#### Abstract

Motivation: Glycosylation is one of the most heterogeneous and complex post-translational modifications. Liquid chromatography coupled mass spectrometry (LC-MS) is a common high throughput method for analyzing complex biological samples. Mass spectrometry data is contains intricate structure, requiring several transformations before it can be used to identify biological molecules, requiring automated tools to analyze samples in a high throughput setting. Existing tools for interpreting the resulting data do not take into account related glycans when evaluating individual observations, limiting their sensitivity. Results: We developed an algorithm for assigning glycan compositions from LC-MS data that can use information encoding biosynthetic network relationships between glycans using Laplacian regularized least squares, and optionally prior information about expected glycan families, built atop an a set scoring functions that use glycan chemical properties. Our method was able to identify as many, or more glycan compositions compared to previous approaches, and demonstrated greater sensitivity with regularization. Our network definition was tailored to N-glycans but the method may be applied to glycomics data from other glycan families like O-glycans or heparin where the relationships between compositions can be expressed as a graph.

# Application of Network Smoothing to Glycan LC-MS Profiling

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# 1 Introduction

Glycosylation is one of the most pervasive and diverse forms of post-translational modification (Varki (2017)). Their study is of great importance for understanding broad classes of biological processes. Mass spectrometry (MS) is a powerful tool for glycan analysis (Zaia (2008)). MS of unseparated mixtures results in poor dynamic range for detection of glycans Peltoniemi et al. (2013). An online separation method like liquid chromatography (LC) or capillary electrophoresis (CE) improves dynamic range and reproducibility of glycan detection, at the cost of increased analysis time and workflow complexity.

There are many tools for interpreting glycan mass spectral datasets (Yu et al. (2013); Peltoniemi et al. (2013); Kronewitter et al. (2014); Goldberg et al. (2009); Maxwell et al. (2012); Ceroni et al. (2008); Frank and Schloissnig (2010)) for both unseparated and separated experimental protocols. These programs address instrument-specific signal processing requirements. For example SysBioWare (Frank and Schloissnig (2010)) performs sophisticated baseline removal prior to fitting peaks, while GlyQ-IQ (Kronewitter et al. (2014)) was written cleaner Fourier Transform MS (FTMS) that does not require such a baseline removal step. Tools that build on the THRASH implementation from Decon2LS (Jaitly et al. (2009); Yu et al. (2013); Maxwell et al. (2012)) are unable to deal with variable baseline noise or extreme dynamic range.

Each tool also has its own format for defining glycan structures or compositions, some even bundling a large database with their software to remove the burden from the user to build a list of candidates themselves (Yu et al. (2013); Kronewitter et al. (2014); Goldberg et al. (2009)) while others define methods for building glycan databases as part of the program (Maxwell et al. (2012); Ceroni et al. (2008)). Many of these tools are designed for specific glycan subclass such as N-glycans or glycosaminoglycans, limiting their vocabulary of possible monosaccharides to just those commonly found in that subclass (Yu et al. (2013); Kronewitter et al. (2014); Peltoniemi et al. (2013); Goldberg et al. (2009)). Often, these tools are tailored for analysis of a particular derivatization state, adduction conditions, or neutral loss pattern (Yu et al. (2013); Peltoniemi et al. (2013); Maxwell et al. (2012)). Work has been done to construct a standardized namespace and representation for glycans, including both structures and compositions (Tiemeyer et al. (2017); Campbell et al. (2014)). This data is publicly accessible, including a programmatic query interface using SPARQL over HTTPS (Aoki-Kinoshita et al. (2015)). Tools that can communicate with these services have the potential to lead researchers to find deeper connections from cross-referenced information, and other researchers can more readily find and use their work.

These spectral processing and glycan library properties are reflected in the scoring function that each program uses to discriminate glycan signal from the background noise and contaminants. Several methods have been developed using different facets of the observed data. Yu et al. (2013) used the isotopic pattern goodness-of-fit while Peltoniemi et al. (2013) used intensity features of associated  $MS^n$ scans to. Kronewitter et al. (2014) combined several features of the  $MS^1$  evidence, including elution profile peak shape goodness-of-fit, isotopic fit, mass accuracy, scan count, and in-source fragmentation correlation. Some of these methods are well-defined and invariant from instrument to instrument in this era of high resolution mass spectrometry, but others are tightly coupled to the experimental equipment. Missing from this list are methods to target a glycan's intrinsic properties, such as charge state distribution or facility in acquiring adducts, which can increase the number of spurious assignments when not accounted for. We propose a new scoring function which is able to combine those properties which are independent of experimental setup with these glycan-aware features.

As observed by Goldberg  $et\ al.\ (2009)$ , there is also value in including related glycan composition identifications in how much confidence one assigns to a another glycan composition assignment. They use a method to exploit the known biosynthetic rules of N-glycans to connect peaks in a MALDI spectrum which could be assigned to a particular N-glycan by intact mass alone. Their method using the maximum weighted subgraph of the biosynthetic network in one of their three had demonstrably better performance than chance with their expert system annotation method. Kronewitter  $et\ al.\ (2014)$  considered a similar idea with more emphasis on handling in-source fragmentation observed in LC-MS and LC-MS/MS experiments.

We extend this notion of a glycan family to cover more sectors of the biosynthetic landscape which we term "neighborhoods", and present an algorithm for learning the importance of each neighborhood from observed data, which can in turn be used to improve glycan composition assignment performance. We also apply our method using three different glycan composition search spaces to show how the underlying database can influence results.

Table 1: Glycan Composition Rule Table

Monosaccharide	Lower Limit	Upper Limit	Constraints
HexNAc	2	9	
$\mathbf{Hex}$	3	10	
$\mathbf{Fuc}$	0	4	$\mathbf{HexNAc} > \mathbf{Fuc}$
$\mathbf{NeuAc}$	0	5	$(\mathbf{HexNAc} - 1) > \mathbf{NeuAc}$

Table 2: Samples Used

Sample Name	Instrument	Derivatization	Adduction	Source
20150930-06-AGP	QTOF	Native	Formate (1)	Khatri <i>et al.</i> (2016a)
20141031-07-Phil-82	QTOF	Native	Formate (3)	Khatri et al. (2016a)
20141103-02-Phil-BS	QTOF	Native	Formate (3)	Khatri <i>et al.</i> (2016a)
20151002-02-IGG	QTOF	Native	Formate (2)	Khatri <i>et al.</i> (2016b)
$20141128-11-Phil-82^{1}$	QTOF	Deutero-reduced and	Ammonium (3)	Khatri et al. (2016a)
		Permethylated		
AGP-DR-Perm-glycans-1 <sup>1</sup>	Orbitrap	Deutero-reduced and	Ammonium (3)	Khatri et al. (2016a)
		Permethylated		
AGP-permethylated-2ul-inj-	Orbitrap	Reduced and Perme-	Ammonium (3)	Khatri et al. (2016a)
55-SLens <sup>1</sup>		thylated		
Perm-BS-070111-04-Serum <sup>1</sup>	Orbitrap	Reduced and Perme-	Ammonium (3)	Yu et al. (2013); Hu and Mechref (2012)
		thylated		

<sup>&</sup>lt;sup>1</sup> Included  $MS^n$  Scans

## 2 Methods

# 2.1 Glycan Hypothesis Generation

In eukaryotes, N-glycans start with a common, conserved core of **HexNAc2 Hex3**, building up to **HexNAc2 Hex9** (Stanley et al. (2009)). This structure is refined by sequentially removing monosaccharides and replacing them with more complex structures through a series of glycosylase and glycosyltransferase reactions, the enumeration of, which as shown in Akune et al. (2016), yields over a million of possible N-glycan topologies. These topologies define the geometry of the glycan, and it's protein binding properties. Neither  $MS^1$  nor collisional tandem MS of glycans can capture the full tree or graph structure of an N-glycan, so we reduced the topology to a count of each type of residue, a composition.

Starting with the core motif, we generated all combinations of monosaccharides ranging between the limits in Table 1 to build a glycan composition database, which produced 1240 distinct compositions. To perform a side-by-side comparison we also extracted the glycan list from Yu et al. (2013) derived from the biosynthetic rules in Krambeck and Betenbaugh (2005) with 319 compositions, and another database using all N-glycans from GlyTouCan (Tiemeyer et al. (2017)) containing only [Hex, HexNAc, Fuc, Neu5Ac, sulfate], with 275 distinct compositions. As previous analysis of Influenza A virus samples detected sulfated N-glycans (Khatri et al. (2016a)), we also created a combinatorial database with up to one sulfate included, for a total of 2480 compositions. As our algorithm treats HexNAc and HexNAc(S) as distinct entities, for all monosaccharides with post-attachment substituents such as sulfate and phosphate, we detached the substituent from the core monosaccharide. Our implementation is able to interpret IUPAC trivial names and compositions thereof with standard substituent and unambiguous backbone modifications, permitting a wide range of possible glycan compositions.

#### 2.2 LC-MS Data Preprocessing

We analyzed samples from several sources, including both QTOF and Orbitrap instruments as shown in Table 2. For details on sample preparation and data acquisition, please see the source citations. We converted all datasets to mzML format (Martens et al. (2011)) using Proteowizard (Kessner et al. (2008)) without any data transforming filters. We applied a background reduction method based upon (Kaur and O'Connor (2006)), using a window length of 2 m/z. Next, we picked peaks using a simple Gaussian model and iteratively charge state deconvoluted and deisotoped using an averagine (Senko et al. (1995)) formula appropriate to the molecule under study. For native glycans, the formula was H 1.690 C 1.0 O 0.738 N 0.071, for permethylated glycans, the formula was H 1.819 C 1.0 O 0.431 N 0.042. We used an iterative approach which combines aspects of the dependence graph method (Liu et al. (2010)) and with subtraction. All samples were processed using a minimum isotopic fit score of 20 with an isotopic strictness penalty of 2.

#### 2.3 Chromatogram Aggregation

We clustered peaks whose neutral masses were within  $\delta_{mass} = 15$  parts-per-million error (PPM) of each other. When there were multiple candidate clusters for a single peak, we used the cluster with the lowest mass error. Next, we sorted each cluster by time, creating a list of aggregated chromatograms. To account for small mass differences, we found all chromatograms which are within  $\delta_{mass} = 10$  PPM of each other and which overlap in time and merge them. These mass tolerances were selected empirically, and can be adjusted as needed by the user.

# 2.4 Glycan Composition Matching

For each chromatogram, we searched the glycan database for compositions whose masses were within  $\delta_{mass}=10$  PPM for QTOF data, 5 PPM for FTMS data. We merged all features matching the same composition. Then, for each mass shift combination, we searched the glycan database for compositions whose neutral mass were within  $\delta_{mass}$  of the observed neutral mass - mass shift combination mass, followed by another round of merging chromatograms with the same assigned composition. We reduced the data by splitting each feature where the time between sequential observation was greater than  $\delta_{rt}=0.25$  minutes and removed features with fewer than k=5 data points. The same chromatogram may be given multiple assignments and designated multiple mass shifts, and chromatograms without glycan assignments may use chromatograms with glycan assignments as mass shifted components. This ambiguity information was propagated through each merge and split step. We termed these remaining assigned and unassigned chromatograms candidate features.

#### 2.5 Feature Evaluation

We computed several metrics to estimate how distinguishable each candidate feature was from random noise. The metrics are mentioned in List 1, but for more information see Section S1.

#### List 1: Chromatographic Feature Metrics

- 1. Goodness-of-fit of chromatographic peak shape to a model function (Yu and Peng (2010); Kronewitter et al. (2014)).
- 2. Goodness-of-fit of isotopic pattern to glycan composition weighted by peak abundance (Maxwell et al. (2012)).
- 3. Observed charge states with respect to glycan composition and mass.
- 4. Time gap between  $MS^1$  observations detecting missing peaks and interference.
- 5. Adduction states with respect to glycan composition and mass.

These metrics are bounded in [0,1). Any observation for which any metric was observed below 0.15 was discarded as having insufficient evidence for consideration. The observed score s for each candidate feature is the sum of the logit-transformation of these metrics. This produces a single value bounded in  $[0,\infty)$ , whose distribution we assume is asymptotically normal. A value of s < 8 reflects a low confidence match, with confidence increasing as s does. As these metrics are tied to reliable detection of the the glycan by the mass spectrometer, they are dependent upon glycan abundance and sample quality and the resolution of the mass spectrometer used.

# 2.6 Glycan Composition Network Smoothing

Evidence for individual glycan compositions can often be enough to claim that composition had been detected. Lower abundance may score poorly in one or more features, leading to the glycan composition being discarded. Other methods have demonstrated that it is advantageous to use relationships between glycans based on biosynthetic or structural rules to adjust the score of a single glycan assignment (Goldberg et al. (2009); Kronewitter et al. (2014)). We propose a method based on Laplacian regularized least squares (Belkin et al. (2006)) to use evidence from glycan compositions related over a network to smooth it's evaluation of glycan composition feature matching.

#### 2.6.1 Glycan Composition Graph

For each database of theoretical glycan compositions we create, we define each composition to be a coordinate vector in a  $\mathcal{Z}^{+c}$  space where c is the number of components in any glycan composition, and represented by a node in an undirected glycan composition graph  $\mathcal{G}$ . Under this interpretation, we can compute the  $L_1$ -distance between two glycan compositions. For any two glycan compositions  $g_u, g_v$ , if  $L_1(g_u, g_v) = 1$  we add an edge connecting  $g_u$  and  $g_v$  to  $\mathcal{G}$  with weight w = 1.

Name	Bounds
High Mannose	$\mathbf{HexNAc} = 2 \land \mathbf{Hex} \in [3, 10] \land \mathbf{NeuAc} = 0$
Hybrid	$\mathbf{HexNAc} \in [2,4] \land \mathbf{Hex} \in [2,6] \land \mathbf{NeuAc} \in [0,2]$
Bi-Antennary	$\mathbf{HexNAc} \in [3,5] \land \mathbf{Hex} \in [3,6] \land \mathbf{NeuAc} \in [1,3]$
Asialo-Bi-Antennary	$\mathbf{HexNAc} \in [3,5] \land \mathbf{Hex} \in [3,6] \land \mathbf{NeuAc} \in [0,1]$
Tri-Antennary	$\mathbf{HexNAc} \in [4,6] \land \mathbf{Hex} \in [4,7] \land \mathbf{NeuAc} \in [1,4]$
Asialo-Tri-Antennary	$\mathbf{HexNAc} \in [4, 6] \land \mathbf{Hex} \in [4, 7] \land \mathbf{NeuAc} \in [0, 0]$
Tetra-Antennary	$\mathbf{HexNAc} \in [5,7] \land \mathbf{Hex} \in [5,8] \land \mathbf{NeuAc} \in [1,5]$
Asialo-Tetra-Antennary	$\mathbf{HexNAc} \in [5,7] \land \mathbf{Hex} \in [5,8] \land \mathbf{NeuAc} \in [0,0]$
Penta-Antennary	$\mathbf{HexNAc} \in [6, 8] \land \mathbf{Hex} \in [6, 9] \land \mathbf{NeuAc} \in [1, 5]$
Asialo-Penta-Antennary	$\mathbf{HexNAc} \in [6, 8] \land \mathbf{Hex} \in [6, 9] \land \mathbf{NeuAc} \in [0, 0]$
Hexa-Antennary	$\mathbf{HexNAc} \in [7, 9] \land \mathbf{Hex} \in [7, 10] \land \mathbf{NeuAc} \in [1, 6]$
Asialo-Hexa-Antennary	$\mathbf{HexNAc} \in [7, 9] \land \mathbf{Hex} \in [7, 10] \land \mathbf{NeuAc} \in [0, 0]$
Hepta-Antennary	$\mathbf{HexNAc} \in [8, 10] \land \mathbf{Hex} \in [8, 11] \land \mathbf{NeuAc} \in [1, 7]$
Asialo-Hepta-Antennary	$\mathbf{HexNAc} \in [8, 10] \land \mathbf{Hex} \in [8, 11] \land \mathbf{NeuAc} \in [0, 0]$

Table 3: N-Glycan Neighborhoods

#### **Neighborhood Definition** 2.6.2

Our definition of distance connects glycan compositions which differ by a single monosaccharide, but we can assert larger collections of glycan compositions are related. We define neighborhoods for N-glycans using intervals over monosaccharide counts defined in Table 3. These neighborhoods are arranged to span particular epitopes or biosynthetically related subtypes of N-glycans, such as sialylation state or branching pattern.

Glycan compositions may belong to zero or more neighborhoods, as there are unusual glycan compositions which do not satisfy any neighborhood's rules, and several neighborhoods intentionally overlap to express broad relationships between groups.

We define a matrix **A** as an  $n \times k$  matrix where  $A_{i,k}$  to be the degree to which  $g_i$  belongs kth neighborhood:

$$A_{i,k} = \frac{1}{|\text{neighborhood}_k|} \sum_{g^* \in \text{neighborhood}_k} L_1(g_i, g^*)$$
(1)

To reduce the impact of neighborhood size on the elements of A, the columns of A are first normalized to sum to 1, and then the rows of **A** are normalized to sum to  $1^1$ .

We assume that members of the same neighborhood will share a central tendency,  $\tau$ .

#### Laplacian Regularization

We combine the observed score s and the structure of  $\mathcal{G}$  to estimate a smoothed score  $\phi$  that combines the evidence for each individual glycan composition as well as its relatives. As s is the size of the set of observed glycan composition p while  $\phi$  is

of size n, we partition  $\phi$  into a block vector  $\begin{bmatrix} \phi_o \\ \phi_m \end{bmatrix}$  with dimensions  $\begin{bmatrix} p \\ n-p \end{bmatrix}$ . Let  $\mathbf{L}$  be the weighted Laplacian matrix of  $\mathcal{G}$ , which is an  $n \times n$  matrix. To ensure  $\mathbf{L}$  is invertible, we add  $\mathbf{I}_n$  to  $\mathbf{L}$ . We partition  $\mathbf{L}$  into blocks  $\begin{bmatrix} \mathbf{L_{oo}} & \mathbf{L_{om}} \\ \mathbf{L_{mo}} & \mathbf{L_{mm}} \end{bmatrix}$ . We also partition  $\mathbf{A}$  into  $\begin{bmatrix} \mathbf{A}_o \\ \mathbf{A}_m \end{bmatrix}$  and  $\tau_o = \mathbf{A}_o \tau$ ,  $\tau_m = \mathbf{A}_m \tau$ .

We find the  $\phi$  that minimizes the expression

$$S(\mathbf{L}, \phi, \tau) = \begin{bmatrix} \phi_o - \tau_o, & \phi_m - \tau_m \end{bmatrix} \begin{bmatrix} \mathbf{L_{oo}} & \mathbf{L_{om}} \\ \mathbf{L_{mo}} & \mathbf{L_{mm}} \end{bmatrix} \begin{bmatrix} \phi_o - \tau_o \\ \phi_m - \tau_m \end{bmatrix}$$
(2)

$$\ell = (\mathbf{s} - \phi_{\mathbf{o}})^t (\mathbf{s} - \phi_{\mathbf{o}}) + \lambda \mathcal{S}(\mathbf{L}, \phi, \tau)$$
(3)

where  $\lambda$  controls how much weight is placed on the network structure and  $\tau$ .

To obtain the optimal  $\phi$ , we take the partial derivative of  $\ell$  w.r.t  $\phi_m$ 

$$0 = \frac{\partial \ell}{\partial \phi_{m}} \left( (\mathbf{s} - \phi_{\mathbf{o}})^{t} (\mathbf{s} - \phi_{\mathbf{o}}) + \lambda \mathcal{S}(\mathbf{L}, \phi, \tau) \right)$$
(4)

$$\hat{\phi}_m = -\mathbf{L_{mm}}^{-1} \mathbf{L_{mo}} (\phi_o - \tau_o) + \tau_m \tag{5}$$

<sup>&</sup>lt;sup>1</sup> The stated reduction is not well tested, and the change may well be minimal because all that really happens is the weight of the column for each row is weighted by a shrinking function of column size. It may be better if we don't manipulate A at all.

NI: 11 1 1	Phil-BS			Serum		
Neighborhood $\tau$	Combinatorial + Sulfate	glySpace	Krambeck	Combinatorial	glySpace	Krambeck
high-mannose	18.010	16.949	17.001	20.148	19.568	19.720
hybrid	13.378	15.418	13.700	19.889	18.963	20.056
bi-antennary	0.000	0.000	0.000	16.654	13.808	17.594
asialo-bi-antennary	14.135	12.708	12.913	22.221	21.852	21.827
tri-antennary	0.000	0.000	0.000	24.600	19.313	23.644
asialo-tri-antennary	14.476	10.918	10.823	20.474	20.514	19.764
tetra-antennary	0.000	0.000	0.000	18.630	18.438	17.675
asialo-tetra-antennary	14.351	0.000	6.186	4.118	5.437	2.568
penta-antennary	0.000	0.000	0.000	12.822	14.383	11.681
asialo-penta-antennary	11.582	0.000	4.892	0.000	0.000	0.000
hexa-antennary	0.000	0.000	0.000	0.000	0.000	0.000
asialo-hexa-antennary	11.089	0.000	6.612	0.000	0.000	0.000
hepta-antennary	0.000	0.000	0.000	0.000	0.000	0.000
asialo-hepta-antennary	3.120	0.000	1.351	0.000	0.000	0.000
$\hat{\lambda}$	0.99	0.99	0.99	0.99	0.99	0.99
$\hat{\gamma}$	11.39	18.35	14.42	19.32	19.37	20.67

Table 4: Estimated values of smoothing parameters  $\tau$ ,  $\lambda$ , and  $\gamma$  for each dataset and database

and w.r.t.  $\phi_o$ 

$$0 = \frac{\partial \ell}{\partial \phi_o} \left( (\mathbf{s} - \phi_o)^t (\mathbf{s} - \phi_o) + \lambda \mathcal{G}(\mathbf{L}, \phi, \tau) \right)$$
 (6)

$$\hat{\phi}_o = \left[ \mathbf{I} + \lambda \left( \mathbf{L_{oo}} - \mathbf{L_{om}} \mathbf{L_{mm}^{-1}} \mathbf{L_{mo}} \right) \right]^{-1} (\mathbf{s} - \tau_o) + \tau_o$$
(7)

To use this method, we must provide values for  $\lambda$  and  $\tau$ . While these values could be chosen based on the expectations of the user for a given experiment, we provide an algorithm for selecting their values in Section S 4. These methods use the topology of the glycan composition graph and the distribution of observed scores, and cannot fully capture boundary cases or related but disconnected parts of the graph.

## 3 Results

We demonstrated the performance of our algorithm using released influenza hemagglutinin data set 20141103-02-Phil-BS and a serum glycan data set Perm-BS-070111-04-Serum. Please refer to section S5 for all other datasets. For each comparison, the unregularized case is not smoothed, effectively  $\lambda = 0$ , the partially regularized case uses the grid search fitted values of  $\tau$  but uses a fixed  $\lambda = 0.2$ , and the fully regularized case uses the grid search fitted values of both  $\tau$  and  $\lambda$ .

#### 3.1 Chromatogram Assignment Performance for 20141103-02-Phil-BS

The fitted parameters for the network constructed for 20141103-02-Phil-BS are shown in Table 4. The assigned chromatograms are shown in Figure 1. We observe up to seven branch structures in this sample, consistent with these N-glycans being derived from an avian context (Stanley et al. (2009); Khatri et al. (2016a)).

The comparison of assignment performance with differing degrees of smoothing for each database are shown in Figure 2 and Table 5. We observed the greatest number of assignments using the combinatorial database including sulfate.

#### 3.2 Chromatogram Assignment Performance for Perm-BS-070111-04-Serum

The fitted parameters for the network constructed for *Perm-BS-070111-04-Serum* are shown in Table 4. The assigned chromatograms are shown in Figure 3.

The comparison of assignment performance with differing degrees of smoothing is shown in Figure 4. We observe the greatest number of total true identifications using the partially regularized Combinatorial database. However, the Combinatorial database also has many more false positives, with a ROC AUC of 0.817. These false positives do not appear in the biosynthetically constraind Krambeck database which maximizes its ROC AUC in the partially regularized condition at 0.883. After removing all ambiguous matches, the Krambeck database also has nearly the same number of true matches as the Combinatorial database.

Figure 1: Chromatogram Assignments and Quantification for 20141103-02-Phil-BS Using the Combinatorial + Sulfate database.

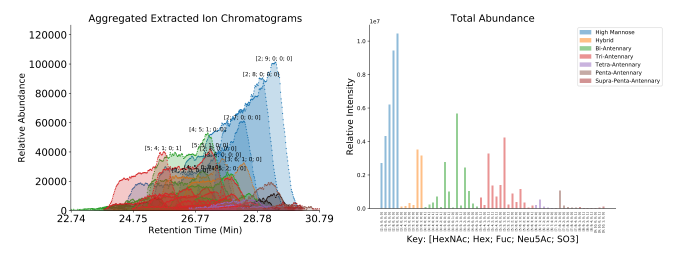
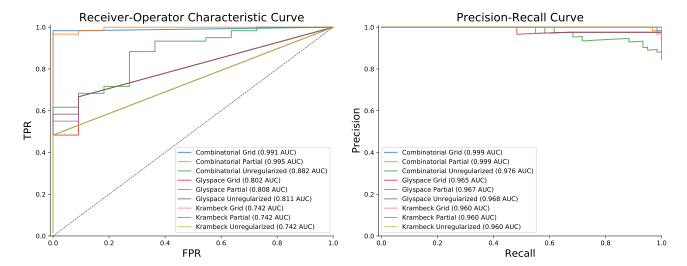


Figure 2: Performance Comparison with and without Network Smoothing for 20141103-02-Phil-BS



# 4 Discussion

We demonstrated that the regularization method improved the sensitivity and specificity of glycan composition assignment for LC-MS based experiments. The method used similar assumptions about the importance of common substructural elements of N-glycans to Goldberg  $et\ al.\ (2009)$ , but we extend this concept with the addition of a procedure for learning the relationship strengths and use broader groups of structures.

The experimental results from the original analysis of 20141103-02-Phil-BS and 20141031-07-Phil-82 82 demonstrated that while both strains expressed predominantly high-mannose glycosylation, 20141103-02-Phil-BS expressed more larger complex-type structures (Khatri et al. (2016a)). In our findings shown in Figure 1, we recapitulate these results while reducing the number of false assignments, Table 5. There are substantial differences in both the mass spectral processing and scoring schemes which contribute to these results, but the regularization procedure is responsible for recovering many low abundance features from this comparison. As these samples are derived from chicken eggs, we have observed larger branching patterns than are observed in normal mammalian tissue (Stanley et al. (2009)). There is evidence for this in the 20141103-02-Phil-BS with **HexNAc9 Hex10**-based compositions suggesting a seven branch pattern, though this cannot be determined without high quality  $MS^n$ data. The  $\tau$  fit for Phil-BS (shown) and Phil-82 (supplement) have smaller values in the neighborhoods of their largest glycan compositions as these features tended to be low in abundance and not high scoring in their own right, but were partially supported by the overlap with the next largest neighborhood, as expected. We observed the best performance with the Combinatorial + Sulfate database, which produced more than half-again as many true matches than the other two databases. It produced several false matches as well, but the smoothing process removed these while boosting the score of other low abundance matches which were consistent with higher scoring matches.

The Krambeck database performed identically in all smoothing conditions as it was only able to match the common species, not including cases that were multiply fucosylated or sulfated. It had no false matches ranked alongside its true

Name	ROC AUC	True Matches <sup>1</sup>
Combinatorial Unregularized	0.882	56
Combinatorial Partial	0.995	57
Combinatorial Grid	0.991	57
GlySpace Unregularized	0.811	40
GlySpace Partial	0.808	38
GlySpace Grid	0.802	31
Krambeck Unregularized	0.742	28
Krambeck Partial	0.742	28
Krambeck Grid	0.742	28
Khatri et al. (2016a)	-	46

<sup>&</sup>lt;sup>1</sup> Selected at  $\phi_o > 5.0$ 

Table 5: Performance Comparison for 20141103-02-Phil-BS

Name	ROC AUC	True Matches $^1$	Non-Ambiguous Matches
Combinatorial Unregularized	0.677	86	60
Combinatorial Partial	0.817	87	61
Combinatorial Grid	0.811	87	61
GlySpace Unregularized	0.788	59	50
GlySpace Partial	0.802	60	51
GlySpace Grid	0.808	60	51
Krambeck Unregularized	0.866	70	59
Krambeck Partial	0.883	70	59
Krambeck Grid	0.883	70	59
Yu et al. (2013)	-	$72^{2}$	59

<sup>&</sup>lt;sup>1</sup> Selected at  $\phi_o > 5.0$ 

Table 6: Performance Comparison for Perm-BS-070111-04-Serum

matches so smoothing could not change its performance. The glySpace-derived database produced more true matches, but also lacked some of these more fucosylated and complex compositions. Some of the compositions included by the glySpace-derived database were lower scoring, but the chosen value of  $\gamma$  for that database was greater than 18, causing the fitted values of  $\tau$  omit the larger, less abundant complex-type N-glycans. This caused smoothing to lower the scores of these real matches rather than raise them, as with the Combinatorial + Sulfate database.

As we show in Figure 4, regularization improves the predictive performance of the identification algorithm on Perm-BS-070111-04-Serum for all databases. We reproduce the majority of the glycan assignments from Yu et al. (2013), but the ambiguity caused by ammonium adduction as shown in Figure 3c makes a direct comparison of composition assignment lists difficult. Our algorithm requires a minimum amount of MS1 information in order to compute a score, which some of the assignments in the original published results do not possess, which are omitted from the count in Table 6. After accounting for ambiguity, we were able to assign all of the compositions previously reported using the Krambeck database, which was used by Yu et al. (2013), and with the combinatorial database. The glySpace-derived database did not contain all of these compositions, but performed competitively with the combinatorial database's ROC AUC. The combinatorial database contained a small number of glycan compositions which were not in Krambeck but which were consistent with other glycan compositions observed nearby in retention time. The combinatorial database also benefited most substantially from smoothing, discarding many false positives while retaining many more true positives at the same false positive rate compared to the other databases. These invalid glycan compositions can match LC-MS features at any point in the elution profile, though in this dataset the majority of these matches appear to be in the time range between 10 and 22 minutes, and similar glycan compositions that are biosynthetically valid elute later on in the experiment. Therefore a for a retention-time aware approach to evaluating glycan composition assignments, as described in Hu et al. (2016) could also be useful, but this is likely dependent upon the experimental workup and separation technique used.

While the biosynthetically constrained Krambeck database performed better on *Perm-BS-070111-04-Serum*, it did not contain all of the reasonably assignable glycan compositions, and it performed poorly on 20141103-02-Phil-BS with a false negative rate of 50% compared to the combinatorial database. This is because the necessary enzymatic pathways were either not considered in the original authors' model because either the enzyme was excluded for simplicity (Krambeck *et al.* (2009)) or because the particular enzymes used were not within the scope of the model used (Spiro and Spiro (2000); Ichimiya *et al.* (2014)). This highlights the importance of selecting a good reference database, though a post-processing step such as the we

<sup>&</sup>lt;sup>2</sup> This count only includes those cases where sufficient MS1 scans were available for a direct comparison

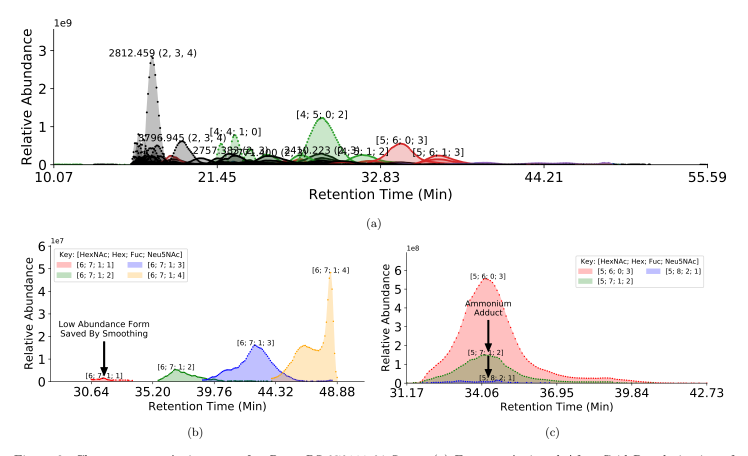


Figure 3: Chromatogram Assignments for *Perm-BS-070111-04-Serum* (a) Features Assigned After Grid Regularization of *Perm-BS-070111-04-Serum* (b) Low scoring features which may be discarded based on individual evidence alone may be more reasonable to accept given evidence from related composition, such as our network smoothing method (c) This sample contains heavy ammonium adduction which introduces ambiguity in intact mass based assignments

described here can help mitigate using too large a database, but not a too small one.

In this work, we used the same network neighborhood imposed over different underlying sets of composition nodes, and the connectivity of those networks did not take into account the biosynthetic process. It may be possible to obtain better performance by defining network connectivity according to enzymatic relationships. This may also alter how the neighborhoods are defined and how  $\bf A$  is parameterized, and in turn how  $\tau$  is learned. Similarly, this procedure depends upon the scoring functions used, so selecting another set of functions for the data to fit may lead to different parameter values.

Lastly, while these case studies have demonstrated the algorithm's ability to learn network parameters from the data, an expert can define  $\tau$  and  ${\bf A}$  themselves or obtain a model fitted on related data and apply it directly without a fitting step. An expert could use this model specification to impose prior beliefs on the evaluation process, and adjust  $\lambda$  to control the importance of the these beliefs. Similarly, one could also use the derivation of  $\hat{\phi}_m$  to estimate the score for an unobserved glycan composition, given  ${\bf A}$  and  $\tau$ .

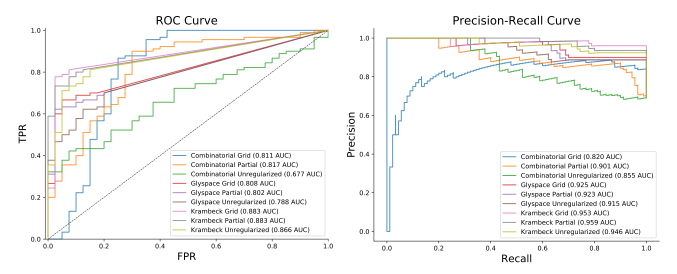
We used our glycoinformatics toolkit to produce a richer abstraction of glycans and monosaccharides, including producing standard-compliant textual representations of these structures and compositions. We produced a text file containing all of the glycan compositions found in the Krambeck and Combinatorial database but not the glySpace-derived database in the above samples (see supplemental information XXX), and have submit it to GlyTouCan (Tiemeyer et al. (2017)) for registration so that future researchers can use these structures.

# 5 Conclusions

In this study, we demonstrated the advantages of our application of Laplacian Regularization to smooth LC-MS assignments of glycan compositions across multiple experimental protocols (Hu and Mechref (2012); Khatri *et al.* (2016a)). Our algorithm's performance is competitive with existing tools for analyzing the same type of data, with the added benefit of more flexible evaluation process and broader range of understood monosaccharides. Our tools integrate with glySpace and allows users to leverage existing glycomics repositories to build databases where applicable.

All of the methods demonstrated in this paper are available as part of the open source, cross-platform glycomics and glycoproteomics software GlycReSoft, freely available at http://www.bumc.bu.edu/msr/glycresoft/.

Figure 4: Performance Comparison with and without Network Smoothing for Perm-BS-070111-04-Serum



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