Homepeptide Repeats: Implications for Protein Structure, Function and Evolution

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Abstract

Analysis of protein sequences from Mycobacterium tuberculosis H37Rv (Mt H37Rv) was performed to identify homopeptide repeat-containing proteins (HRCPs). Functional annotation of the HRCPs showed that they are preferentially involved in cellular metabolism. Furthermore, these homopeptide repeats might play some specific roles in protein–protein interaction. Repeat length differences among Bacteria, Archaea and Eukaryotes were calculated in order to identify the conservation of the repeats in these divergent kingdoms. From the results, it was evident that these repeats have a higher degree of conservation in Bacteria and Archaea than in Eukaryotes. In addition, there seems to be a direct correlation between the repeat length difference and the degree of divergence between the species. Our study supports the hypothesis that the presence of homopeptide repeats influences the rate of evolution of the protein sequences in which they are embedded. Thus, homopeptide repeat may have structural, functional and evolutionary implications on proteins.

Keywords: Homopeptide repeats; Disordered regions; Replication slippage; Protein domains; Rate of evolution

Introduction

Amino acid repeats are frequently found in protein sequences and are generated by repetitive elements in the genome like long terminal repeats (LTRs) and non-LTRs [1]. These repeats can be further classified into three distinct groups – homopeptide, dipeptide and sequence repeats, based on the number of amino acid residues repeated in the protein sequence [2]. Homopeptide repeats/single amino acid repeats are strings of a single amino acid residue occurring two or more times directly adjacent to each other. A dipeptide repeat is a pair of non-identical amino acid residues tandemly repeated in a linear sequence. Finally, a sequence repeat is an amino acid motif (made up of different combinations of amino acids) that is repeated several times in the protein sequence.

A homopeptide repeat is believed to have arisen either from sequential expansion of short codon repeat through replication slippage or from accumulation of point mutations in the coding sequences [2]. Several studies have shown that accumulation of guanine or cytosine at the third position of every codon in the genome is the fundamental cause for repeat expansion through replication slippage [3–5]. Homopeptide repeats are often embedded in low complexity regions, which also include interrupted and non-tandem repeats [6]. Hydrophobic residues are underrepresented in the repeats with the exception of leucine (L) which is over-represented in bacterial repeats and occurs in large numbers in signal peptides from humans [7]. Zhang et al. examined the location of homopeptide repeats in the protein sequences from Arabidopsis thaliana and Oryza sativa and identified a negative correlation...
between the amino acid distribution, their position and their functions [8]. The positional bias of some repeats to the N-terminal regions implies that these regions are more active in generating repeat sequences.

A recent work by Faux et al. revealed that all repeats (irrespective of the type of amino acid residue involved) play an important role in the processes that require the assembly of large multi-protein complexes [9]. Bjorklund et al. found that tandem repeats have a variety of binding properties and are involved in protein–protein interactions as well as binding to ligands such as DNA and RNA [10]. Moreover, homopeptide repeat that mediates protein–protein interactions can also facilitate network evolution [11]. Many parasitic organisms (Eukaryote or Bacteria) possess surface antigens that are made up of amino acid repeats. When forming an interface between host and pathogen, these repetitive proteins may act as virulence factors and get involved in immune invasion and cytoadherence [12].

Homopeptide repeats are also actively associated with the development of many diseases. For example, poly-glutamine (poly-Q) and poly-alanine (poly-A) repeats are involved in the development of Huntington disease and ocuulopharyngeal muscular dystrophy (OPMD), respectively [13]. Similarly, poly-glycine (poly-G) repeats play an important role in protein targeting [14] and poly-R repeats are involved in binding of viral proteins to RNA [15,16].

In Mycobacterium tuberculosis H37Rv (Mtb H37Rv), 21% of the proteins are hypothetical ones whose function has not yet been determined (results not shown). About 10% of the coding capacity has been dedicated to two Mycobacterium-specific protein families of unknown function, namely the Proline-Glutamate (PE) and Proline-Proline-Glutamate (PPE) families. Proteins belonging to these two families have conserved proline (P) and Q residues in their N-termini [17]. There are many structural and functional studies on Mycobacterium gene products as a whole, but still, no sufficient data is available about the large number of homopeptide repeats present in the proteins.

Although homopeptide repeats have been studied and characterized in many other proteomes, information is unavailable about their structure, function and evolution particularly about those present in prokaryotes. The present study provides an overall picture of the homopeptide repeats present in Mtb H37Rv proteome and the repeats were analyzed in relation to their implication in protein structure, function and evolution.

Results and discussion

During the present analysis, homopeptide repeat-containing proteins (HRCPs) are found to be 289 (a total of 310 repeats), which is about 7.7% of the Mtb H37Rv protein sequences (see Materials and Methods for details). The proportion of the repeat-containing proteins in Mtb H37Rv is similar to that of yeast; whose protein sequences contain 7.6% homopeptide repeats [18].

Faux et al. previously reported that in prokaryotes, homopeptide repeats are commonly made up of serine (S), G, A and P, while in eukaryotes, the amino acids Q, asparagine (N), A, S and G are often seen in homopeptide repeats [9]. Figure 1 illustrates the rate of occurrence of homopeptide repeats based on their repeat frequencies. From the figure, it is evident that four distinct classes of homopeptide repeats containing A, G, arginine (R) or P are abundant in Mtb H37Rv. These results are consistent with that of Faux et al. Green and Wang proposed that the repeats formed by hydrophilic amino acids are abundant in the protein sequence databases [19]. However, it is interesting to note that a majority of the homopeptide repeats observed in the present study are composed of hydrophobic amino acid residues which in turn could affect the overall hydrophobicity of the protein [20]. In addition, Marcotte and his co-workers proposed that R repeats are highly depleted in protein sequences [21]. Nevertheless, in the Mtb H37Rv genome, R repeats constitute about 8% of the repeats present (Figure 1). Arginine being a charged amino acid may play an important role in protein–protein interactions.

Functional annotation of HRCPs in Mtb H37Rv

Experimental evidence suggests that HRCPs have biased functions [8,9,18]. Functional annotation of the HRCPs is carried out using Yeast Protein Database (YPD) to evaluate their abundance in different classes of proteins. Results showed that, in yeast, homopeptide repeats are overrepresented in transcription factors, protein kinases and transporter proteins [9]. These data are consistent with

Figure 1 Total number and percentage of homopeptide repeats in Mtb H37Rv

A pie chart describing the total number and the percentage of the homopeptide repeats. The residues A (blue), G (green), R (red) and P (violet) occur most often among the homopeptide repeats.
studies by Gerber et al. indicating that the insertion of either poly-Q or poly-P tracts enhances the transcriptional activation of the GAL4/VP16 fusion construct in yeast [22]. Furthermore, analysis on the HRCPs from Homo sapiens, rodents, Drosophila melanogaster, Caenorhabditis elegans, Oryza sativa and Arabidopsis thaliana all showed similar observations [9,23].

In *Mtb* H37Rv, a total of 289 HRCPs exhibit have functional annotations. However, and the functions of these HRCPs are not well established. 58% (168 out of 289) of the HRCPs were assigned to four protein families – polymorphic GC-rich repetitive sequence (PGRS) proteins, PPE proteins, PE proteins and hypothetical proteins (Figure 2). However, about 22% (65 HRCPs) of the HRCPs are involved in cellular metabolism and 12% (34 HRCPs) of the proteins participated in transport and signaling process (grouped as “Transport” in Figure 2). Only 3% (10 HRCPs) of HRCPs belong to either transcription or translation apparatus, indicating that in *Mtb* H37Rv, repeats do not exhibit bias towards transcriptional or translational proteins. This may be due to the fact that bacteria have simple intracellular mechanisms. Thus, from the present study, it appears that the selective enrichment of homopeptide repeats towards transcription factors and signaling proteins does not extend to *Mtb* H37Rv protein sequences.

### Identification of homopeptide repeats in 3D structure

Previous studies reported that compared to the protein sequence database, there is a deficiency of simple sequence repeats in PDB [25,26]. Homopeptide repeats encoded by iterations of a single codon likely result from strand slippage and substantial increase in multiple codons within the homopeptide coding regions which makes the homopeptides important for the structure of the proteins containing them [27]. The 3D structures of only five HRCPs from *Mtb* H37Rv are available in PDB (Table 1). Among these five structures, two repeats interact with a metal ion (histidine tracks in zinc and ferric uptake regulator protein with PDB ID 2O03). Poly-A repeats are well-known for their helical structure [28] and in Mycobacterial protein sequences, poly-A stretches form helical structures too.

As most of the HRCPs in *Mtb* H37Rv are yet to be crystallized, an extensive search for the homopeptide repeats in PDB was carried out using the BSDD server [24]. Simultaneously the corresponding secondary structures were analyzed using the STRIDE software [29]. Serine, glycine and arginine repeats have high probability of forming β turn conformation, while leucine is involved in α helical conformation. Protein domains tend to exhibit stable yet flexible conformations when present in the short linker groups. Thus, homopeptide expansions are tolerated when they occur in the linker regions. In *Mtb* H37Rv, histidine repeat in the linker region is found in serine/threonine pro-

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**Figure 2 Functional annotation of HRCPs in *Mtb* H37Rv**

The homopeptide repeats were analyzed and categorized into eight classes. The number of repeat containing proteins has been provided on the X-axis and their corresponding functional class is provided on the Y-axis. The highest number of repeat containing proteins is the metabolic and hypothetical domains (Table S1). To deduce the functions of these repeats in the domains, the three-dimensional (3D) structures of the HRCPs were given as input to the BSDD server [24]. The server identified two distinct domains, including peptide (small fragments) containing arginine repeats within the RNA-binding domain, and biotin carboxylase containing glycine within ATP-binding domain.
Table 1 Structural homopeptide repeats present in Mtb H37Rv protein sequences

| PDB ID | Protein description               | Repeat type | Secondary structure |
|--------|-----------------------------------|-------------|---------------------|
| 2O03   | Zinc uptake regulator protein     | HHHHH       | β-sheets            |
| 1RWI   | Serine/threonine protein kinase    | HHHHHHH     | β-turns             |
| 1U5H   | Citrate lyase beta subunit        | AAAA        | α-helix             |
| 2BPQ   | Anthranilate phosphoribosyl transferase | AAAAA    | α-helix             |
| 3EKL   | Fructose 1,6-bis phosphate aldolase | TTTTT     | β-hairpins          |

Figure 3 The poly-histidine tract occurs in the linker regions of protein kinase PknD. The poly-H track (residues 265–270) was revealed in the linker regions of Ser/Thr protein kinase PknD (PDB ID: 1RWI). Poly-H was shown in black.

Table 2 Average RONN score for the homopeptide repeats in the functionally annotated HRCPs

| Repeat type | Number of repeats | Average RONN score |
|-------------|-------------------|--------------------|
| Alanine (A) | 78                | 0.49               |
| Arginine (R)| 11                | 0.64               |
| Proline (P) | 12                | 0.76               |
| Glycine (G) | 6                 | 0.52               |
| Histidine (H)| 1              | 0.51               |
| Valine (V)  | 8                 | 0.33               |
| Threonine (T)| 3               | 0.47               |
| Leucine (L) | 2                 | 0.32               |
| Serine (S)  | 1                 | 0.52               |
| Glutamine (Q)| 1              | 0.91               |
| Phenylalanine (F)| 1             | 0.19               |

0.5, suggesting that these repeats are disordered, hence lack a definite tertiary structure. In addition, two repeat types made up of A or T demonstrate an average score of 0.47 (can be considered as 0.5). Thus, it is reasonable to conclude that 90% of the homopeptide repeats found in Mtb H37Rv are disordered since 9 out of 11 repeat types observed have a RONN score greater than or equal to 0.5 (Tables 2 and S2).

Repeats present in the disordered regions

Disordered regions are parts within the protein molecule that do not fold into a stable secondary structure. These regions may vary in size [30,31] and are involved in many biological processes such as regulation, signaling and cell cycle control [32]. To check if homopeptide repeats are disordered, the HRCPs were given as input to the RONN software. An amino acid residue is determined as disordered if the average probability of disorder (also known as RONN score) is greater than 0.5 [33]. As shown in Tables 2 and S2, six homopeptide repeat types made up of R, P, G, H, S or Q have a RONN score greater than 0.5, suggesting that these repeats are disordered, hence lack a definite tertiary structure. In addition, two repeat types made up of A or T demonstrate an average score of 0.47 (can be considered as 0.5). Thus, it is reasonable to conclude that 90% of the homopeptide repeats found in Mtb H37Rv are disordered since 9 out of 11 repeat types observed have a RONN score greater than or equal to 0.5 (Tables 2 and S2).

Amino acid usage, frequency of amino acid repeats and its usage

We next tested whether there is a linear correlation between the frequency of amino acid repeats and the usage of the amino acid in the protein sequences in Mtb H37Rv. The compositional analysis of all the protein sequences shows that A, G, L and V are found in abundance (Figure 4). In the present study, valine is not further considered since the overall count of valine rich repeats is low. There is a significant difference in the amino acid usage and the frequency of the corresponding homopeptide repeat present in all the Mtb H37Rv protein sequences. The percentage of the homopeptide repeats is plotted against the composition of the corresponding amino acids in the protein sequences in Figure 4. From this figure, it is evident that only four amino acid residues (A, G, P and R) have a higher tendency to form repeat sequences. Although the other amino acid residues (like L) are present relatively high, they possess a very low propensity to form homopeptide repeats. These results are consistent with the hypothesis proposed by Zhang et al. that poly-G repeats are solely responsible for the high glycine content in Oryza sativa and Arabidopsis thaliana [8]. These data suggest that there is no linear relationship between the frequency of the amino acid repeats and amino acid composition [1]. Thus, it is possible that other than the amino acid content, many factors are responsible for the occurrence and frequencies of the homopeptide repeats.

Difference in repeat size

Comparative studies on repeat size difference between species have dealt only with a small number of genes. It is therefore unclear whether the rapid change reported previ-
ously is representative of homopeptide repeats in general or whether conserved homopeptide repeats also exist [18]. To address this question, we compared the homopeptide repeats from four homologous species. Repeat size difference between homologous proteins indicates a degree of repeat conservation. The size difference ranges between 0 and 1, where 0 represents complete conservation of the repeats and 1 implies complete absence of the repeats between two homologous sequences [34].

We examined the HRCPs from *Mtb* H37Rv and their putative homologs from three other organisms including *Escherichia coli* (Bacteria), *Sulfolobus acidocaldarius* DSM639 (Archaea) and *Homo sapiens* (Eukaryote). We found that the repeat sequences followed a regular pattern of conservation and divergence between the four species (Figure 5). The repeat size difference between *Mtb* H37Rv and *E. coli* is low, compared to that between *Mtb* H37Rv and *S. acidocaldarius* DSM 639 or *H. sapiens* (Table S3 and Figure 5). It is interesting to note that the repeats are more conserved in *Mtb* H37Rv and *E. coli* and least in *H. sapiens*, demonstrating that there exists a direct relationship between the repeat size difference and the order of divergence.

**Evolutionary rate analysis**

Low-complexity regions that include homopeptide repeats evolve much faster than the rest of the protein sequence [35] and homopeptide repeats are present within functionally and structurally more evolutionarily active regions [36]. It was hypothesized that repeat expansion and contraction may provide a mechanism for rapid morphological evolutionary changes [37] and that this expansion and contraction may be facilitated by replicative slippage [38,12]. An evolutionary analysis of 2838 open reading frames from three Saccharomyces species showed that fast evolving low-complexity sequences outnumbered conserved sequences by a ratio of 10–1 [35]. Previous studies on Hox proteins by Casillas et al. showed that the long homopeptides are present all along the protein except in the highly conserved regions [36]. These repeat regions are the origin of most of the indels and thus are responsible for the high amino acid evolution of Hox proteins.

Previous studies show that homopeptide repeats have an influence on the evolutionary rate of the protein sequences in which they are embedded [25,37,39]. To see whether this holds true in Mycobacteria, orthologous proteins from

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**Figure 4** Comparison of amino acid usage in homopeptide repeats and proteome in *Mtb* H37Rv

The amino acid usage (%) in both homopeptide repeats (homerepeats) and in the protein sequences (proteome) was provided here. The amino acid residues A, G, P and R exhibit peak usage in both homopeptide repeats and the total protein sequences.

**Figure 5** Repeat size differences between proteins from *Mtb* H37Rv, *E. coli*, *S. acidocaldarius* and *H. sapiens*

The repeat size difference between *Mtb* H37Rv and *E. coli* is low, compared to that between *Mtb* H37Rv and *S. acidocaldarius* DSM 639 or *H. sapiens*. nun
seven Mycobacterial species are included in the present study. The occurrence of homopeptide repeats is consistent among the seven Mycobacterial species, however, their frequencies are not similar (Table 3). Twenty orthologous proteins contained several homopeptide repeats in common. Multiple sequence alignment for all 20 orthologous proteins showed that the homopeptide repeats observed in these proteins are conserved among the seven Mycobacterial species. An example of the alignment is provided in Figure 6 for mmpS3. The repeats in the protein sequences are conserved at different levels, which is evident from the expansion and contraction of the repeats in the seven species, e.g., the proline tract found in mmpS3 shown in Figure 6. Furthermore, we constructed the phylogenetic tree based on the multiple sequence alignment of all 20 proteins. The evolutionary rates with repeat blocks, E(R), and without repeat blocks, E(A), were calculated for all 20 proteins using PROTDIST and FITCH programs [40] (Table 4). Paired t-test analysis showed that the evolutionary rates of all 20 proteins with repeat blocks are significantly different (at 0.05 level) from that of the proteins without repeat blocks. In addition, a binomial testing was performed for all 20 proteins and the resulting Z-score (4.504) supported the significant difference (at 0.01 level). Thus, E(R) and E(A) result from different populations and the mean difference observed between them is not a consequence of coincidence or random sampling. Taken together, these data indicate that the differences observed in the experiment are not a consequence of coincidence or random sampling. Therefore, it is reasonable to conclude that proteins with homopeptide repeats E(R) evolve faster than the proteins without repeats E(A).

### Conclusion

The present analysis shows that the homopeptide repeats are common in Mtb H37Rv and four major classes are identified, namely, poly-A, poly-G, poly-R and poly-P con-
taining repeats. Structural and functional analyses show that each class of homopeptide repeats adopt a distinct secondary structure and are mainly found in proteins involved in cellular metabolism, transport and signaling. Analysis of the homopeptide repeat frequency and the amino acid usage suggests that the amino acid content is not the only factor responsible for changes in homopeptide repeat frequencies. Finally, homopeptide repeats could have a profound effect on the evolutionary rate of the protein containing it. Thus, it can be concluded that homopeptide repeats play a vital role in the structure, function and evolutionary rate of the homopeptide repeat containing proteins.

Materials and methods

Identification and distribution of homopeptide repeats in Mtb H37Rv

All the protein sequences from Mtb H37Rv (3988 protein sequences) were downloaded from the local FTP site maintained at the Bioinformatics centre, Indian Institute of Science. Locally developed PERL scripts were used to detect the homopeptide repeats in the protein sequences. Homopeptide repeats defined here refer to continuous occurrence of single amino acid residues for five or more times. The cut-off size of five residues was chosen because of its significantly low probability of occurrence by chance.[41]

Repeat frequency = Number of X repeat/total number of homopeptide repeats present in the protein sequences, where X represents any amino acid residue.

Functional annotation of HRCPs

Functional annotation of HRCPs was done using NCBI BioSystems [42] and COG databases [43]. NCBI BioSystems database was used to detect the protein structures involved either in a biological or disease pathway. In the case of COG, the most straightforward application is the prediction of individual protein function using the COGNNITOR program. For each HRCP, the NCBI RefSeq_ID (Reference Sequence IDentifier), description, homopeptide tract and function were recorded.

Scanning for domains containing homopeptide repeats

The HRCP sequences were searched against the Pfam database [44] to identify the functional domains present. Two programs on the PDBsum [45] server namely, LIGPLOT [46] and NUCPLOT [47], was used to analyze the molecular interactions. LIGPLOT gives the schematic representation of the interactions between the ligand and the amino acid residues, while NUCPLOT shows the protein-nucleic acid interactions. Here again, the RefSeq_ID, description, repeat type, domain name and Pfam identification were recorded for each homopeptide repeat present within the protein domain.

Identification of homopeptide repeats in 3D structure

To identify the secondary and tertiary structures of the HRCPs, the PDB archive was searched with the standalone versions of BLASTP [48] and STRIDE softwares [29].

Identification of homopeptides present in the disordered regions of the proteins

Functionally annotated HRCP sequences were probed for disordered regions. RONN [33] software was used to detect the degree of disorder in the HRCP sequences.

Calculation of the amino acid usage

The amino acid usage and the homopeptide repeat fraction (number of homopeptide repeats/total number of proteins) in Mtb H37Rv were calculated using locally developed PERL scripts. The amino acid frequency of all the protein sequences was calculated using the formula:

Amino acid frequency = Number of X residues in the protein sequence/total number of amino acid residues in the protein sequences; where X represents any amino acid residue.

The amino acid frequencies of all the protein sequences were compared with the homopeptide repeat fraction observed.

Calculation of the repeat size difference

To investigate the purifying selection pressure on homopeptides, putative homologous HRCPs from Mtb H37Rv, E. coli (Bacteria), Sulfolobus acidocaldarius DSM639 (Archaea) and Homo sapiens (Eukaryote) were detected using the BLASTP [47] search (against the nr database). The putative homologous sequences were aligned using ClustalW [49] and scanned for the homopeptide tract. The numbers of amino acids in homopeptide tracts of the three homologs were recorded.

Evolutionary rate analysis

HRCPs from Mtb H37Rv were given as query to the program BLASTP [48]. Using H37Rv as the template, 20 orthologous proteins were detected in six other Mtb species (Mycobacterium marinum, Mycobacterium smegmatis MC2155 strain, Mycobacterium ulcerans Agy99 strain, Mycobacterium vanbaalenii PYR-1 strain, Mycobacterium sp. MCS strain and Mycobacterium gilvum PYR – GCK strain). These orthologous protein sequences were aligned using ClustalW [49] and the alignments were scanned to identify the homopeptide repeats of at least five residues
long. The multiple sequence alignment of the proteins was provided as input to the PROTDIST program of the PHYLIP (the PHYLogeny Inference Package) software [40] for the construction of distant matrices (based on Jones-Taylor-Thornton distance model), which were then used to construct phylogenetic trees using the FITCH program in the same package. The sum of the branch lengths in the phylogenetic tree provided the evolutionary rates for the proteins with repeat sequences. Further, to investigate the influence of the repeats in the evolution of the protein, the repeat blocks were removed and the above mentioned procedure was applied to the protein sequences.

Authors' contributions

KS conceived and supervised the project, and critically analyzed the database. MUK, BB, SP, JJ and KS collected the data and performed various analyses. MG performed the statistical analysis for validation. BB drafted the manuscript. MKV critically reviewed the manuscript and participated in discussion, curation and validation. KS revised the manuscript. All authors read and approved the final manuscript.

Competing interest

The authors have no competing interest to declare.

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Supplementary material

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