A peer-reviewed version of this preprint was published in PeerJ on 8 January 2015.

View the peer-reviewed version (peerj.com/articles/724), which is the preferred citable publication unless you specifically need to cite this preprint.

Singh GP. (2015) Association between intrinsic disorder and serine/threonine phosphorylation in *Mycobacterium tuberculosis*. PeerJ 3:e724
https://doi.org/10.7717/peerj.724
Association between intrinsic disorder and serine/threonine phosphorylation in *Mycobacterium tuberculosis*

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Abstract

Serine/threonine phosphorylation is an important mechanism to regulate protein function. In eukaryotes phosphorylation occurs predominantly in intrinsically disordered regions of proteins. While serine/threonine phosphorylation and protein disorder are much less prevalent in prokaryotes, *M. tuberculosis* has both high serine/threonine phosphorylation and disorder. Here I show that, similar to eukaryotes, serine/threonine phosphorylation sites in *M. tuberculosis* are highly enriched in intrinsically disordered regions, indicating similarity in substrate recognition mechanism of eukaryotic and *M. tuberculosis* kinases. Serine/threonine phosphorylation has been linked to the pathogenicity and survival of *M. tuberculosis*, thus better understanding of how its kinases recognize their substrates could have important implications in understanding and controlling the biology of this deadly pathogen.
Introduction

Reversible phosphorylation of serine and threonine residues is a widespread post-translational modification in eukaryotes, with more than a third of proteins phosphorylated during their lifetime [1]. Phosphorylation can modify protein interactions, enzyme function, localization or degradation. While traditionally, regulation and signal transduction in bacteria was thought to be mediated by histidine and aspartate phosphorylation in two-component systems, recently the occurrence and importance of phosphorylation of serine/threonine (S/T) residues has gained much attention [2–4]. Large scale mass-spectrometry based analysis has revealed S/T phosphorylation in a number of bacteria [2]. *M. tuberculosis* shows the highest rates of phosphorylation among studied bacteria with 8% of the proteins identified as phosphorylated [5].

One of the most interesting findings about serine/threonine phosphorylation in eukaryotes is its association with intrinsically disordered regions [6–11]. Intrinsically disordered regions lack a well-defined three-dimensional structure and are characterized by low content of hydrophobic amino acids and high net charge [12]. These characteristic physiochemical properties allow accurate prediction of disordered regions across proteomes [13]. Disordered regions are often associated with binding to multiple partners in a transient manner [14–19]. These regions may undergo disorder to order transition upon binding, with decrease in conformational entropy. This uncouples binding affinity and specificity, allowing highly specific interactions to be reversible [20,21]. It has been proposed that due to high surface accessibility and transient mode of interactions, disordered regions are ideally suited for regulation by reversible phosphorylation.
Disordered regions are highly abundant in eukaryotes with about a third of the proteins predicted to have at least one long (> 30 residues) disordered region and about 19% of the residues in disordered state [22]. In contrast, most bacteria have much less disorder in their proteome, with about 4% proteins with long disordered regions and about 6% disordered residues [22]. The association between disorder and serine/threonine phosphorylation has not been investigated in prokaryotes.

Among bacteria, *M. tuberculosis* has one of the highest disorder content, with about 10% disordered residues [22]. Thus, *M. tuberculosis* has both high level of S/T phosphorylation and disorder content. This prompted me to ask whether the association seen between S/T phosphorylation and disorder in eukaryotes might be present in *M. tuberculosis*, which indeed was found to be the case.

**Materials and Methods**

Data on *M. tuberculosis* phosphoproteins and phosphosites was obtained from Prisic *et al.* [5]. This study identified 301 phosphoproteins and 500 S/T phosphorylation events. For 215 of these sites, the specific residue that was modified could be identified with high confidence. *M. tuberculosis* proteome was obtained from Tuberculist database [23]. For disorder prediction I utilized IUPred program [24]. IUPred method is based on the observation that disordered regions do not form sufficient favorable interactions to fold, and thus have high estimated energy content [25]. I also utilized ESpript program [26], which is conceptually different from IUPred. This method is machine learning based predictor which was trained on experimentally characterized disordered regions (missing regions in X-ray structures in PDB). I also used MFDp2 disorder
prediction tool, which is an ensemble disorder prediction tool [27,28]. Secondary structure
prediction was done at Network Protein Sequence Analysis (NPSA) server [29] using consensus
approach [30]. To calculate statistical significance of enrichment of pS/T sites in disordered and
loop regions, I also used randomization test. For example, 84 pS/T sites are observed as
disordered by IUPred program out of total 215 pS/T. For testing enrichment, I randomly selected
equal number of S/T sites from each phosphoprotein as observed (1000 randomizations), and
counted number of randomizations in which 84 or more sites were observed in disordered
regions. This was not observed in any of 1000 randomization; hence $p$-value is less than 1e-3.
To analyze conservation of S/T sites, seven mycobacterium genus were chosen which are least
similar to $M. tuberculosis$ with respect to BLAST sequence similarity score ($Mycobacterium sp.
MCS, M. vanbaalenii, M. smegmatis, M. chubuense, Mycobacterium sp. JDM601, M. gilvum and
M. abscessus$). Orthologs of $M. tuberculosis$ in mycobacteria were identified using reciprocal
best blast approach [31], and aligned using Clustal Omega [32]. Alignment positions with gaps
were excluded from the analyses. Out of 215 sites, 135 sites were present in proteins with
orthologs in all 7 species. Of these 135, 103 sites were without gaps using Clustal Omega.
Positions with replacement of serine with threonine and vice-versa were considered as
conserved. Phosphosite data for other bacteria was obtained from respective publications [33–
40].

**Results**

Mass-spectrometry based analysis had previously revealed 301 phosphoproteins in $M.$
tuberculosis with 500 S/T phosphorylation sites [5]. First, I tested whether phosphoproteins in
$M. tuberculosis$ are more likely to be disordered (i.e. have long (>=30 residues) disordered
regions). I utilized IUPred program to predict disordered regions at the proteome wide level [25]. Phosphoproteins are about twice as likely to be disordered compared to non-phosphoproteins (29.6 % vs. 13.4% respectively, Fisher $p$ 4e-12). Since longer proteins are also more likely to have long disordered regions, I tested whether phosphoproteins have higher percentage of disordered residues. Phosphoproteins also have higher percentage of disordered residues compared to non-phosphoproteins (16.7% vs. 12.0% respectively, two tailed t-test $p$ 3e-5).

Out of 500 phosphorylation events detected in *M. tuberculosis*, the phosphoresidue could be identified for 215 sites with high confidence [5]. For these sites, I tested whether phosphorylated S/T (pS/T) residues are more likely to be disordered compared to non-phosphorylated S/T (npS/T) residues from the same set of proteins. Overall 39.1% of pS/T sites are disordered compared to 22.4% of npS/T sites (Fisher test $p$ 6e-8 and randomization test $p < 1e-3$, Figure 1). The results are very similar when another disorder prediction method, Espiritz[26] was used (52.6% pS/T disordered compared to 27.8% npS/T sites, Fisher test $p$ 8e-14, randomization test $p <1e-3$, Figure 1). A more recent disorder predictor MFDp2 [27,28] also gives similar results (43.7% pS/T disordered compared to 19.9 % npS/T sites, Fisher test $p$ 6e-15, randomization test $p <1e-3$, Figure 1). Disordered regions are also characterized by high irregular secondary structure (coil regions). Thus, I tested whether pS/T residues are enriched in coil regions of proteins. pS/T residues are more likely to occur in predicted coil regions compared to npS/T residues (70.2% pS/T sites in coil compared to 55.7% npS/T sites in coil, Fisher test $p$ 2e-5 and randomization test $p <1e-3$, Figure 2). There is also depletion of pS/T residues in beta-sheet regions (4.2% pS/T sites in sheet compared to 11.3% npS/T sites in sheet, Fisher test $p$ 4e-4 and randomization test $p <1e-3$, Figure 2), while no significant difference was found for helix regions.
(22.8% pS/T sites in helix compared to 27.1% npS/T sites in helix, Fisher test $p$ 0.2 and randomization test $p$ 0.12, Figure 2).

Disordered regions evolve faster than ordered regions [41], but pS/T residues evolve slower than npS/T in eukaryotes [42,43], so I tested whether *M. tuberculosis* pS/T and npS/T differ in conservation among seven mycobacterial proteomes (see methods). Conservation of pS/T is not significantly different from npS/T residues among these mycobacterial proteins (5.0 species vs. 5.2 species respectively using, Wilcox test $p$ 0.92, number pS/T sites 103). Separately analyzing disordered and ordered sites does not reveal significant difference in conservation between pS/T and npS/T sites (Wilcox $p$ 0.85 and 0.87 respectively). Thus pS/T are not more conserved than npS/T among mycobacteria.

Prisci *et al.* [5] conducted *in vitro* phosphorylation of 13-mer synthetic peptides corresponding to *in vivo* phosphorylation sites with different purified kinases. They could find phosphorylation of about half of these peptides. I tested the whether different kinases have differential preference for predicted disordered phosphoacceptors. I find that PknA has slightly higher preference for disordered phosphoacceptors than other kinases (Figure 3); however the uncorrected Chi-square test $p$-value is 0.04. Other kinases show uncorrected $p$-values > 0.05.

**Discussion**

Here I show the enrichment of localized serine/threonine phosphosites in disordered regions of proteins in *M. tuberculosis*. This preference is similar to that observed in eukaryotes. Association
between protein disorder and phosphorylation may offer similar advantages as those proposed in eukaryotes, including binding to multiple partners and transient mode of interaction [6,14–21], which are a prerequisite for regulatory interactions. Thus *M. tuberculosis* and eukaryotic serine/threonine phosphorylation dependent regulation may be more similar than generally appreciated. While most bacteria have low amount of protein disorder, *M. tuberculosis* has high disorder content. I suggest that high disorder content might be the reason for association between disorder and phosphorylation in *M. tuberculosis*. This suggestion is supported by the observation that other bacteria with high disorder content (*Streptomyces coelicolor* and *Halobacterium salinarum*) also show association between disorder and S/T phosphorylation, but bacteria with low disorder content do not (Supplemental Table 1) except for *Thermus thermophilus*, where enrichment of phosphorylation in disordered regions with marginal statistical significance was observed (Supplemental Table 1). Previously, a large fraction of phosphosites in *Thermus thermophilus* were observed in loop regions [44], albeit the statistical test for enrichment was not performed.

I find that PknA has a slightly higher preference for disordered phosphoacceptor sites on synthetic 13-mer substrate peptides under *in vitro* conditions than other kinases (Figure 3). However these results should be taken with the caveat that the structure of a peptide in *in vitro* conditions might be very different from the *in vivo* structure in the context of the full protein. Thus differential preferences of different kinases towards disordered substrates under *in vivo* conditions remain an open question.
I did not find pS/T sites to be more conserved than npS/T sites among mycobacteria, this suggests that phosphorylation network in mycobacteria evolves fast and may contribute towards niche specific adaptation. Important questions for future include, whether disordered and ordered S/T phosphosites are functionally different, and whether different kinases differ in their preference for disorder in their substrates under physiological conditions. Incorporation of disorder information might also be useful to predict novel S/T phosphosites [45], as has been shown in eukaryotes [6,46]. S/T kinases and their substrates have been linked to the survival, pathogenesis and virulence of *M. tuberculosis* [3,47,48], thus these finding may facilitate understanding the basic biology this deadly pathogen.

**Acknowledgements**

I would like to thank Dr. Shampa Ghosh from Bionivid Technology, Dr. Rahul Modak and Dr. Avinash Sonawane from KIIT University for critical reading of the manuscript.

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**Figure Legends**

**Figure 1.** Phosphorylated serine/threonine sites in *M. tuberculosis* are more likely to be **disordered**. Both phosphorylated and non-phosphorylated serine/threonine sites are from the same set of proteins. The disorder was predicted using IUPred, Espiritz and MFDp2 methods. The Fisher test \( p \) values are 6e-8, 8e-14 and 6e-15 respectively. pS/T - phosphorylated serine/threonine, npS/T- non-phosphorylated serine/threonine.

**Figure 2.** Association between predicted secondary structure and phosphorylation of serine/threonine sites in *M. tuberculosis*. Phosphorylated serine/threonine sites occur preferentially in coil regions, while being significantly depleted in sheet regions and show no significant difference in helix regions. The Fisher test \( p \) values are 2e-5, 4e-4 and 0.2 respectively. pS/T - phosphorylated serine/threonine, npS/T- non-phosphorylated serine/threonine.

**Figure 3.** Percentage of disordered phosphoacceptors are shown for different kinases. The phosphorylation was carried out by purified kinases under in vitro conditions on synthetic 13-mer peptides corresponding to in vivo phosphorylation sites (Prisic et al.). Uncorrected Chi-square \( p \) values were significant (<0.05) only for PknA.
Figure 1
Figure 2

A bar graph showing the percentage of serine/threonine sites in different structural states: Coil, Sheet, and Helix. The graph compares pS/T and npS/T states.
Figure 3