	2.16e-11
#5	DHTLSQMR	3.19	4.88e-11
#6	NAISDNVK	3.05	5.12e-11

At a high level:



Evaluation: Consistent FDR estimation across methods by using an entrapment strategy

- Methods are agnostic to pseudo targets
- ☐ Pseudo targets leads to loss of statistical power, only for evaluation purposes

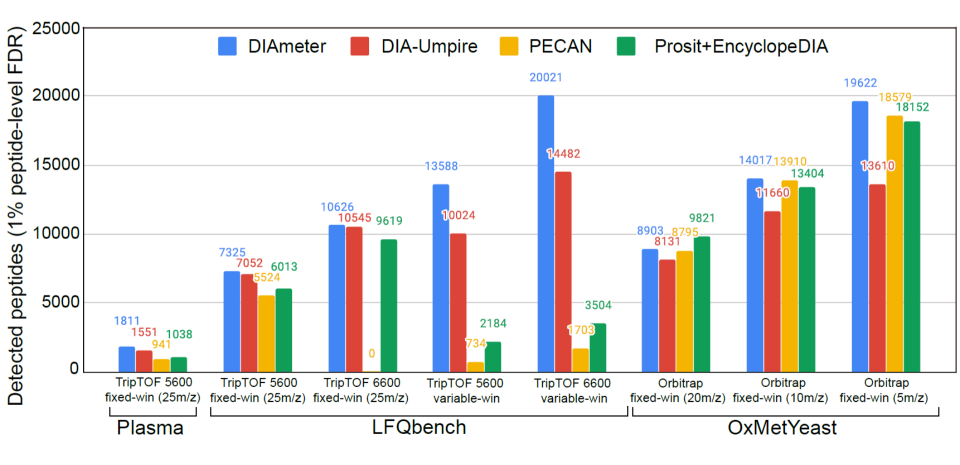


DIAmeter is evaluated on datasets across diverse settings

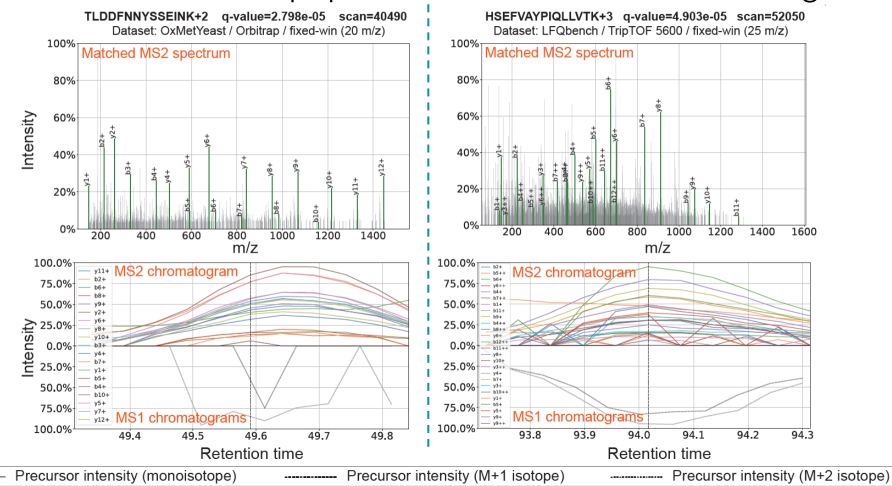
Yeast dataset with enriched methionine oxidation (OxMetYeast) ☐ Orbitrap instrument
☐ Acquired by using different isolation window sizes
Multispecies mixture dataset (LFQbench) ☐ TripTOF 5600/6600 instrument ☐ Acquired by using fixed/variable isolation window Navarro et al. Nature Biotechnology (2016)
Human blood plasma dataset (Plasma) ☐ TripTOF 5600 instrument ☐ Acquired by using variable isolation window ☐ High dynamic range

Rosenberger et al. Nature Biotechnology (2017)

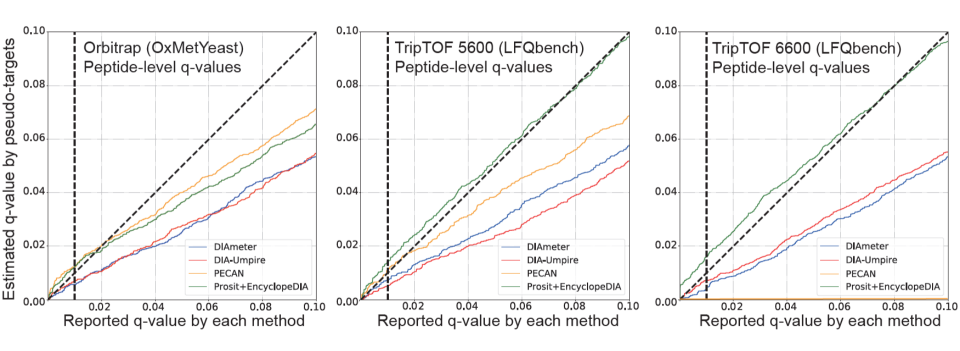
DIAmeter confidently detects more peptides at 1% FDR



DIAmeter can detect peptides with undetectable MS1 signal

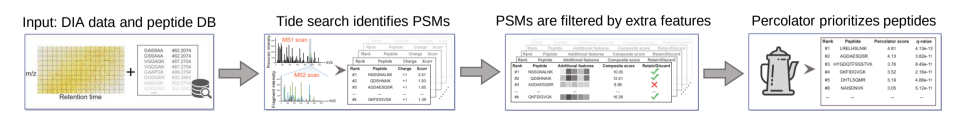


DIAmeter accurately controls the FDR



Conclusions

- □ DIAmeter is more sensitive than other methods, without relying on spectral library
- DIAmeter is robust in diverse settings
- □ DIAmeter can detect peptides with undetectable MS1 signal
- ☐ Availability: http://crux.ms/



Acknowledgements

•Noble lab members:





