Disorder-to-order transition in PE–PPE proteins of Mycobacterium tuberculosis augments the pro-pathogen immune response

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Keywords
folding-upon-binding; immune modulation; immunogenicity; Mycobacterium tuberculosis; pathogenesis; protein–protein interaction

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(Received 3 June 2019, revised 11 October 2019, accepted 22 October 2019)
doi:10.1002/2211-5463.12749

A growing body of evidence supports the hypothesis that intrinsically disordered proteins often mediate host–pathogen interactions and modulate host functions for pathogen survival and virulence. Mycobacterium tuberculosis (M. tb) has evolved largely through reductive evolution, with a few exceptions such as the glycine–alanine-rich PE–PPE/PGRS protein family, which has been expanding in pathogenic mycobacteria. Here, our analyses of the M. tb proteome and secretome revealed that the PE–PGRS subfamily is enriched for disordered regions and disordered binding sites, pointing to their importance in host–pathogen interactions. As a case study, the secondary structure of PE35–PPE68 and PE32–PPE65 of the pathogenesis-related RD1 and RD8 regions was analyzed through Fourier-transform infrared spectroscopy. These disordered proteins displayed a considerable structural shift from disordered to ordered while engaged in the formation of complexes. While these proteins are immunogenic individually and enhance the pro-pathogen response, their corresponding complexes enhanced the responses manifold as displayed here by PE35 and PPE68. It is likely that M. tb exploits such disorder–order structural dynamics as a strategy to mount a pro-pathogen response and subvert host defense for productive infection. This functional gain also serves as a means to compensate genomic content loss due to reductive evolution.

Abbreviations
BCG, bacillus Calmette–Guérin; BSA, bovine serum albumin; E. coli, Escherichia coli; ELISA, enzyme-linked immunosorbent assay; ESAT-6, early secreted antigenic target; ESX, ESAT-6 secretory system; FTIR, Fourier-transform infrared spectroscopy; HIV, human immunodeficiency virus; IDR, intrinsically disordered protein; IDP, intrinsically disordered region; IgG, immunoglobulin G; LPS, lipopolysaccharide; Mce, mycobacterial cell entry protein; MmpL, mycobacterial membrane protein large; PE, proline–glutamic acid; PGRS, polymorphic glycine-rich sequence; PPE, proline–proline–glutamic acid; RD, region of difference; TLR2, Toll-like receptor 2.
Mycobacterium tuberculosis (M. tb), the etiological agent of tuberculosis (TB), causes millions of deaths worldwide. The emergence of drug resistance and co-infection with HIV has further worsened the situation. The molecular mechanism(s) of pathogenesis and the associated host–pathogen interactions are yet to be elucidated, thereby hampering the development of new drugs or vaccines to target this bacterial pathogen. Thus, there is an immediate need for novel approaches to tackle this pathogen, which has co-evolved with humans and thus possibly adapted well to counter host resistance [1].

Mycobacterium tuberculosis has undergone reductive evolution; that is, its genome size has decreased with gain in pathogenesis [2,3]. The only exception to this tendency is few protein families such as toxin anti-toxin system and novel PE–PPE/PGRS family of proteins that are specific to mycobacteria. This gene family has expanded with the gain in pathogenesis and accounts for about 10% of the coding capacity of the M. tb genome [4,5]. Some of the genes are organized in a defined operonic pattern within the genome, where a PE gene is followed by a PPE gene [6] with few genes associated with members of the ESX family that are important virulence factors and T-cell antigens. Many members of the PE–PPE/PGRS family are membrane attached and localize to the cell surface where they are involved in host–pathogen interactions. Some members of the family are involved in modulating the host immune response [7–12], immune quorum sensing, and virulence [13,14]. Comparative genome analysis of M. tb H37Rv and BCG, a non-pathogenic attenuated strain of Mycobacterium bovis, revealed the absence of some genomic regions in BCG denoted as regions of difference (RDs). The exclusive presence of some RDs in pathogenic strains suggests that their encoded proteins are important virulence factors. There are 16 RDs, of which RD1 has been well studied. RD1 is absent in all the BCG strains, but present in all virulent strains where it codes for the PE35, PPE68, and ESAT-6 genes along with other genes belonging to the ESX family that are important for attachment to and invasion of host cells.

Other M. tb protein families known to be associated with pathogenesis include the Mce (mammalian cell entry) and MmpL families (mycobacterial membrane protein large) [28]. While the Mce proteins play an important role in the entry and survival of the bacteria in the host [29], the MmpL proteins are multisubstrate efflux pumps. We therefore selected the PE–PPE, PE–PGRS, Mce, and MmpL families, along with the entire secretome of M. tb [19] for analysis of structural disorder and potential interaction sites. Computational analysis revealed that PE–PGRS family had highest disorder along with some other PE and PPE family proteins. To complement our computational study, we unsuccessfully tried to clone, express, and purify PE–PGRS proteins. Considering the higher disorder content in some PE–PPE proteins, we further experimentally investigated two interacting protein pairs from the PE–PPE family, namely PE32/PPE65 and PE35/PPE68. Besides being disordered, these protein pairs are present within the RD regions and thus are suitable candidates to study the folding-upon-binding mechanism to gain insight into structure-mediated pathogenicity. In our previous study, we demonstrated that the PE32–PPE65 protein complex is more immunogenic than its individual protein components [30]. We now describe a step-up
approach to determine the role of structural flexibility in the functions of such PE–PPE protein pairs. Our results suggest that the presence of intrinsically disordered regions and host-like interaction sites in M.tbc proteins [31] could be a key strategy employed by the pathogen for rapid adaptive changes, host invasion, hijacking of host machinery, and diversifying the functions of its proteins.

Materials and methods

Dataset assembly

The M.tbc H37Rv proteome was obtained from NCBI. The selected families were complemented by the PE10, PE11, PPE40, PE_PGRS63, and PE_PGRS58 sequences from Tuberculist database (http://tuberculist.epfl.ch/), and the secretome was obtained from Ref. [19] (Table S1).

Computational protein analysis

We predicted structural disorder for each member of the investigated families and reference set using the IUPred (long) algorithm [32]. Residues with a predicted disorder propensity > 0.5 were considered as disordered. PE–PPE family disorder analysis was also carried out using the RONN prediction tool. For detecting disordered binding regions, we used the ANCHOR method [33].

Expression and purification of proteins

The expression and purification of proteins were done according to previously published protocol [30,34]. Briefly, expression constructs of all four genes, PE32-pET28a, PPE65-pET28a, PE35-pET28a, and PPE68-pET28a, were transformed into Escherichia coli BL21(DE3) strain. Culture was induced with 1 mM IPTG for 3 h at 37 °C for the expression of recombinant proteins. Purification of recombinant proteins was carried out by sonicating bacterial cells and solubilizing recombinant proteins using 0.3% sarkosyl in phosphate-buffered saline. Recombinant proteins were purified using Ni-NTA affinity column and eluted with 200 mM imidazole. Eluted proteins were dialyzed for 24 h at 37 °C for the expression of recombinant proteins. Purification of recombinant proteins was carried out by sonicating bacterial cells and solubilizing recombinant proteins using 0.3% sarkosyl in phosphate-buffered saline. Recombinant proteins were purified using Ni-NTA affinity column and eluted with 200 mM imidazole. Eluted proteins were dialyzed for 24 h with buffer replaced at different time points with fresh (100 mM) phosphate-buffered saline. Dialyzed proteins were concentrated using Centricons with 3 kDa cutoff. Protein was treated with phosphate-buffered saline. Dialyzed proteins were concentrated using Ni-NTA affinity column and eluted with 200 mM imidazole-buffered saline. Recombinant proteins were purified utilizing recombinant proteins using 0.3% sarkosyl in phosphate-buffered saline. Recombinant proteins were purified using Ni-NTA affinity column and eluted with 200 mM imidazole. Eluted proteins were dialyzed for 24 h with buffer replaced at different time points with fresh (100 mM) phosphate-buffered saline. Dialyzed proteins were concentrated using Centricons with 3 kDa cutoff. Protein was treated with phosphate-buffered saline.

Fluorescence spectra measurements

Fluorescence emission spectra of all four recombinant proteins (0.5 mg·mL⁻¹) were carried out in a Jasco fluorometer (FP-6200) (Jasco International Co., LTD, Tokyo, Japan). The excitation wavelength was set at 280 nm, and emission spectra were measured in the wavelength range 300–340 nm in quartz cuvette of path length 3 mm. All the experiments were carried out at 25 ± 0.1 °C with both excitation emission slits set at 5 nm. The fluorescence intensity of the blank solution was subtracted from that of each sample.

FTIR spectroscopy

Protein samples of PE32, PE35, PPE65, and PPE68 were purified to high degree of purity (Fig. S1) and concentrated for successful FTIR experiments. To generate protein complex, both the proteins were mixed in equal amount in 1× PBS and incubated for 1 h at 4 °C. Proteins were stored at −20 °C for future use. Protein concentrations were estimated using BCA protein estimation kit. Phosphate-buffered saline (100 mM) was used as protein storage and experimental buffer for FTIR. Concentrated 3 µL samples of the proteins were placed on a SensIR ATR reflection element. FTIR spectra were recorded at 4 cm⁻¹ resolution on an Agilent Cary 630 spectrometer (Agilent Technologies, Inc., Santa Clara, CA, USA) equipped with a DTGS detector. The experiments were performed on the liquid sample at room temperature using the spectrometer software MICRO-LAB (Agilent Technologies, Inc., Santa Clara, CA, USA). For each sample, a 64-scan single-beam spectrum of the buffer was collected. For the protein sample, spectrum in the absorption mode (64 scans each) was recorded.

Secondary structure analysis was carried out by curve fitting using the opus software (Opus Consulting Solutions Inc, Alpharetta, GA, USA). The absorption bands at low wavenumber were free from features of noise as judged from the peaks above 1800 cm⁻¹ and water vapor at below 1750 cm⁻¹. A straight baseline passing through 1715 and 1600 cm⁻¹ was subtracted before the curve fitting. The baseline was again modified by the least squares curve-fitting program, which allows for a horizontal baseline to be adjusted as an additional parameter to obtain the best fit. The second-derivative spectrum was used to determine the initial peak positions for curve fitting, and the peaks were fitted using Voigt functions [35,36]. In curve fitting, each component is a mixture of Lorentzian and Gaussian line shapes. The area under the entire band was considered 100%, and each component after fitting was expressed as a percent fraction.

Immunization of mice

C57BL/6j mice (8–10 weeks age) were procured from National Institute of Nutrition, Hyderabad, India. The experimental protocol was approved by the Institute Animal Ethics Committee, and during all the procedures, Institute Animal Ethical Guidelines were followed. These animals
were housed in plastic cages with wire lid and provided with access to unlimited pellet diets and water. A photoperiod of 12 h was provided. Experiment was performed in two parts: In one part, two groups of mice (five mice in each group) were immunized with individual proteins, and the third was taken as control; and in the second part, two groups were taken with same number of mice, one immunized with PE–PPE complex and the other taken as control. Mice were immunized subcutaneously with purified proteins (20 µg of protein in PBS) without any adjuvant in LPS-free condition to minimize Th1/Th2 bias inherent to adjuvants. Purified proteins were passed through polymyxin B bead column to remove endotoxins. Mice injected with PBS served as controls. Injection was done subcutaneously at the base of the tail. A booster dose of respective proteins was given at the 15th day of primary immunization. Mice were sacrificed after 4 weeks of primary immunization.

Serum collection, splenocyte isolation, and single-cell preparation

Mice were euthanized, and blood was collected through cardiac puncture and transferred to a 1.5-mL vial for coagulation at room temperature for 1 h. Later on, serum was collected by centrifugation of coagulated blood at 1500 g. Spleens were isolated, crushed, and passed through a 40-µm cell strainer to get a single-cell suspension. Cells were pelleted down at 300 g for 10 min, washed with PBS, and resuspended in RBC lysis buffer. The process of resuspension of cells in RBC lysis buffer was repeated till there was no sign of RBC in the pellet. The cells were washed twice with PBS and once with incomplete DMEM and finally resuspended in DMEM containing 10% FBS.

Splenocyte proliferation assay

Cells were seeded in 96-well plates and treated with different concentrations of proteins (0.2, 1, and 2 µg·mL⁻¹) for 3 days at 37 °C and 5% CO₂. [³H]thymidine (0.5 µCi·mL⁻¹) was added to each well and incubated at 37 °C for 24 h. Cell harvester was utilized for harvesting the cells, and beta scintillation counter was used to measure cpm.

Cell culture and ELISA for cytokine levels and B-cell response

Splenocytes (0.1 × 10⁶ cells per well) were seeded in 96-well plates in DMEM containing 10% FBS. Protein treatments with different concentrations were given for 72 h, and supernatants were collected and stored at −20 °C. Secreted cytokine levels in culture supernatant were measured using PeproTech (Rocky Hill, NJ, USA) standard ELISA development kits for TNF-α, IL-6, IL-4, and IL-10 and BioLegend ELISA MAX™ (San Diego, CA, USA) standard kit for IL-1β. ELISA was carried out using the manufacturer’s protocol. Briefly, capture antibody was coated on 96-well ELISA plates in PBS (PeproTech) or bicarbonate coating buffer and kept at room temperature or 4 °C overnight. Plates were washed with PBS-T (0.05% Tween-20) thrice. Three hundred microlitre of 1% BSA solution was used as blocking buffer, while 0.1% BSA was used as assay diluent. After 2 hours of blocking, the plate was washed four times with 300 µL of wash buffer and samples along with standards were added and incubated at room temperature for 2 h. Plates were again washed four times with wash buffer, and detection antibody was added. Plates were washed four times, and avidin-conjugated antibody was added for 30 min at room temperature. After four washes, ABTS/TMB substrate was added for color development. 2 N H₂SO₄ was added to stop the reaction. For PeproTech kit, absorbance was measured at 405 nm, whereas for the BioLegend kit, absorbance was taken at 450 nm. B-cell response against individual proteins was assayed using antibody against total IgG as described earlier [37].

Intracellular cytokine staining and extracellular staining of surface markers

Splenocytes (1 × 10⁶) were seeded per well in 96-well plates and restimulated with 10 µg·mL⁻¹ proteins for 8 h in the presence of GolgiPlug™/GolgiStop™ (BD Biosciences, San Jose, CA, USA). Cells were harvested, washed, and stained for the CD4 and CD8 cell surface markers with anti-CD4 and anti-CD8 antibodies. Cells were fixed with 4% paraformaldehyde, permeabilized with 0.02% Triton X-100, and then stained for intracellular IFN-γ and IL-2 (FITC for CD4, Alexa Fluor 488 for CD8, APC for IFN-γ, and PE for IL-2) Antibodies were procured from BD Biosciences.

Statistical evaluation of immunological experimental results

PRISM 5 (GraphPad Software, San Diego, CA, USA) was used for statistical analysis, and the results were expressed as mean ± SD. One-way ANOVA statistical comparisons were performed on the results to obtain P values. The value of P < 0.05 was considered statistically significant.

Results

PE–PPE/PGRS proteins of the three subfamilies are disordered to different extents

Computational analysis by the IUPred and RONN disorder prediction methods revealed that the PE–PGRS subfamily is significantly more disordered than the other selected families or the rest of the proteome.
Composition bias is a common feature of intrinsically disordered proteins. These disordered proteins or disordered regions in proteins are usually enriched with charged amino acids such as Arg, Gln, Glu, Lys, Pro, and Ser and deficient in hydrophobic amino acids such as in Phe, Trp, Tyr, Cys, Ile, Leu, and Val. However, amino acids such as Ala, Gly, His, Asn, and Asp are known to be enriched in some intrinsically disordered proteins. Enrichment in disorder was also observed for secreted proteins that potentially modulate host cell processes through regulatory and signaling interactions. The same two groups of proteins were also enriched in ANCHOR-predicted disordered binding sites (Fig. 1B), which are disordered patches likely mediating energetically favorable interactions with globular protein partners. While PE proteins were rather depleted in disordered regions, PPE proteins had similar disorder levels as the rest of the proteome. PE domain of PE proteins is around 100 residues in

![Image](74x383 to 538x562)

**Fig. 1.** Computational analysis of *Mycobacterium tuberculosis* protein families and selected groups of proteins. We compared (A) the IUPred-predicted protein disorder content and (B) density of ANCHOR binding sites of the selected six groups of proteins with the rest of the proteome. The number of proteins in each group is indicated above the respective box in brackets. Boxes contain 25–75% of the data with the median indicated as a horizontal line in the box. Lower and upper whiskers indicate 10% and 90% of the data, respectively. Significance was calculated by the Kruskal–Wallis test coupled with Dunn’s multiple comparisons post hoc tests for each investigated property, and significant differences are marked by stars (*P < 0.05; **P < 0.01, ***P < 0.001, ****P < 0.0001). (C) Domain maps of the shortlisted proteins for experimental study: PE32, PPE65, PE35, and PPE68 are depicted in light gray with Pfam entities shown as darker gray bars with the name and residue boundaries indicated. ExPASy-PROSITE predicted ala-rich low-complexity regions marked in green.
length, which attains an almost completely helical conformation in PE–PPE complexes [see Protein Data Bank (PDB) entry 4KXR]. As most of them probably show high helix propensities in their free forms, their low predicted disorder content is not surprising. PPE proteins are more variable in terms of their length and sequence. They all have an N-terminal PPE domain that can form a stable complex with a specific PE partner domain. As a complex, they mediate interactions with components of the type VII secretion system [38]. Besides their PPE domains, they have variable C-terminal regions that can be short (as in PPE41), or might contain transmembrane regions (as in PPE37), other domains, or disordered regions (as in PPE37 and PPE68). Although the PPE family does not have a very high level of disorder on average, some of its members, like the ones investigated in this study, do have extended disordered regions that could play a major role in pathogenesis. In case of the MmpL proteins that are transmembrane efflux pumps and Mce membrane proteins, the low predicted disorder content and lack of ANCHOR binding sites are not surprising, as transmembrane regions generally receive very low disorder probability values.

**Computational analysis of the PE32, PPE65, PE35, and PPE68 proteins**

Among the four proteins studied, PE32 and PPE65 are more structured according to IUPred (Fig. S3A). However, RONN predicted 15.1% and 54.96% structural disorder in these proteins, respectively (Fig. S3B). PE35 had little disorder, while PPE68 had a highly disordered C-terminal region according to both IUPred and RONN, also interspersed with a number of ANCHOR binding sites (Fig. S3C,D). The interaction partners of the four proteins were checked in the IntAct [39] and STRING [40] databases. PE32 and PPE65 interact with each other, based on their genomic locations, and also with the EsxW and EsxI proteins. Besides interacting with each other and ESX-1 specialized secretion system, proteins EssA, B, H and Snn2, PE35, and PPE68 also bind the human TLR2 receptor [34]. The proteins also have alanine-rich low-complexity regions and domains as shown in Fig. 1C and have a number of predicted high-probability post-translational modification sites, mostly N-myristoylation and phosphorylation sites, the biological relevance of which is questionable. We also aligned the PE proteins, PE32 and PE35, to PE25 for which a structure is available in the PDB in complex with PPE41. The three proteins are identical in length but show remarkable sequence variation (Fig. S4A). The alignment of PPE65 and PPE68 to PPE41 shows that both PPE65 and PPE68 have long C-terminal extensions compared to PPE41 that mainly consists of the PPE domain, and that these extensions do not show any detectable sequence similarity (Fig. S4B). The complete lack of sequence similarity outside the PPE domains of these proteins implies high rates of evolutionary changes and a high level of functional specialization within this subfamily.

**Protein–protein interaction studies using fluorescence spectroscopy**

PE–PPE genes whose products form a functional protein complex are generally co-operand and thus co-expressed [41]. So far, PE25–PPE41 and PE8–PPE15 pairs have been copurified and cocrystallized [17,18] probably due to the fact that PPE41 mainly consists of the PPE domain and lacks extended C-terminal regions. We investigated the structures and interactions of two other pairs: PE32 and PPE65, as well as PE35 and PPE68. The proteins were expressed in E. coli, and recombinant proteins were purified as per earlier method [30]. The purity of the proteins was checked through SDS/PAGE: Single bands were observed at their respective positions compared to the protein ladder (Fig. S1). As PE32 and PE35 proteins lack tryptophan residues, we used fluorescence spectroscopy to analyze the interaction of PE32 with PPE65 and PE35 with PPE68 (Fig. 2). A clear shift in the emission spectrum of PPE65 was observed when different concentrations of PE32 were added (Fig. 2A). The emission spectrum of PPE68 was similarly monitored while adding PE35 protein, and a pronounced change in fluorescence intensity was observed (Fig. 2B).

**Secondary structure analysis of PE32, PPE65, and the PE32–PPE65 complex**

Infrared bands in amide I region (1700–1600 cm⁻¹) arise mainly from the C=O stretching vibrations of the protein backbone. Different types of secondary structures give different C=O stretching vibrations in amide I region due to variations in molecular geometry and hydrogen bond patterns [42]. For each protein sample, the spectrum was obtained against the buffer background. The protein absorbance spectrum was further checked with the buffer spectrum and subtracted in an iterative manner until a straight baseline was obtained in the 2000–1750 cm⁻¹ spectral region. To resolve the extensively overlapping component bands in amide I region that arise from various secondary structural elements, second-derivative analysis with 5-point
smoothing was applied. The data obtained from second-derivative analysis were used to determine the number of bands (and their position) to resolve the protein spectrum into different secondary structure components. The band area for each component peak was used to calculate the relative contribution of that component to a particular protein’s secondary structure.

Using curve-fitting analysis on recombinant PPE65 (Fig. 3B), more than 66% of random coil and unstructured region content was observed, implying that the protein is loosely folded. The band observed below 1630 cm$^{-1}$ (1630–1615 cm$^{-1}$) indicated unstructured regions or protein aggregation. The bands near 1643 cm$^{-1}$ (1650–1640 cm$^{-1}$) and 1675 cm$^{-1}$ (1680–1662 cm$^{-1}$) arise from random coil conformations. The band near 1653 cm$^{-1}$ (1660–1650 cm$^{-1}$) signals alpha helical structure, while the band observed above 1680 cm$^{-1}$ (1700–1680 cm$^{-1}$) is assigned to beta sheets or beta turns. We also observed a fair amount of unstructured regions in PE32 (Fig. 3A); however, this protein was more structured in comparison with PPE65. When we mixed PE32 and PPE65 (in 1 : 1 ratio) to see the structural effects of the interaction, more than 66% ordered structure (36% alpha helices and 32% beta sheets) was observed. Thus, a structural shift from disorder to order occurred upon complex formation. However, significant amount of unstructured regions (> 30%) was still retained in the complex (Fig. 3C,D and Table S2), likely involving the C-terminal half of PPE65. The band observed near 1709 cm$^{-1}$ is tentatively assigned to Asp, Glu, or Asn, whereas the band at 1615 cm$^{-1}$ is assigned to Asn. This indicates that these amino acid side chains play a key role in complex formation. These results also potentially explain why previous expression and purification attempts for most members of this family in
isolation could be problematic, potentially precluding structural and biophysical studies.

**Secondary structure analysis of PE35, PPE68, and the PE35-PPE68 complex**

The other co-operonic protein pair, PE35 and PPE68, which were previously shown to interact with each other and the human TLR2 receptor, and to display increased immunogenic activity as a complex, was also used for FTIR analysis. PE35 displayed a highly disordered structure with 41% of regular secondary structures (Fig. 4A), to which alpha helices and beta sheets contributed 23% and 18%, respectively. The much longer PPE68 had a similar content of secondary structure (43%) as PE35, while in agreement with the applied disorder prediction methods, a significant amount of unstructured regions was also observed (Fig. 4B). Analysis of the structure content of the PE35-PPE68 complex revealed a major transition from disorder to order upon complex formation (Fig. 4C,D and Table S2), but with significant regions (40%) still retaining their disordered states in the complex, likely involving the C-terminal disordered half of PPE68. The peak around 1714 cm\(^{-1}\) indicates the involvement of charged residues such as Asp/Glu/Asn/Gln in the interaction between the two proteins. In all, comparing the spectra of the individual proteins and the corresponding complexes showed major structural shifts toward a more structured state, but also indicated the presence of disordered regions in the complex state for both protein pairs.

**PE35 and PPE68 induce cell proliferation individually but suppress polyfunctional T cells**

To determine the functional significance of the biophysical features reported above, experiments were designed, based on animal immunization studies, to determine the immunogenicity of the individual proteins. Different concentrations of PE35 and PPE68 proteins (0.2, 1, and 2 \(\mu\)g-mL\(^{-1}\)) were used to restimulate cultured splenocytes for 72 h which were then also treated with \([H^3]\) thymidine for 24 h. Harvested cells were used to count \(\beta\) activity using beta scintillation counter as a measure of cell proliferation. We found that proliferation in PPE68 restimulated cells was very high, while when applying a lower dose of PE35, it was not significant, implying that PE35 may be less antigenic. Clearly, individual PE35 and PPE68 proteins induced cell proliferation in splenocytes to different extents (Fig. S5). An earlier study with the PE35/PPE68 pair in macrophage cells has shown that these proteins suppress IL-12 secretion but enhance IL-10 secretion in a dose-dependent manner [27]. We restimulated splenocytes with the respective individual proteins in culture and determined the intracellular levels of cytokines (IL-2 and IFN-\(\gamma\)) through FACS. We observed a clear decrease in the number of IFN-\(\gamma^+\)/IL-2+ CD4\(^+\) cells. However, the number of double-positive CD8\(^+\) cells did not change significantly (Fig. 5A,B). These results from animal immunization experiments point to the low immunogenicity of individual proteins as opposed to the complex.
The PE35–PPE68 complex displays enhanced downregulation of Th1 cytokines

After investigating how the individual proteins modulate immune response, we investigated how the protein complex behaves. Interestingly, we observed a significant decrease in IFN-γ+/IL-2+ double-positive CD4+ and CD8+ cells when immunized with the complex of PE35–PPE68 (Fig. 6A,B). ELISA performed with the supernatant of restimulated splenocytes showed an increase in the level of IL-4 and IL-10 cytokines in a protein complex concentration-dependent manner (Fig. 7A,B) with a significant decrease in the level of IL-6 (Fig. 7C). These results demonstrate that the PE35–PPE68 complex shifts the balance toward Th2-dominated immune response, favoring the survival of the pathogen inside the host.

B-cell response against PE35–PPE68 pair in mice

Earlier observations with a PE–PPE pair showed that the PE protein was less immunogenic [41]. We predicted B-cell antigenicity using the DNASTAR tool [43], which predicts PE35 to be less immunogenic, but PPE68 to be highly immunogenic with several antigenic patches (Fig. S6). Evaluation of the total IgG response against the respective proteins revealed no significant response against PE35 alone, significantly higher response against PPE68 alone, and an even higher response for the PE35–PPE68 complex (Fig. 7D).

Discussion

Being an intracellular pathogen, M. tb potentially employs a repertoire of pathogenic effector proteins which can modulate host proteins to aid host cell entry and favor its survival inside macrophages. Given the fact that pathogenic M. tb strains have undergone reductive evolution [44], and based on our earlier observations [37,41], we hypothesized that they might employ disordered proteins or disordered protein regions to ensure that their relatively small proteomes still have a sufficient level of functional diversity and potential to undergo adaptive changes for maintaining survival and pathogenesis. These disordered proteins utilize their inherent plasticity to adopt entirely different structure (and subsequently, function) upon binding to different partners, thus increasing the interactome without requiring multiple proteins. Pathogens are known to employ disordered proteins to disrupt and subvert the host cell cycle and normal
cellular signaling processes [45]. Emerging evidences have validated that the flexibility of proteins in other phyla also adds to their functionality particularly in context with multiple interactions. We have recently published the role of unstructured C terminus of mycobacterial PPE37 in host–pathogen interaction [31]. A recent report also validates that secreted proteins, which are harbingers for interaction with host, from pathogens are also enriched in disordered amino acid stretches [46]. Moreover, even in eukaryotic systems, the disorder in a protein is now considered a central characteristic for binding promiscuity to multiple ligands [47,48].

Computational analyses revealed that the PE–PGRS subfamily is the most disordered among all the investigated pathogenesis-related protein families and that the secreted proteins of the bacterium are also more disordered than the rest of its proteome. This relatively higher level of structural disorder in PE–PPE/PGRS proteins supports their anticipated functions in immune evasion and host–pathogen interactions, since many known pathogenic effector proteins are disordered. Disorder-mediated aggregation propensity and susceptibility to proteolytic degradation make them very difficult to express and purify in recombinant form in isolation, leading to only the PE25–PPE41 and PE8–PPE15 pairs being successfully crystallized [17,18]. Thus, expansion of PE/PPE family in pathogenic mycobacteria seems justified as these proteins have accumulated multiple structural features that aid pathogenicity. The structural disorder is expected to enable these proteins as key signaling hubs that could modulate protein machinery of pathogen as well as host for its effective survival.
Biophysical studies revealed that PE35/PPE68 and PE32/PPE65 have unstructured regions to different extents and undergo disorder-to-order transition upon interaction. The observed infrared absorption bands from mainly charged amino acid side chains imply that they play a fundamental role in stabilizing the protein complex. The interaction of PE32 with PPE65 and PE35 with PPE68 also validates the biological relevance of the co-operative arrangement of their genes.

Our computational and experimental analyses suggest that while PE proteins show a considerable sequence variation, their short length and structural features seem relatively well-conserved. We found that in their unbound forms, PE proteins are disordered to different extents, but based on the PE25–PPE41 complex structure and our observations, they appear to fold upon binding to their cognate PPE partner domains almost completely. On the other hand, PPE proteins are longer and only their N-terminal PPE domains seem to take part in forming the complex with PEs and in interacting with Esx secretion system components [38]. While the observed increase in secondary structure content upon interaction reaffirms that PE and PPE domains cofold and get stabilized in PE–PPE complexes, we have measured a relatively large unstructured content in both investigated complexes, implying that the C-terminal variable halves of the PPE65 and PPE68 proteins retain their disordered states. This suggests the likelihood of multiple additional interactions that need to be understood to gain a holistic view of the mechanism of pathogenesis.
This transition toward structural order upon interaction of PE and PPE suggests gain of function. Investigating the immunogenicity of the proteins demonstrated that PE35 and PPE68 are antigenic, with PPE68 emerging as a strong B-cell antigen. PE35 proved to be a weak B-cell antigen, in agreement with earlier observations regarding the PE counterparts of other PE–PPE partners and with the PE domain of PE–PGRS33 [49]. PE/PPE proteins help in the formation of the complex required for transport of the proteins across the membrane [18,50]. Our results, however, showed that PE35/PPE68 plays an important role in immune modulation, which was earlier also reported for PE32/PPE65 [30]. The complex of PE35–PPE68 displayed gain of function by exhibiting significantly higher antibody response in mice, along with significantly enhanced suppression of IFN-γ+/IL-2+–double-positive CD4+ and CD8+ cells. A significant increase in the level of IL-4 and IL-10 cytokines along with a significant decrease in the level of IL-6 in the supernatant of restimulated splenocytes was evident in a protein dose-dependent manner when using the complex of PE/PPE for immunization as well as for in vitro stimulation. These observations indicate that PE–PPE complexes modulate immune response toward Th2 type, leading to increased anti-inflammatory and reduced pro-inflammatory cytokine production that eventually aids the escape/survival of the bacterium within the host. Although individual proteins were also immunogenic and displayed anti-inflammatory behavior, the stable PE–PPE complex had a greater impact. Therefore, the observed structural transition expectedly led to the observed gain of function that could be one of the contributing factors for enhanced pathogenicity (Fig. 8). Earlier observations on PE/PPE also depicted additive immunological effect as a pair [34,37], and we now provide a biophysical perspective of the observed effect. The stability accorded to the complex enables modulation in interaction with host proteins to enhance virulence. This could have profound implications in vaccine-based and other pathogenesis-related studies as disorder-to-order content in a protein is likely to affect the possible outcome of the interaction.

In summary, infection and intracellular survival strategies of M.tb may partly rely on disordered proteins, as its pathogenesis-related PE–PPE/PGRS proteins are enriched in structurally disordered regions. The repetitive and disordered nature of PE–PPE/
PGRS proteins likely enables fast evolutionary changes and thus may mask the pathogen from host immune response, similar to what has been described for merozoite surface protein 2 (MSP2) of the malaria pathogen *Plasmodium falciparum* [51]. Our results suggest that *M. tb* is able to counterbalance reductive evolution by employing this specific family of proteins, many of which contain disordered regions. These disordered regions achieve a more ordered state upon PE-PPE interaction or likely retain partially disordered states even after the formation of the specific complex. Being exposed on the pathogen surface and showing high levels of functional specialization among family members through variable segments, PE–PPE protein complexes successfully influence the host immune system toward favoring pathogen survival. Furthermore, the presence of multiple pairs/variants of such complexes [38] in the pathogenic toolbox of *M. tb* likely allows for using them in alternative combinations on its cell surface, thereby further hampering effective host immune response. Future studies should aim at a better understanding of the role of disordered proteins in *M. tb*, which will, in turn, help in deciphering the molecular mechanisms of the most prevalent infectious disease, and in developing improved strategies for its effective control.

**Acknowledgements**

JA thanks University Grants Commission for Senior Research Fellowship and EMBO for EMBO Short-Term Fellowship. MK thanks Council of Scientific and Industrial Research (CSIR) for Senior Research Fellowship. RP was supported by the EMBO fellowship (ALTF 702-2015). The Medical Research Council (MC_U105185859) and the Lister Institute Research Prize Fellowship supported MMB. SEH and NZE thank the Department of Biotechnology, Ministry of Science and Technology, Government of India (MoS&T, GoI), for a Center of Excellence grant. SEH is a JC Bose National Fellow of the Department of Science and Technology, Ministry of Science and Technology, Government of India (MoS&T, GoI), and Robert Koch Fellow of the Robert Koch Institute, Berlin, Germany. This work was funded by Centre Of Excellence Phase 2 Grant (BT/PR12817/COE/34/23/2015), (BT/ PR23099/NER/95/632/2017) and (BT/PR23155/NER/ 95/634/2017) by the Department of Biotechnology, Ministry of Science and Technology, to SEH and NZE.

**Conflict of interest**

The authors declare no conflict of interest.

**Author contributions**

JA, JAS, MK, and RP performed the experiments. NZE, MMB, and SEH designed the project. JA, MK, MMB, NZE, RP, and SEH helped in writing the manuscript. JAS, SK, AS, NZE, and SEH helped in the interpretation of results.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Fig. S1. SDS/PAGE image of the purified PE32, PPE65, PE35, and PPE68 proteins.

Fig. S2. Structural analysis of the PE–PPE/PGRS protein family using RONN. Each subfamily was divided into three groups, i.e., completely disordered members with more than 80% disorder, partially disordered members with disorder between 20–80% and ordered proteins having less than 20% structural disorder.

Fig. S3. Structural disorder predictions for PE32, PE35, PPE65, and PPE68. (A) Predicted IUP red disordered regions and disordered binding sites using the ANCHOR tool in PE32 and PPE65. (B) Predicted disordered regions in PE32 and PPE65 using RONN. (C) Predicted IUP red disordered regions and disordered binding sites in PE35 and PPE68 using ANCHOR. (D) Disordered regions in PE35 and PPE68 as predicted by the RONN tool.

Fig. S4. Alignments of our investigated PE and PPE proteins to those with available complex structure. The (A) PPE41, PPE65 and PPE68 proteins as well as the (B) PE25, PE32 and PE35 proteins were aligned using Clustal Omega 1.2.4. The regions of the PPE and PE domains are marked by light grey and yellow backgrounds, respectively.

Fig. S5. Individual recombinant PE35 and PPE68 proteins induce cell proliferation. Different concentrations of individual recombinant proteins were used for re-stimulating the splenocytes in culture for 72 hrs. [3H] thymidine (0.5 mci/ml) was added to each well and incubated at 37 °C for 24 hours. Cells were harvested and β scintillation counter was employed for counting...
Beta activity. Beta activity was significant for the cells re-stimulated with 1 µg/ml or higher dose of PE35 while it was significant for 0.2 µg/ml or higher dose of PPE68, suggesting PPE68 to be more antigenic. Data represents experiments with three mice in each group. Statistical significance was determined by one-way ANOVA and data represented as mean ± SD. p value <0.05 was considered statistically significant.

**Fig. S6.** Antigenicity index of PE35 and PPE68. Antigenicity plots depicting the antigenic regions of PE35 and PPE68.

**Table S1.** Identifiers of the protein family members and secretome proteins used in the analyses.

**Table S2.** Secondary structure content in the investigated recombinant proteins and their complexes in percentage.