Sequence analysis

Discovery of phosphorylation motif mixtures in phosphoproteomics data

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ABSTRACT

Motivation: Modification of proteins via phosphorylation is a primary mechanism for signal transduction in cells. Phosphorylation sites on proteins are determined in part through particular patterns, or motifs, present in the amino acid sequence.

Results: We describe an algorithm that simultaneously discovers multiple motifs in a set of peptides that were phosphorylated by several different kinases. Such sets of peptides are routinely produced in proteomics experiments. Our motif-finding algorithm uses the principle of minimum description length to determine a mixture of sequence motifs that distinguish a foreground set of phosphopeptides from a background set of unphosphorylated peptides. We show that our algorithm outperforms existing motif-finding algorithms on synthetic datasets consisting of mixtures of known phosphorylation sites. We also derive a motif specificity score that quantifies whether or not the phosphoproteins containing an instance of a motif have a significant number of known interactions. Application of our motif-finding algorithm to recently published human and mouse proteomic studies recovers several known phosphorylation motifs and reveals a number of novel motifs that are enriched for interactions with a particular kinase or phosphatase. Our tools provide a new approach for uncovering the sequence specificities of uncharacterized kinases or phosphatases.

Availability: Software is available at http://cs.brown.edu/people/braphael/software.html.

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Supplementary information: Supplementary data are available at Bioinformatics online.

1 INTRODUCTION

Modification of proteins via phosphorylation is a primary mechanism for signal transduction in cells. Members of signaling pathways include kinases that phosphorylate proteins at tyrosine, serine or threonine residues and phosphatases that dephosphorylate proteins. Both kinases and phosphatases recognize their substrates in part through patterns, or motifs, present near the phosphorylation site in the amino acid sequence of the substrate. A number of such motifs have been identified and are recorded in databases, such as PhosphoSite Plus (www.phosphosite.org), Scansite (Obenauer et al., 2003), Mini-motif Miner (Balla et al., 2006) and PhosphoMotif Finder (Amanchy et al., 2007). However, many phosphorylation sites are not associated with a kinase and/or phosphatase. For example, over 54% of the 11 176 documented phosphorylation modifications in the Human Protein Reference Database (HPRD) (Peri et al., 2003) have no known upstream enzyme. Part of the reason for this knowledge gap is that discovery of phosphorylation sites is presently more efficient than identification of the kinase that phosphorylates the site. Traditional experimental methods that measure kinase–substrate interaction (such as co-immunoprecipitation or fluorescence resonance energy transfer) require some hypothesis as to the kinase involved in the phosphorylation. Computational methods to predict kinase–substrate interactions from structural information (Brinkworth et al., 2003) or from other known kinase substrates (Blom et al., 2004) have also been introduced, but such additional information is not always available.

Recently, algorithms have emerged to accurately predict substrates of some well-characterized kinases. NetworKIN (Linding et al., 2007) integrates phosphorylation motifs and protein interaction networks, relying on well-characterized kinase motifs from NetPhosK (Blom et al., 2004) and Scansite (Obenauer et al., 2003). NetPhorest (Miller et al., 2008), classifies the upstream enzymes of phosphorylation sites using 125 different sequence-based classifiers, and NetworKIN now integrates these classifiers. For algorithms such as these, knowledge of motifs for additional kinases and phosphatases would be useful.

A growing source of phosphorylation sites is phosphoproteomic experiments that simultaneously measure hundreds to thousands of phosphorylated residues in a cell (Cao et al., 2007; Olsen et al., 2006; Rush et al., 2005; Wolf-Yadlin et al., 2006). In a phosphoproteomics experiment, peptides containing a phosphorylated residue are identified from a purified sample via mass spectrometry (Hoffert and Knepper, 2007). The resulting phosphopeptides measured under the same experimental condition provide data for the identification of sequence motifs that indicate an interaction with a specific kinase or phosphatase that was active during the experiment.

The motif-finding problem arising from phosphoproteomics data is the following. Given a collection of peptides each containing

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a specific phosphorylated residue (serine, threonine or tyrosine), find a set of repeated patterns, or motifs, that are overrepresented in the phosphorylated peptides relative to a background set of unphosphorylated peptides. Note that all phosphorylated peptides share the phosphorylated residue, and thus the peptides can be aligned on the phosphorylated residue. This problem differs from many of the standard motif-finding problems in that the data is a mixture of motifs, representing the numerous kinase–substrate interactions that were measured in the experiment. Most motif-finding programs are optimized to solve a different problem, that of identifying and ranking relatively short motifs in a collection of long, unaligned background sequence, a problem motivated by the discovery of transcription factor binding sites (e.g. see Bailey and Elkan, 1994, 1995; Brazma et al., 1996; Buhler and Tompa, 2002; Jonassen et al., 1995; Lawrence et al., 1993; Rigoutsos and Floratos, 1998; Tompa et al., 2005). While these methods might output multiple motifs, usually in the form of a ranked list, these lists often contain variations of the same motif, and do not explicitly identify a mixture. A method called Motif-X (Schwartz and Gygi, 2005) was introduced for motif identification in aligned phosphoproteomics data and has since been employed in several studies, such as Xue et al. (2006). However, the type of motifs produced by Motif-X, patterns consisting of either a single letter at each position or a ‘wild-card’ character matching any position, are quite restrictive. Moreover, the greedy iterative approach used by Motif-X limits the motif mixtures that will be found. Finally Motif-X ignores the possibility of overlapping motifs.

We present the Motif Description Length (MoDL) algorithm for the discovery of mixtures of protein phosphorylation motifs. MoDL is based on the principle of minimum description length (MDL) (Grünwald, 2007) from information theory, and produces a set of motifs that succinctly describes the biases in sequence composition in a foreground set of phosphorylated peptides in comparison to a background set of unphosphorylated peptides. MoDL compares favorably to other published motif-finding programs including the aforementioned Motif-X, Teiresias (Rigoutsos and Floratos, 1998), MDL-Pratt (Brazma et al., 1996; Jonassen et al., 1995), MEME (Bailey and Elkan, 1994, 1995) and Gibbs Motif Sampler (Lawrence et al., 1993).

We also derive a motif specificity score (MSS) that measures whether or not the phosphopeptides containing a motif instance have a significant number of interactions with a specific kinase or phosphatase recorded in a protein–protein interaction database. A kinase/phosphatase-motif pair with a high MSS identifies a potential upstream kinase/phosphatase that targets the substrate motif. We apply our MoDL algorithm to several phosphoproteomic datasets and identify both known and novel phosphorylation motifs. Some of these motifs have high MSSs, implying that the motif describes the preference of a kinase or phosphatase to phosphorylate particular amino acid peptides. Application of MoDL and the MSS to large-scale phosphoproteomic datasets provides a new approach for uncovering the sequence specificities of uncharacterized kinases or phosphatases.

2 METHODS
Consider a collection of $N$ phosphopeptides of fixed length aligned such that the common phosphorylated residue is in the center position. Additionally, consider a background matrix obtained from residue frequencies at each position in unphosphorylated peptides of the same fixed length with the same center residue. The goal is to describe the $N$ phosphopeptides as a mixture of instances of an unknown number of motifs and sequences that contain no recognizable motifs.

Motif definitions fall into two major classes: pattern-based motifs that represent consensus sequences and profile-based motifs that represent the frequencies of each amino acid residue occurring at each position using position-specific scoring matrices (PSSMs) (Stormo, 2000). We will use the former model in the motif discovery stage, and define a motif to be a pattern consisting of conserved positions that match any letter from a specified list (denoted by brackets ‘[ ]’) and wild-card positions (denoted by ‘.’) that match any letter.¹ Each motif has a single phosphorylated residue, which we denote by an underlined character. Once motif instances are identified in a dataset, we output a profile, or PSSM, constructed from these instances. Other pattern-based motif-finding algorithms, such as Motif-X (Schwartz and Gygi, 2005) and Multiprofiler (Keich and Pevzner, 2002) similarly convert the derived motifs into a scoring matrix. The use of patterns to represent motifs during discovery might appear to be disadvantaged compared with PSSMs, but in one comprehensive comparison of motif-finding algorithms for transcription factor binding site identification (Tompa et al., 2005), the best performing algorithm was a pattern-based method.

We use description length as a metric to evaluate how well a set of motifs describes a collection of phosphopeptides. Description length is an information-theoretic quantity that measures the amount of information (in terms of bits) required to represent (or encode) a dataset. To encode a collection of phosphopeptides, we must identify each residue in each phosphopeptide as originating from either an instance of a particular motif or as originating from the background (‘non-motif’) peptides. In addition, we must encode the motifs themselves. By finding the set of motifs with MDL, we annotate the peptides as motif instances in a way that maximizes the redundancy in the representation. MoDL aims to find such a motif set (Fig. 1A).

Let $X$ be an $N \times L$ matrix containing the phosphopeptides of length $(L+1)$ with the known center phosphorylated residue removed. Let $P$ be an $20 \times L$ matrix where $P(i,j)$ gives the frequency of residue $i$ in position $j$ in a larger set of unphosphorylated peptides (also with the common center residue removed). MoDL computes the motif total number of bits required to encode both $M$ and $X$ using $M$. We first describe how to compute the description length for a particular set, $M$, of motifs for a dataset $X$, and then describe a greedy approach to approximate the MDL.

2.1 Computing description length
Our aim is to find a set of motifs $M^* = \{m_1, ..., m_k\}$ that minimizes the total number of bits required to encode both $M$ and $X$.

$$M^* = \arg \min_M \Lambda(X, M).$$

Note that since $P$ is independent of the choice of $M$, we do not explicitly encode it.

The number of bits $\Lambda(X, M)$ required to encode $X$ and $M$ is the sum of bits $\Lambda(M)$ required to encode the motif set and the number of bits $\Lambda(X|M)$ to encode the data described by the motif set. Computing $\Lambda(M)$ requires encoding the motif set $M = \{m_1, ..., m_k\}$, which we do by concatenating the encodings of the individual motifs $\Lambda(m_i)$ preceded by an encoding of the value of $k$ (assume some upper bound $K \geq k$ on the number of motifs). We assume that the peptides $x_1, ..., x_N$ are independent, and so the description length of $X$ is the sum of the description length of each peptide.

$$\Lambda(X, M) = \log_2 K + \sum_{i=1}^{k} \Lambda(m_i) + \sum_{i=1}^{N} \Lambda(x_i|M).$$

¹In most known protein phosphorylation motifs, there are few conserved positions. In contrast, many algorithms for transcription factor binding site identification assume that most positions are conserved.
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Fig. 1. Overview of the MoDL algorithm and MSS calculation. MoDL uses the description length, a measure of the amount of information (bits) required to describe the input phosphopeptides using a motif set \( M \) and the background distribution. MoDL attempts to find the optimal motif set with minimum description length. (A) With an empty motif set (i.e. no motifs), each peptide must be described explicitly from the background distribution, yielding high description length (left column). On the opposite extreme, each phosphopeptide can be described as a unique motif, but the resulting motif set yields high description length (right column). The optimal motif set includes only motifs that match several phosphopeptides, and minimizes the total description length required to represent both the motifs and the phosphopeptides (center column). After the optimal motif set is determined, the individual motifs are ranked according to the increase in description length when a motif is removed from the set. (B) Computing the MSS between a kinase and a motif group, the proteins containing a motif instance. The proteins are colored according to the motif instances they contain at one or more phosphorylation sites, and gray proteins contain no motif instances. To find the MSS for the blue motif \( \text{D..Y[S,D]} \), we consider all proteins in the motif group (blue). A kinase will have a high MSS if the number of interactions between the kinase and the motif group (solid lines) is significantly greater than the number of interactions between the kinase and proteins not in the motif group (dotted lines).

We will briefly describe how to compute \( \Lambda(m) \) and \( \Lambda(x|M) \); see Supplementary Material for a detailed derivation. Each motif \( m \) is encoded in three parts: (i) a vector that indicates whether each position is a conserved position or a wildcard position, (ii) the residues for each conserved position, described as either a list of indices (the list method) or as a 20-length binary vector (the vector method) and (iii) a vector specifying the encoding method used for each conserved position. The encoding method that requires the smallest number of bits is used for each conserved position.

To compute \( \Lambda(x|M) \), peptide \( x \) is encoded in three parts: (i) a vector specifying the motif instances that \( x \) contains (or '0' if it contains no motif instances), (ii) the background residues that are not part of any motif (encoded using \( b \)) and (iii) the conserved positions (encoded using the motif instances that \( x \) contains). It is possible for multiple motifs to represent a single conserved position if a peptide \( x \) contains more than one motif instance. We choose the motif that requires the least number of bits to represent each conserved position.

### 2.2 The MoDL algorithm

Finding the motif set \( M^* \) with MDL is complicated by the fact that the space of motif sets is very large: the number of motifs is exponential in the alphabet size 20 and the peptide width \( L \). Considering motifs that only appear in the data reduces the search space, but the number is still too large to directly compute. The MoDL algorithm builds a motif set \( M \) from an initial set of simple candidate motifs using a greedy iterative approach. Candidate motifs are single-letter motifs (motifs with only one residue in one conserved position) that appear in the phosphopeptides. The empty motif set is initialized to \( M^{(0)} = \{ \} \) at time \( t = 0 \). At iteration \( (t+1) \), we construct...
a set \( W \) of potential motif sets from the motif set \( \mathcal{M}^{(1)} \) by removing a motif from \( \mathcal{M}^{(1)} \), adding a motif to \( \mathcal{M}^{(1)} \) or merging two motifs in \( \mathcal{M}^{(1)} \). Merging two motifs \( m_i \) and \( m_j \) means taking the union of the list of residues for each conserved position \( t \) and updating the vector of conserved positions and the vector of encoding methods. We build \( W \) by performing the following five operations:

1. Remove: Remove a motif from \( \mathcal{M}^{(1)} \).
2. Add: Add a candidate motif to \( \mathcal{M}^{(1)} \).
3. Add/Remove: Add a candidate motif to \( \mathcal{M}^{(1)} \) and remove another motif from \( \mathcal{M}^{(1)} \).
4. Merge: Merge a candidate motif with a motif from \( \mathcal{M}^{(1)} \).
5. Merge/Remove: Merge a candidate motif with a motif from \( \mathcal{M}^{(1)} \) and remove another motif from \( \mathcal{M}^{(1)} \).

The motif set \( \mathcal{M}^{t+1} \in W \) with the lowest description length is chosen. We repeat these steps for \( t_{\text{max}} = 50 \) iterations, or until the description length has not decreased for \( t \) iterations (we set \( t = 10 \)). As a final step, we rank each motif by the increase in the description length when the motif is removed from the motif set \( \mathcal{M}^{(1)} \).

The worst-case running time of each iteration is \( O(K^2) \), where \( K \) is the maximum number of motifs and \( R \) is the cardinality of the set of candidate motifs. Note that since \( P \) is constructed from the data \( X \), the number of iterations \( t \) is the only user-defined parameter.

### 2.3 Computing the Motif Specificity Score (MSS)

A motif identified by MoDL suggests that there is a kinase or a phosphatase that prefers to bind to peptides containing an instance of the motif. We refer to the subset of phosphoproteins containing an instance of a motif as a motif group (proteins with multiple measured phosphopeptides are considered as one protein in the motif group). If we had lists of known substrates for each kinase and phosphatase, we would expect to find a kinase or phosphatase whose list of substrates overlapped considerably with a motif group. Since every protein has an association with itself of maximum score 1, but the majority of these edges do not signify protein–protein interaction. In this way, STRING does not provide information about autophosphorylation, a common feature of signaling pathways. Despite these difficulties, however, we find that a number of kinase/phosphatase-motif pairs give statistically significant MSSs.

### 3 RESULTS

#### 3.1 Benchmarking on synthetic data

We performed three experiments to compare MoDL’s ability to extract multiple motifs with other well-known algorithms. In the first experiment, we planted instances of two motifs, \( H.G[EV][KN]D\ldots C \ldots [CR]G \) and \( \ldots Y \ldots P \), into a set of background peptides and ran a number of motif-discovery algorithms, including Motif-X (Schwartz and Gygi, 2005), Teiresias (Rigoutsos and Floratos, 1998), MDL-Pratt (Brazma et al., 1996; Jonasson et al., 1995), the Gibbs Motif Sampler (Lawrence et al., 1993), and MEME (Bailey and Elkan, 1994, 1995). MoDL outperforms all other methods, returning a nearly perfect reconstruction of the planted motifs (see Supplementary Material). The closest competitors were MEME and the Gibbs Motif Sampler, two algorithms that use PSSMs and thus further comparison of MoDL to MEME was performed.

In the second experiment, we constructed a synthetic dataset consisting of 10 instances of each of two simple motifs, \( D.\ldots Y \) and \( [TL]\ldots PP \). For each instance, the non-conserved positions were chosen uniformly according to the background distribution. We then added 0, 5, 10, 15 and 20 peptides chosen from the background distribution to the motif instances, yielding datasets consisting of 0%, 20%, 33%, 43% and 50% background peptides. For each of these five synthetic datasets, we compared MoDL’s performance to MEME by evaluating the labelling of peptides produced by the motif-finding program to the true labelling. We used the metrics of precision, recall, the fraction of pairs of peptides with the same motif label that are both instances of one planted motif, and F-score, the harmonic mean of precision and recall. (see Supplementary Material for the calculation of precision and recall). MoDL obtains a higher average F-score over these datasets with lower SD: MoDL (0.9200 ± 0.0447) versus MEME (0.7496 ± 0.1702). Further, MoDL’s recall is 1.0 for all five datasets while MEME’s average recall is 0.8467 ± 0.1193. With 0% background sequences, MoDL’s precision is 1.0 compared with MEME’s precision of 0.8901 and even with 43% background sequences MoDL’s precision is 0.8182 with recall equal to 1.0, while MEME’s precision and recall are 0.3701 and 0.633, respectively.

In the third experiment, we compared the performance of MoDL, Motif-X, and MEME on datasets from Scansite (Obenauer et al., 2003), a collection of phosphorylation sites and PSSMs for a number of well-characterized kinases. We constructed 20 datasets from different combinations of six different Scansite tyrosine motifs ABL, EGFR, PLC\(\gamma\), SRC, FYN and LCK (see Supplementary Table S2). For every combination of three Scansite motifs from these six, we built a dataset consisting of the top 25-scoring peptides for each motif and added 75 background peptides to construct datasets with 150 peptides. In 10 out of 20 datasets, MoDL gives both higher
We applied MoDL to three phosphoproteomic datasets: human HER2 signaling (Wolf-Yadlin et al., 2006), mouse Mast Cell signaling (Cao et al., 2007) and various cancer cell lines (Rush et al., 2005). Each dataset consisted of peptides of length 13 with a measured phosphorylated tyrosine in the 7th position, and background sets were constructed from all 13mers with a tyrosine in the 7th position in the corresponding species proteome. The HER2 signaling dataset included (Wolf-Yadlin et al., 2006) two cell lines (parental ‘P’ and ‘24H’) that were stimulated separately with either epidermal growth factor (EGF) or Heregulin (HRG). See Supplementary Material for further details.

For each dataset, MoDL returned a ranking of one or two phosphorylation motifs (Table 1). We compared these motif sets to the motifs returned by Motif-X (Schwartz and Gygi, 2005), another program specifically designed to find phosphorylation motifs in aligned datasets. Briefly, Motif-X considers statistically significant residues in particular positions (called residue/position pairs) according to a binomially distributed model. The algorithm is divided into a motif-building step and a data reduction step. In the motif-building step, statistically significant residue/position pairs are identified by a greedy recursive search. In the set reduction step, all peptides that contain an instance of the motif identified in the motif-building step are removed. These steps are repeated until the motif-building step fails to return a motif due to the lack of statistically significant residue/position pairs. The Motif-X score is the negative logarithm of the P-value for each motif. Note that Motif-X prunes the dataset after a motif is discovered by removing all peptides containing instances of a motif. Because of this pruning, the score of a motif found at a later iteration could be artificially low if some of its instances were removed earlier.

In all datasets, MoDL returned higher scoring motifs according to Motif-X’s score. For example, in the Mast cell dataset, the highest scoring motif found by MoDL, [DE]..Y [ADENSTY], exceeds Motif-X’s high-scoring motif D..Y by 50 orders of magnitude. We emphasize that MoDL does not explicitly optimize the Motif-X score.

The Motif-X score quantifies the significance of a single motif, while MoDL optimizes the expressiveness of a set of motifs. We therefore examined the compression of each motif set \( \mathcal{M} \), defined as the difference \( \Delta(X,\{|\})=\Lambda(X,\mathcal{M}) \) in the number of bits required with a measured phosphorylated tyrosine in the 7th position, and background sets were constructed from all 13mers with a tyrosine in the 7th position in the corresponding species proteome. The HER2 signaling dataset included (Wolf-Yadlin et al., 2006) two cell lines (parental ‘P’ and ‘24H’) that were stimulated separately with either epidermal growth factor (EGF) or Heregulin (HRG). See Supplementary Material for further details.

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to encode the data with no motifs and the number of bits required to encode the data with $\mathcal{M}$. Not surprisingly, in all cases MoDL produced motif sets with larger compression than Motif-X. In some cases, such as the HER2 HRG datasets, the differences are quite large.

The NPM-ALK motif $H.G[EV][KN]P\_Y.C..[CR]G$ returned by MoDL has the largest number of conserved positions of all motifs discovered in the studied datasets. This motif corresponds to a linker peptide between zinc-finger domains (Jantz and Berg, 2004), and five zinc-finger proteins contain multiple instances of the motif: ZNF670, ZNF91, TIP20, ZNF24 and ZNF264. Additionally, instances of $H.G[EV][KN]P\_Y.C..[CR]G$ appear in 11 peptides from the five zinc-finger proteins that were not measured phosphorylation sites. While there is evidence of serine and threonine phosphorylation in the linker peptides (Jantz and Berg, 2004), our result indicates that tyrosine residues are also phosphorylated in the linker peptides (Jantz and Berg, 2004). In the mast cell dataset, the highest scoring motif $[DE]..[IL]..[IL]Y$ is the same motif that has a high MSS for LYN (1.7007). This MSS is significant with a false discovery rate of 0.15, 8 out of 11 proteins in the dataset have interactions recorded in STRING, we predict that other proteins in the motif group are also substrates for ABL. In particular, GRF1 is one of the eight proteins in the $Y$..$P$ motif group in the HER2 24H HRG dataset. There is no recorded interaction with ABL in STRING. However, the BCR-ABL fusion protein is known to phosphorylate GRF1 at the site Y1106 (Goss et al., 2006), the same site measured in the experiment. Another motif found by MoDL is $[DENS][DNPRS].Y$ which has a high MSS for PTPN11, a known LYN antagonist in breast cancer metastasis (Coopman et al., 2000).

### 3.3 Motif Validation on Experimental Data

For each motif returned by MoDL, we computed the MSS for kinases and phosphatases present in the underlying signaling networks to predict upstream enzymes that target the motif groups. See Supplementary Table S4 for the list of kinases and phosphatases tested.

Three of the motifs had significant MSSs in the HER2 datasets (Table 2). The motifs $Y..P$ and $[PV]$ have high MSSs for ABL kinase, which is part of the known ABL consensus motif $A.V[TV][YAAP]$ (Songyang and Cantley, 1995). In addition to the known interactions recorded in STRING, we predict that other proteins in the motif group are also substrates for ABL. In particular, GRF1 is one of the eight proteins in the $Y$..$P$ motif group in the HER2 24H HRG dataset. There is no recorded interaction with ABL in STRING. However, the BCR-ABL fusion protein is known to phosphorylate GRF1 at the site Y1106 (Goss et al., 2006), the same site measured in the experiment. Another motif found by MoDL is $[DENS][DNPRS].Y$ which has a high MSS for PTPN11, a known LYN antagonist in breast cancer metastasis (Coopman et al., 2000).

### Table 1. Motifs identified by each algorithm in each dataset

<table>
<thead>
<tr>
<th>Dataset</th>
<th>MoDL Motif set</th>
<th>Motif-X score</th>
<th>Compression (bits)</th>
<th>Motif-X Motif set</th>
<th>Compression (bits)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HER2 P. EGF-stimulated 229 peptides</td>
<td>[DE]..Y</td>
<td>31.65</td>
<td>122.92</td>
<td>D..Y</td>
<td>8.32</td>
</tr>
<tr>
<td>HER2 P. HRG-stimulated 191 peptides</td>
<td>[DE]..Y</td>
<td>30.18</td>
<td>100.80</td>
<td>E..Y</td>
<td>7.47</td>
</tr>
<tr>
<td>HER2 24H. EGF-stimulated 225 peptides</td>
<td>[ADEN][ADLP]..Y</td>
<td>47.75</td>
<td>140.17</td>
<td>D..Y</td>
<td>8.83</td>
</tr>
<tr>
<td>HER2 24H. HRG-stimulated 209 peptides</td>
<td>[DENS][DNPRS]..Y</td>
<td>43.65</td>
<td>119.09</td>
<td>D..Y</td>
<td>8.22</td>
</tr>
<tr>
<td>Mast cell 142 peptides</td>
<td>[DE]..[ADESTY]</td>
<td>64.75</td>
<td>137.69</td>
<td>E..Y</td>
<td>8.87</td>
</tr>
</tbody>
</table>

Motifs returned by MoDL are ranked by the increase in description length when the motif is removed from the motif set. The motifs returned by Motif-X are ranked by Motif-X score. The Motif-X score is the negative logarithm of the binomial p-value computed using the percentage of phosphopeptides with the motif and the percentage of background peptides with the motif. The compression for a motif set $\mathcal{M}$ is the difference in description length $\Lambda(X;\{\}) - \Lambda(X;\mathcal{M})$. See Supplementary Table S3 for a full table of all datasets.
Table 2. MSSs for the motifs discovered in the HER2 datasets (false discovery rate = 0.05)

<table>
<thead>
<tr>
<th>Dataset</th>
<th>MoDL</th>
<th>MEME</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Motif</td>
<td>Kinase/Phosphatase</td>
</tr>
<tr>
<td>HER2 P. EGF-stimulated</td>
<td>ABL</td>
<td>Y..[PV]</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HER2 24H HRG-stimulated</td>
<td>ABL</td>
<td>Y..P</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HER2 24H HRG-stimulated</td>
<td>PTPN11</td>
<td>[DE]..Y[ADESTY]</td>
</tr>
<tr>
<td></td>
<td>SYK</td>
<td>[DENS][DNPRS].Y</td>
</tr>
<tr>
<td></td>
<td>ZAP70</td>
<td>[DENS][DNPRS].Y</td>
</tr>
</tbody>
</table>

MSSs were computed for the proteins that appear in the STRING database. The motifs logos were created using WebLogo (Crooks et al., 2004). Motifs and interacting proteins for MEME are in Supplementary Table S5.

4 DISCUSSION

We introduced the MoDL algorithm for the discovery of protein phosphorylation motifs, and showed that MoDL outperforms other algorithms for identification of mixtures of motifs in phosphoproteomics datasets. In particular, MoDL outperforms Motif-X even when using Motif-X’s scoring function, and generally produces motifs with higher specificity. Unlike other motif-finding methods, the MoDL algorithm requires no user-defined parameters besides criteria for termination. We showed that MoDL more accurately identifies motifs in synthetic datasets compared with both pattern-based motif-finding algorithms, such as Motif-X, Teresias and MDL-Pratt, and profile-based algorithms, such as MEME. Note that our comparisons to other algorithms like MEME are not indictments of these methods, but more a reflection of the fact that many motif-finding algorithms are not optimized for the problem of separating a set of sequences into a mixture of instances of an unknown number of motifs. For example, MEME is optimized for the problem of finding motifs in unaligned sequences where motifs are expected to be relatively rare compared with the sequence length.

We also compared the performance of MoDL and MEME on these datasets. On the HER2 P. EGF and HER2 24H HRG datasets MEME returned significant (false discovery rate \(\leq 0.05\)) motifs for ABL. The MSS for ABL in these two datasets are 1.5453 and 1.4227, respectively, which are much lower than the corresponding MSS of 3.7628 and 3.4672 obtained by MoDL (Table 2 and Supplementary Table S5).

Many of the motifs identified by MoDL are short and do not have many conserved positions, both consistent with earlier studies (Schwartz and Gygi, 2005) and various motif databases (Amanchy et al., 2007; Balla et al., 2006; Obenauer et al., 2003). Our identification of the linker sequence in the zinc-finger proteins shows that longer and/or highly conserved motifs are identified by MoDL when they are present. The relatively small number of conserved positions in many of the phosphorylation motifs imply that de novo prediction of phosphorylation sites and/or substrates of a particular kinase/phosphatase from sequence motifs in proteins sequence alone will likely yield many false positives.

We introduced the MSS to evaluate whether a kinase or phosphatase preferentially interacts with proteins containing a given motif. While, the MSS is an imperfect measure of kinase/phosphatase enrichment because it depends on the quality of the underlying protein–protein interaction network, we were able to identify several kinase/phosphatase-substrate interactions in the HER2 datasets that are known to be active in this signaling pathway. We also obtain novel predictions of interactions that were not recorded in the STRING database including the phosphorylation of GRF1 by ABL. In contrast to the NetworKIN algorithm (Linding et al., 2007) – which links kinases to their substrates using STRING and well-characterized motifs in Scansite (Obenauer et al., 2003) – our approach focuses on using protein–protein interactions to validate newly discovered motifs identified in a proteomics experiment. We also used the MSS to directly compare motifs returned by MoDL and motifs returned by MEME on experimental data, suggesting further uses of this statistic for comparing motif-finding programs.
Discovery of phosphorylation motif mixtures

An important caveat to discovery of phosphorylation motifs in mass spectrometry datasets is that sample preparation steps used to enrich for phosphopeptides might introduce biases in the phosphorylation sites that are measured. The three datasets we reported used immobilized metal affinity chromatography (IMAC) and immunoaffinity purification (IAP), while other affinity-based methods like metal oxide affinity chromatography (MOAC) are becoming popular (see Hoffert and Knepper, 2007, for a review). A recent comparative study (Bodenmiller et al., 2007) found that ∼35% of the phosphopeptides were common to all three techniques, suggesting that the biases of each technique are not overwhelming.

Framing the motif-finding problem as one of minimizing description length is a promising approach, and further improvements to the MoDL algorithm are possible. In particular, we observed that MoDL returns at most three motifs in phosphopeptide data, and thus it is possible that MDL might be too restrictive of a measure in that two or more biologically distinguished motifs might be merged into a single motif. Notably, the motif [DE]_X[ADESTY] in the mast cell dataset includes several distinct single-residue motifs for the SRC kinase family as listed in Schwartz and Gygi (2005). Thus, it might be useful to introduce a user-defined parameter of the number of motifs to return, allowing the user to incorporate prior knowledge about the number of interactions expected in a dataset. MoDL can also be modified to incorporate user-defined variable-length gaps as MDL-Pratt provides (Brazma et al., 1996). Finally, incorporating the MSS in the motif discovery stage will explicitly identify motifs with high MSSs in the protein–protein interaction network.

We demonstrate that the combination of sequence motifs identified in phosphoproteomics data from a single experimental condition with high-throughput protein interaction networks is a promising approach for linking kinases/phosphatases to their substrates in an experimentally stimulated signaling pathway. **Funding:** Career Award at the Scientific Interface from the Burroughs Wellcome Fund (to B.J.R.); National Institutes of Health (grant 2P20RR15578 to A.R.S.); Beckman Young Investigator Award (to A.R.S.); NSF Graduate Fellowship (to A.R.).

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