The many faces of the helix-turn-helix domain: 
Transcription regulation and beyond

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Abstract

The helix-turn-helix (HTH) domain is a common denominator in basal and specific transcription factors from the three superkingdoms of life. At its core, the domain comprises of an open tri-helical bundle, which typically binds DNA with the 3rd helix. Drawing on the wealth of data that has accumulated over two decades since the discovery of the domain, we present an overview of the natural history of the HTH domain from the viewpoint of structural analysis and comparative genomics. In structural terms, the HTH domains have developed several elaborations on the basic 3-helical core, such as the tetra-helical bundle, the winged-helix and the ribbon-helix–helix type configurations. In functional terms, the HTH domains are present in the most prevalent transcription factors of all prokaryotic genomes and some eukaryotic genomes. They have been recruited to a wide range of functions beyond transcription regulation, which include DNA repair and replication, RNA metabolism and protein–protein interactions in diverse signaling contexts. Beyond their basic role in mediating macromolecular interactions, the HTH domains have also been incorporated into the catalytic domains of diverse enzymes. We discuss the general domain architectural themes that have arisen amongst the HTH domains as a result of their recruitment to these diverse functions. We present a natural classification, higher-order relationships and phylogenetic pattern analysis of all the major families of HTH domains. This reconstruction suggests that there were at least 6–11 different HTH domains in the last universal common ancestor of all life forms, which covered much of the structural diversity and part of the functional versatility of the extant representatives of this domain. In prokaryotes the total number of HTH domains per genome shows a strong power-equation type scaling with the gene number per genome. However, the HTH domains in two-component signaling pathways show a linear scaling with gene number, in contrast to the non-linear scaling of HTH domains in single-component systems and sigma factors. These observations point to distinct evolutionary forces in the emergence of different signaling systems with HTH transcription factors. The archaea and bacteria share a number of ancient families of specific HTH transcription factors. However, they do not share any orthologous HTH proteins in the basal transcription apparatus. This differential relationship of their basal and specific transcriptional machinery poses an apparent conundrum regarding the origins of their transcription apparatus.

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1. Introduction

The general paradigms for the processes of transcription initiation and regulation first emerged from the pioneering studies on gene expression in bacteria and phages [1]. Transcription in bacteria was found to be catalyzed by a single multi-subunit RNA polymerase, which is recruited to promoters by means of a DNA-binding protein, the sigma factor that recognizes specific sequences upstream of genes [2–4]. The sigma factor and the RNA polymerase, together, constitute the basal transcription apparatus that is required for the baseline transcription of all genes. Early studies, especially in the Bacillus subtilis sporulation model, suggested that there may be more than one sigma factor that might recruit the catalytic core of the RNA polymerase to alternative sets of genes. This provided a mechanism for regulating the broad changes in gene expression, which correlate with the different developmental or differentiation states of a bacterium [5,6]. A number of early studies on metabolic regulation in bacteria also indicated that there are other regulatory DNA-binding proteins that act as switches to control the expression of specific smaller sets of genes. These sets of genes are often collinear on the chromosome, and encode components of a common pathway for the utilization of a particular environmental metabolite (e.g. lactose), or constitute interacting components of a developmental pathway (e.g. lytic or lysogenic development of phages). These regulatory proteins, termed the specific transcription factors, were found to belong to two distinct functional types: (1) those which negatively regulated transcription of their target gene (repressors) and (2) those which positively regulated transcription of their target genes (activators) [1]. The affinities of the specific transcription factors for their target sequences on DNA were often found to be dependent on their interaction with low-molecular weight compounds (effectors), which bound to them, or phosphorylation and other post-transcriptional modifications [1]. When transcription in eukaryotes was first investigated, several differences in the subunit composition and architecture of the basal transcription machinery and specific transcription factors were noted [7,8]. However, the basic regulatory mechanisms in bacterial transcription, which were brought to light in early studies, remained applicable in a generic sense across the entire Tree of Life [1].

The pioneering investigations of Matthews, Ohlendorf, Sauer, Doolittle and co-workers in 1982 provided the first glimpse of the features unifying diverse transcription regulators [9–13]. They showed that the phage lambda transcription regulators, cro and the cI repressor, and lacI, the lactose operon repressor, shared a similar tri-helical DNA-binding domain. The 2nd and the 3rd helices of this tri-helical domain constituted a Helix-Turn-Helix motif, and this motif was shown to be a critical determinant of their interaction with DNA. Thus, these DNA-binding domains came to be
referred as the helix-turn-helix (HTH) domain. Sequence and secondary structure analysis by these pioneering workers suggested that the HTH domain was a common DNA-binding motif found in several other bacterial repressors as well as activators, such as the cAMP-dependent catabolite activator protein. They perceptively suggested that all these DNA-binding domains have descended from a common ancestor, through duplication and divergence, thereby generating the diversity of transcription regulators that regulate bacterial and phage genes \([9,11]\). Subsequent sequence analysis revealed that DNA recognition by sigma factors was also mediated by HTH domains, similar to those observed in the specific transcription factors \([14–16]\). In the second half of the 1980s the conserved domains of transcription factors regulating eukaryotic development and differentiation, namely the homeodomains and Myb domains, were also noted to possess the HTH fold \([17,18]\). These and other investigations suggested the general significance of this module in DNA–protein interactions across a wide phylogenetic spectrum \([18–21]\).

An explosion of structural studies in the 1990s, while strengthening the basic structure–function relation between the HTH domain and DNA binding, also produced a large amount of data regarding the diversity of DNA–protein interactions mediated by different versions of the HTH domain. A number of sequence and structural analysis studies also uncovered HTH modules in several specific eukaryotic transcription factors, chromatin proteins like histone H1, and basal transcription factors such as TFII B and TFII E \([22–30]\). These findings lead to the idea that the HTH domain is probably one of the most ancient conserved features of transcription apparatus, which was already present in the last universal common ancestor of all extant life forms (LUCA). Studies during this period also showed that although archaea and eukaryotes possess a similar basal transcription machinery, the specific transcription factors of the former are clearly closer to those of the bacteria than the eukaryotes \([30–34]\). The other major development in the later half of the 1990s was the birth of the genomic era, unleashing the power of comparative genomics. Starting with the earliest comparative genomic studies it became apparent that the HTH domain was a highly prevalent domain in prokaryotes \([35]\). Comparative analysis also helped in identifying several major monophyletic assemblages of HTH transcription factors, each distinguished by their own distinctive sequence and structure features (for example see Refs. \([36–41]\)). These classes often showed one or more distinctive domain architectures – i.e. the fusion of the HTH domain with additional globular domains in the same polypeptide. These globular domains, which are linked to the HTH show a bewildering diversity, and point to the immense variety of functional contexts in which the HTH domain may be deployed. The combined use of genome sequence data and high-throughput expression data also cast light on the multi-level transcriptional regulatory networks in prokaryotes in which these HTH-containing proteins functionally interact to maintain a particular transcriptional state in the cell \([42,43]\).

Over the years, the recruitment of the HTH domains to biological functions beyond transcriptional regulation has also become apparent. Some of these proteins, participating in functions such as DNA repair and RNA metabolism, exploit the nucleic acid-binding properties of the HTH just as in the case of transcriptional regulation (for example see: \([44–48]\)). However, there are other instances where the HTH may be adapted to very different functions, such as mediating specific protein–protein interactions, or as a structural unit of a larger enzymatic domain \([49–51]\).

In this article we aim at providing a synthetic overview of the structural, functional and evolutionary diversity of the HTH domain from the vantage point of over two decades of intense investigation. We currently enjoy unprecedented advantages due to an enormous amount of genomic data, numerous high resolution structures, functional studies and sensitive computational tools to capture the highlights of the natural history of HTH domains. Our focus is principally, but not exclusively, on versions of the domain found in the prokaryotic super-kingdoms. We first discuss the structural diversity seen in this fold, followed by a discussion of the unifying themes in domain architectures of HTH proteins and their significance to different biological functions. We next provide a higher-order natural classification of these domains and discuss their genome-wide demography in light of the general adaptive tendencies that can be inferred from comparative genomics. In this context we also consider the adaptation of the HTH to functional roles beyond transcriptional regulation. Finally, we discuss the relevance of the information gleaned from these diverse areas in reconstructing the origin and evolution of transcription and its regulation.

### 2. Structural scaffold of the HTH domain and its diverse elaborations

The basic HTH domain is a simple fold comprised of three core helices that form a right-handed helical bundle with a partly open configuration. When it is displayed by placing the third helix in the front and in the horizontal orientation, the 3 helices of the domain form an approximately triangular outline (Fig. 1). We use this as the default orientation for all further illustrations and discussions. The characteristic sharp turn, which is a defining feature of this domain, is situated between the 2nd and the 3rd helix, and typically does not
tolerate insertions or distortions. However, the loop between helix-1 and helix-2 shows far greater variability and may accommodate several modifications in the different classes of HTH domains. Furthermore, diverse N- and C-terminal extensions to the core tri-helical bundle are also encountered in several classes of HTH domains. The partly open configuration of the bundle results in a shallow cleft between helix-3 and helix-1 on the side opposite to helix-2. This cleft appears to have acted as a structural niche that favored the evolution of additional structural elements to pack into it via hydrophobic interactions. Thus, most of the extension to the core HTH domain are structural elements that appear to have evolved to generate a more closed configuration by interacting with the cleft (Fig. 1; see below for further discussion). The 3rd helix, known as the recognition helix, typically forms the principal DNA–protein interface by inserting itself into the major groove of the DNA [19,24,52]. Nevertheless, the individual residues involved in DNA contacts may widely vary across the fold. Additional secondary contacts with DNA may be mediated by other parts of the structure or even extensions outside of the core HTH domain, such as a basic patch at the N-terminus of helix-1 [52].

Despite the great sequence diversity observed in this fold, there are a few sequence elements that are widely conserved in members of the fold. The most characteristic of these is the “shs” pattern (where ‘s’ is a small residue, most frequently glycine in the first position, and ‘h’ is a hydrophobic residue) that lies in the turn between helix-2 and helix-3 of the core HTH structure. The other well-conserved signatures is the “phs” (where ‘p’ is a charged residue, most frequently glutamate) that is present in helix-2 (Fig. 1). The conserved hydrophobic residues in these motifs, together with at least two other conserved-hydrophobic residues seen in helix-1 and helix-3 localize to the interior and form the characteristic hydrophobic core that stabilizes the domain (Fig. 1). The conservation of these elements across diverse members of this fold from the 3 superkingdoms of life, taken together with the conserved structure–function associations of this fold strongly supports the monophyletic origin of HTH domains [9,11] from a common ancestor that bore the above-mentioned sequence features.

Based on their distinctive features, members of the HTH fold can be divided into two major structural classes and a few other highly derived structural classes that contain drastic alterations of the core HTH domain.

2.1. HTH domains with a simple three-helical bundle and its extensions

The simplest version of the HTH domain, the basic tri-helical version, is comprised entirely of the three core helices with no additional elaborations (Figs. 1 and 2). This configuration appears to be closest to the ancestral state of the domain and is widely seen across the three super-kingdoms of life. This version is represented by the HTH domains such as the region-3 and region-4 of the sigma factors, the RPB10 subunit of the DNA-dependent-RNA polymerases of the archaeo-eukaryotic lineage, bacterial transcription factors of the FIS family, the eukaryotic Homeo, POU and Myb domains, the eukaryotic BRCA2 tower domain, and the paired module and related HTH domains of transposases and resolvases (Fig. 2). A distorted version of the basic tri-helical HTH domain is also seen in the zinc-chelating domain.
of several retroviral integrases [53], in which helix-3 is packed more closely against the other helices by means of a Zn ion chelated by conserved cysteines and histidines at the ends of helix-1 and helix-3 (Fig. 1). Likewise, in RPB10, a set of zinc-chelating cysteines help in stabilizing an N-terminal loop against helix-3 [54].

The tetra-helical version of HTH domain is an elaboration of the basic tri-helical version and is characterized by an additional C-terminal helix which packs against the shallow cleft formed due to the open configuration of the tri-helical core (Figs. 1 and 2). Several major families of prokaryotic transcription factors display this version of the domain. The multi-helical version, typified by the archaeo-eukaryotic basal transcription factor TFIIB, is a further elaboration of the tetra-helical HTH, wherein two additional helices have been added to the N-terminus of the tetra-helical core, resulting in a larger globular helical bundle (Fig. 2). The Bright (ARID) domain, a eukaryotic DNA-binding domain [55], is another more divergent version of the TFIIB-like HTH domains. This version of the fold is very infrequent in the bacteria, and is represented by a circularly permuted version seen in the sporulation regulator Spo0A from spore-forming Gram-positive bacteria [56]. Other versions, which represent relatively infrequent elaborations of the basic tri-helical version, are the KorB-like HTHs [57] and the FlhD-like HTHs [58]. The former version is characterized by an additional N-terminal helix that packs against the basic 3-helix core and the later contains a C-terminal helical extension that packs very differently from the helical extension seen in the above-mentioned classical tetra-helical forms (Fig. 2).

2.2. The winged HTH domain

The winged HTH (wHTH) domains are distinguished by presence of a C-terminal β-strand hairpin unit (the
The simplest versions of the wHTH domains contain a tight helical core similar to basic tri-helical version followed by the two-strand hairpin (Figs. 1 and 2). However, many wHTH domains display further serial elaborations of the β-sheet. In the 3-stranded version, the loop between helix-1 and helix-2 assumes an extended configuration and is incorporated as the 3rd strand in the sheet, via hydrogen bonding with the basic C-terminal hairpin (Fig. 2). The DNA-binding domain of the bacterial transcription regulator MerR defines an aberrant derivative of a 3-stranded version of the wHTH domain, in which helix-1 has been lost (Fig. 2). In addition to the MerR family of bacterial repressors this version of the fold is seen in a wide variety of phage, bacterial and eukaryotic DNA-binding proteins and translation factors. The topoisomerase II family has two copies of wHTH domains [63]. One of these wHTH domains contains a large insert between helix-1 and helix-2. This insert contains an extended β-sheet structure that forms a brace around double-stranded DNA. Also present in this insert is a second wHTH domain which is circularly permuted with helix-1 occurring to the C-terminus of the wing. The two structurally related ribosomal proteins L11 and S18 represent another distinctive derivative of wHTH domain. While they share a β-strand hairpin between helix-1 and helix-2 with the 4-stranded wHTH, unlike the latter they possess only a single C-terminal strand (Fig. 2). A highly modified HTH domain seen in the catalytic domain of the phage integrases shows a rare insertion between helix-2 and helix-3 of the core domain [63]; however, this insert does not distort the structure of the core. The FF-domain [64] and the C-terminal domain of the PP2C protein phosphatase [65] define another highly modified version of the HTH domain that is currently only known from eukaryotes. This version domain shows the insertion of a helical insert between helix-1 and helix-2 resulting a different packing of the elements and a distortion of the fold (Fig. 2).

3. General and specific aspects of the domain architectures of HTH proteins

HTH domains are combined with other domains in the same protein giving rise to an astonishing array of domain architectures (Table 1 and Fig. 3). Despite the diversity, all the architectures can be classified into a small number of generic architectural classes, the members of each class being unified by certain general organizational and functional principles. These generic architectural classes illustrate how natural selection has convergently engineered similar functional solutions using a relatively small repertoire of domains [66]. Hence, the more highly populated architectural classes
are likely to represent the more successful functional solutions in which the HTH domain has been involved.

3.1. Simple architectures involving the HTH domain

The simplest architectures involving the HTH domain are seen in certain proteins related to the cI repressors (e.g., the archaeal repressors typified by AF1793), most proteins of the MetJ-Arc superfamily and the Fis proteins. These proteins are almost entirely comprised of just a standalone HTH, and might, at best, have some small extensions that play a role in dimerization or interactions with other components of the transcriptional machinery. The eukaryotic ribosomal protein S10 and the histone H1 respectively contain RNA and DNA binding versions of the wHTH domain, which is fused to low-complexity sequence that forms a non-globular tail [67]. Such non-globular extensions that play a role in non-specific nucleic acid contacts and protein contacts during transcription activation are very common in eukaryotic transcription factors with HTH domains [41]. A family of bacterial proteins typified by the B. subtilis sigma D regulator YhlX (SwrB) [68,69] contains a HTH domain fused to a N-terminal transmembrane region (Fig. 3). These HTH proteins might regulate transcription under the influence of signaling events associated with the cell membrane. The next level of

Fig. 3. Examples of domain architectures of proteins with HTH domains. The 3 panels show the HTH domains in one component systems, those in two component systems and those fused to enzymatic domains. Below each protein the name of the protein or its family is indicated. The domain names and abbreviations are as shown in Table 1. Additional abbreviations are H – any HTH domain; wH – winged HTH; Assd – archaeal specific signaling domain; Acyclase – adenylyl cyclases; AATRS – aminoacyl tRNA synthetase; TPR – tetratricopeptide repeats; HAMP – domain present in histidine kinases, adenylyl cyclases, methyl-accepting proteins and phosphatases, chro-chromodomain BTAD-conserved domain found in bacterial signaling proteins, CPS-D1, the domain 1 of the large subunit of the carbamoylphosphate synthetases, MGS- the methylglyoxal synthase domain that binds ornithine in the carbamoylphosphate synthetases, ParB- the OB-fold nucleic domain seen in ParB proteins, B-hel- a conserved beta-barrel seen in the Lhr helicases. The transmembrane regions are indicated by yellow boxes. The grey boxes represent uncharacterized globular domains that are unique to a particular family of proteins. Af: Archaeoglobus fulgidus, Ana: Nostoc sp., Blic: Bacillus licheniformis, Bs: Bacillus subtilis, Bthe: Bacteroides thetaiotaomicron, Ec: Escherichia coli, Llac: Lactococcus lactis, Lmon: Listeria monocytogenes, Mjan: Methanococcus jannaschii, Mlo: Mesorhizobium loti, Mmaz: Methanosarcina mazei, Mcap: Methylcoccus capsulatus, Mthe: Methanothermobacter thermautotrophicus, Mtu: Mycobacterium tuberculosis, Phor: Pyrococcus horikoshii, Poke: Planomicrobium okeanokoites, Tm: Thermotoga maritima, Sent: Salmonella enterica, Spyo: Streptococcus pyogenes, Hs: Homo sapiens, Sc: Saccharomyces cerevisiae.
Table 1
Globular domains frequently linked to the HTH domain in the same polypeptide

<table>
<thead>
<tr>
<th>Domain</th>
<th>Structure</th>
<th>Placement of HTH</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Domains in two-component, phosphorelay and S/T kinase signaling cascades</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>REC (receiver domain)</td>
<td>αβ. [Flavodoxin-like topology], PDB 1NTR</td>
<td>1.19N; 78.35C</td>
<td>Is phosphorylated on an aspartate residue by the histidine kinase dependent phospho-relay system. Typically found fused to OmpR and LuxR-family (in NarL-like proteins) HTH domains</td>
</tr>
<tr>
<td>Histidine kinase</td>
<td>α + β, PDB 1BXD</td>
<td>0.10N; 0.03C</td>
<td>Usually occurs as a standalone subunit, but on few instances is fused to the downstream transcription factor with receiver and HTH domains</td>
</tr>
<tr>
<td>3H</td>
<td>α + β fold</td>
<td>1.00N</td>
<td>A domain sharing a common fold with the HPr domain of the PTS system. It has 3 conserved histidine residues that are likely to be phosphorylated</td>
</tr>
<tr>
<td>FHA</td>
<td>β fold, PDB 1LGQ</td>
<td>0.23N; 0.16C</td>
<td>A phosphoserine/threonine peptide-binding domain. Combinations with HTHs are expanded mainly in Actinomycetes, which have numerous serine/threonine kinases</td>
</tr>
<tr>
<td>HPT (histidine phosphotransfer domain)</td>
<td>α helical domain PDB 1QSP</td>
<td>–</td>
<td>Receives a phosphate on a conserved histidine in the two-component relay. Found fused to OmpR family wHTH domains along with Receiver domains in cyanobacteria</td>
</tr>
<tr>
<td>PRD (PTS Regulatory Domain)</td>
<td>α helical domain PDB 1H99</td>
<td>3.84N</td>
<td>Typically phosphorylated by the EIIb and HPr. It is often found fused to the HTH in the LevR-subfamily with the NtrC-type AAA+ ATPases</td>
</tr>
<tr>
<td>PTSIIA</td>
<td>α + β</td>
<td>–</td>
<td>A conserved domain of the PTS system that receives the phosphate in the phosphor-relay and regulates the sugar permeases</td>
</tr>
<tr>
<td>PTSIIIB</td>
<td>α + β PDB 1BLE</td>
<td>–</td>
<td>A conserved domain of the PTS system that receives the phosphate in the phospho-relay and phosphorylates the PRD domain</td>
</tr>
<tr>
<td><strong>SMBDs typically associated with one-component systems</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ACT (aspartokinase, chorismate mutase, TyrA domain)</td>
<td>α + β sandwich [RRM-like fold], PDB 1PSD</td>
<td>0.19N</td>
<td>Binds various small molecule ligands such as amino acids and purine derivatives</td>
</tr>
<tr>
<td>Lrp-C</td>
<td>α + β sandwich [RRM-like fold]</td>
<td>15.55N</td>
<td>This domain is exclusively found at the C-termini of proteins of the Lrp family and is closely related to the ACT fold. It is likely to be an amino-acid-sensing derivative of the ACT fold which is unique to the Lrp proteins. The amino acid-sensing domain exclusively found fused the ArgR-family of HTH domains</td>
</tr>
<tr>
<td>ArgR-C</td>
<td>α + β, DcoH fold, PDB 1XXA</td>
<td>3.35N</td>
<td>A widespread fold with numerous enzymes that typically function as oxidases or oxygenases. Most members of this fold contain 3 conserved residues that play a role in ligand-recognition, as well as catalysis in the enzymatic versions of the fold. Binds carbohydrates, oxalate and amino acids</td>
</tr>
<tr>
<td>DSBH (Double-stranded β-helix domain)</td>
<td>A double-stranded helix of strands, PDB: 2ARC</td>
<td>6.45N; 14.68C</td>
<td>Domain found at the N-terminus of the Boa1-like archaeal HTHs that may be distantly related to the DSBH domain</td>
</tr>
<tr>
<td>Boa1-N</td>
<td>All β</td>
<td>–</td>
<td>This domain shows an N-terminal all β sub-structure that is likely to form a DSBH structure and a C-terminal all-α unit (Fig. 6). A lineage specific expansion is seen in Nostoc sp. (17 proteins). Many versions contain a conserved cysteine which may be the site for ligand-interaction or attachment of a prosthetic group</td>
</tr>
<tr>
<td>AlcR-N</td>
<td>Two distinct sub-structures with all β and all α sub-structures</td>
<td>1.65C</td>
<td>This domain shows an N-terminal all β sub-structure that is likely to form a DSBH structure and a C-terminal all-α unit (Fig. 6). A lineage specific expansion is seen in Nostoc sp. (17 proteins). Many versions contain a conserved cysteine which may be the site for ligand-interaction or attachment of a prosthetic group</td>
</tr>
<tr>
<td>cNMPBD (cNMP-binding domain)</td>
<td>A double-stranded β-barrel domain, PDB: 1RGS</td>
<td>7.65 C</td>
<td>Binds 3′-5′ cyclic nucleotides and typically found fused to the C-terminus of the Crp-like HTH domains</td>
</tr>
<tr>
<td>CBS (Cystathionine beta-synthase)</td>
<td>αβ; in nearly all cases, a dimer, PDB 1ZFJ</td>
<td>2.61N</td>
<td>A ligand-binding domain that probably binds a wide range of ligands which may include purine nucleotides and cyclic nucleotides. It is found fused in different proteins to both wHTH and cl-like tetra-helical HTH domains</td>
</tr>
<tr>
<td>UTRA (UbIC, transcription regulator associated domain)</td>
<td>α + β, PDB 1FW9</td>
<td>13.39N</td>
<td>Versions of this domain appear to accommodate a wide range of ligands that include amino acids, sugars, fatty acids and alkylphosphonate. Always found fused to a HTH domains of the HutC/FarR subfamily of the GntR family. Shares a fold with the bacterial Chorismate Lyases</td>
</tr>
<tr>
<td>Domain Name</td>
<td>Description</td>
<td>PDB</td>
<td>Structure</td>
</tr>
<tr>
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<td>-----------</td>
</tr>
<tr>
<td>DeoR-C</td>
<td>α/β PDB 1LK7</td>
<td>0.48N</td>
<td>This domain shares a common fold with the phosphosugar isomerases such as D-ribose-5-phosphate isomerase.</td>
</tr>
<tr>
<td>DtxR-N</td>
<td>α-helical domain followed by SH3-like barrel. PDB 2DTR</td>
<td>3.90N</td>
<td>This domain is found exclusively in the iron sensors of the DtxR-Fur family of HTH regulators.</td>
</tr>
<tr>
<td>FadR-C</td>
<td>α-helical. PDB 1H9G</td>
<td>23.58 N; 0.35C</td>
<td>A unique seven-helical fold found fused to the subfamily of the GntR family typified by the FadR protein.</td>
</tr>
<tr>
<td>PBP-I</td>
<td>α/β PDB: 1JWL</td>
<td>29.35N; 1.35C</td>
<td>Members of this fold bind diverse small molecule ligands. Usually found in the N-termini of the LacI family</td>
</tr>
<tr>
<td>PBP-II</td>
<td>α/β PDB: 1AL3</td>
<td>87.97N; 0.61C</td>
<td>Members of this fold bind diverse ligands like N-acetylserin, thiosulphate, amino acids, carbohydrate. It is usually found in the C terminus of LysR family.</td>
</tr>
<tr>
<td>T-OB</td>
<td>All β PDB: 1B9M</td>
<td>1.23N</td>
<td>A modified version of the OB fold that typically functions as an obligate dimer. Found in ModE family where it binds molybdate.</td>
</tr>
<tr>
<td>PAS (Per-Arn-Sim domain)</td>
<td>α/β. PAS-like fold PDB: 2PYP</td>
<td>13.61N; 3.48C</td>
<td>A widely utilized superfamilly of ligand-binding domains that bind a range of ligands such as heme, Flavin nucleotides, cinnamate. The ligand-binding domains of the IclR family are divergent versions of the PAS domain.</td>
</tr>
<tr>
<td>GAF (cGMP phosphodiesterase, Adenylate cyclase, FhlA domain)</td>
<td>α/β. PAS-like fold</td>
<td>0.16N; 2.10C</td>
<td>Another widely utilized superfamilly of ligand-binding domains sharing a common fold with the PAS domain. Binds ligands such as cNMPs, tetrapyroles and formate. Often found at the N-terminus of NtrC-like proteins.</td>
</tr>
<tr>
<td>TraR-N</td>
<td>α/β. PAS-like fold. PDB: 1L3L</td>
<td>1.90C</td>
<td>N-terminal domain of Agrobacterium TraR transcription factor. It is always found at the C-terminus of several transcription factors of the LuxR family.</td>
</tr>
<tr>
<td>PocR</td>
<td>Predicted PAS-like fold</td>
<td>0.23C</td>
<td>Domain found fused to the AraC family HTH domains in the 1,2-propanediol-dependent transcription. It also occurs in other contexts fused to diverse signaling domains. Secondary structure predictions suggest that it adopts a PAS-like fold.</td>
</tr>
<tr>
<td>NTF2</td>
<td>α + β</td>
<td>–</td>
<td>A domain found in several enzymes like Steroid isomerases, Scytalone dehydratase and the carotenoid binding protein. Likely to function as a SMBD fused to certain sigma factors.</td>
</tr>
<tr>
<td>GyrI</td>
<td>α + β (contains a duplication of SH2 modules)</td>
<td>4.55N</td>
<td>Frequently found associated with HTH domains of the MerR family.</td>
</tr>
</tbody>
</table>

**Enzymatic domains**

<table>
<thead>
<tr>
<th>Domain Name</th>
<th>Description</th>
<th>PDB</th>
<th>Structure</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>NadR nucleotidy transferase domain (HIGH)</td>
<td>α/β. HUP fold; PDB 1LW7</td>
<td>0.48N</td>
<td>The catalytic domain functions in the adenylation of the nicotinamide mononucleotide in NAD biosynthesis.</td>
<td></td>
</tr>
<tr>
<td>mir6529-C module with metalloprotease-like and metal-chelating domains</td>
<td>Contains two distinct sub-domains</td>
<td>2.16N</td>
<td>A novel conserved module typically found at the C-terminus of HTH domains of the cI-like family. The N-terminal sub-domain possesses the same fold of as the Zn-dependent metalloproteases but many copies of it may be catalytically inactive as they show disruptions of HEXXH signature. The C-terminal sub-domain contains a conserved group of 4 cysteines, which suggests that it chelates metals. Its predicted secondary structure suggests that it may adopt a PAS-like fold.</td>
<td></td>
</tr>
<tr>
<td>Biotin ligase domain</td>
<td>BirM (Fold: Class II aaRS and biotin synthetases) and BirC (Fold: SH3-like barrel) PDB 1HXD</td>
<td>2.65N</td>
<td>The ligase domain activates biotin to form biotinyl-5'-adenylate which acts as an effector for transcriptional repression. Its regular activity is the transfer of biotin moiety to biotin-accepting proteins.</td>
<td></td>
</tr>
<tr>
<td>PLPDE</td>
<td>PDB 1DJU</td>
<td>9.48N</td>
<td>Pyridoxal phosphate-dependent aminotransferases. They are typically found fused to GntR family HTH domains.</td>
<td></td>
</tr>
<tr>
<td>Sugar isomerase (SIS)</td>
<td>α/β PDB 1M3S</td>
<td>2.39N</td>
<td>Usually associated with regulators of polysaccharide metabolism regulons. An enzyme in Porphyrin biosynthesis pathway. Usually found fused to a wHTH domain of the OmpR family.</td>
<td></td>
</tr>
<tr>
<td>Uroporphyrinogen-III synthase</td>
<td>α + β</td>
<td>12.65N</td>
<td>(continued on next page)</td>
<td></td>
</tr>
</tbody>
</table>
Table 1 (continued)

<table>
<thead>
<tr>
<th>Domain</th>
<th>Structure</th>
<th>Placement of HTH*</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>PRTase</td>
<td>α/β</td>
<td>PDB 1STO</td>
<td>0.77N</td>
</tr>
<tr>
<td>Sugar kinase</td>
<td>α/β. RNAse H fold</td>
<td>6.77N</td>
<td>Sugar kinases involved in polysaccharide metabolism are usually found fused to a distinct subfamily of wHTH domains of the MarR family</td>
</tr>
<tr>
<td>β-D-Xylosidase</td>
<td>α/β (TIM barrel) PDB: 1UHVD</td>
<td>0.58N</td>
<td>This enzyme is found fused to the AraC-like HTH domains in regulators of polysaccharide metabolism in low GC Gram positive bacteria</td>
</tr>
<tr>
<td>MJ0056</td>
<td>α + β</td>
<td>0.39N</td>
<td>Found fused to HTH domain in well-conserved subfamily of MarR transcription factors that are restricted to the archaea. Genes encoding these proteins are often found in a conserved gene-neighborhood with enzymes of the riboflavin biosynthesis pathway. The conservation pattern suggests that this domain is also enzymatic and may be a novel diaminohydroxyphosphoribosylaminopyrimidine deaminase, because a conventional enzyme has not been identified in most archaea</td>
</tr>
<tr>
<td>V4R (Vinyl-4 reductase domain)</td>
<td>α + β</td>
<td>–</td>
<td>This domain might bind intermediates of chlorophyll or porphyrin biosynthesis pathways</td>
</tr>
<tr>
<td>Thioesterase domain</td>
<td>α + β</td>
<td>–</td>
<td>A thioesterase domain proto-typed by the 4-hydroxybenzoyl-CoA thioesterase. This domain is found fused to the HTH domains of the GntR family in several proteins from low-GC Gram positive bacteria. The proteins usually also possess CBS domains</td>
</tr>
<tr>
<td>Cyanate lyase</td>
<td>α + β</td>
<td>–</td>
<td>Always found fused to a cI-like HTH domain. The enzyme detoxifies cyanide by combining it with bicarbonate to produce ammonia and carbon-dioxide</td>
</tr>
<tr>
<td>ThiJ_PfpI</td>
<td>α/β PDB: 1O14</td>
<td>3.97C</td>
<td>Domains of this superfamily possess a diverse range of activities, such as protease, catalase, 5′-phosphoribosylformylglycinamidase-L-glutamine amido-ligase, and redox-dependent molecular chaperone activity. Usually they are found fused to the AraC family HTH domains</td>
</tr>
<tr>
<td>LexA Procase domain</td>
<td>All β</td>
<td>6.71N</td>
<td>Serine protease domains of the signal peptidase fold. They are found fused to both cI-like and wHTH domains</td>
</tr>
<tr>
<td>Carbamoyl phosphate synthetase</td>
<td>α/β fold. PDB 1a9x</td>
<td>–</td>
<td>A dyad of HTH domains are found between the carboxy and carbamate phosphorylating units of the large subunit of enzyme</td>
</tr>
<tr>
<td>PQO biosynthesis protein C domain</td>
<td>α helical. PDB 1RCW</td>
<td>–</td>
<td>Iron-binding redox enzyme involved in biosynthesis of Coenzyme Pyrroloquinoline quinone. Fusions to HTH seen only in cyanobacteria</td>
</tr>
<tr>
<td>S-adenosyl-l-methionine-dependent methyltransferase domain</td>
<td>α/β. Rossmann fold</td>
<td>1.55N; 0.26C</td>
<td>There have been multiple independent fusions to methyltransferases with different substrate specificities. These include fusions of the ArsR family (Pseudomonas PA0547) and MerR (Bacillus BC2672) families with methylases of unknown specificities, plant isoflavone O-methyltransferases and restriction methylases</td>
</tr>
<tr>
<td>Methyl-DNA protein methyltransferase</td>
<td>α/β. A truncated RNAse H fold</td>
<td>8.52C</td>
<td>These enzymes transfer alkyl groups from O-6-methylguanine-DNA to themselves. The methyl acceptor is a conserved cysteine in the wing of the wHTH domain</td>
</tr>
<tr>
<td>AlkA</td>
<td>α + β, TBP-like domain followed by helical HhH domain. PDB 1DIZ</td>
<td>0.48N</td>
<td>An α-helical DNA glycosylase involved in DNA repair. The HTH domains in these proteins belong to the AraC family</td>
</tr>
<tr>
<td>Ribonuclease R (RnaseR)</td>
<td>α + β</td>
<td>3.06N</td>
<td>A multi-domain enzyme involved in the processing of tmRNA</td>
</tr>
<tr>
<td>P-Loop NTPase</td>
<td>α/β. P-loop fold</td>
<td>12.10N; 51.97C</td>
<td>Diverse versions of the P-loop fold are fused to the HTH domain. These include the AAA+ superclass members like Cdc6, MCM and DnaA, STAND NTPases in the AfsR and MalT like regulators and GTPases in the case of the SelB and IF-12 proteins</td>
</tr>
<tr>
<td>DOC</td>
<td>Predicted α-helical</td>
<td>1.65C</td>
<td>A metal dependent enzymatic domain predicted to function as a nuclease</td>
</tr>
</tbody>
</table>
architectural diversification involves tandem duplications of HTH domains. Such versions are encountered in the sigma factors, where regions 3 and 4 correspond to a tandem duplication of HTH domains [70]. The Myb/SANT family of HTH domains in eukaryotic chromatin proteins and transcription factors also tend to show multiple tandem copies [71]. In certain cases, such duplications of the HTH domain in the same protein are accompanied by functional diversification of the copies. For example, in the sigma factors, the HTH domain corresponding to the region-3 is involved in contacting the core RNA polymerase complex, while the HTH domain associated with region-4 binds DNA associated with the -35 element of the promoter [70].

3.2. Combinations of HTH with other nucleic acid binding domains and protein–protein interaction domains

HTH domains are occasionally combined with other domains that interact with macromolecules resulting in multivalent polypeptides. Examples of such architectures are the basal transcription factors, MBF1 and TFIIE, of the archaeo-eukaryotic lineage, which combine the HTH with a rubredoxin-like Zn-ribbon domain [30]. The Zn-ribbon could either function as a second potential nucleic acid binding domain or mediate specific protein–protein interactions with the basal transcriptional machinery. Similarly, combinations of the HTH with another four-helical DNA binding-domain are seen in the plasmid encoded KorB proteins [57]. In the eukaryotes, the HTH is fused to the single-strand DNA-binding OB-fold domains replication factor RP-A, while in the tumor-suppressor protein, BRCA2, a basic tri-helical HTH is inserted within one of its three OB-folds [72]. In a class of widespread bacterial proteins, prototyped by YitL, a wHTH domain is fused with RNA-binding S1 domains, suggesting that this group of proteins may perform an as yet uncharacterized role in bacterial RNA metabolism (Fig. 3). The domain architectures suggest that these multi-domain proteins usually function as adaptors that may bridge two macromolecular complexes, such as specific transcription factors and the basal transcriptional machinery, by way of the multiple interaction surfaces provided by their distinct domains. An alternative general function for these proteins, which is not mutually exclusive of the previous one, is to induce conformational changes in nucleic acids by binding them at multiple sites.

3.3. Combinations of the HTH domain with catalytic domains

The HTH domain is frequently combined with a diverse set of catalytic domains, and there are several
general functional trends associated with such combinations. One common association of the HTH domain with catalytic domains represents the utilization of the domain as a substrate-recognition or localization domain. The HTH is observed to be linked to catalytic domains in several proteins involved in DNA replication (e.g. the FtsK–HerA superfamily ATPase domain [73] in bacterial chromosome pumping protein FtsK [74], or the AAA+ ATPase domain in MCMS and Cdc6/ ORC1, which function in archaeal and eukaryotic replication initiation [75]) and repair (e.g. AlkA-type helical DNA glycosylase domain), certain restriction endonucleases (e.g. FokI [45]) and modification methylases (e.g. ScrFIA methylase) (Fig. 3). In course of this survey we noted that the DprA protein (Smf/Dal/CilB), which bears the cysteine that receives the alkyl group from the catalytic tyrosine of these enzymes that is covalently linked to DNA ([63,78] and LA unpublished). Similarly, in the methyl-DNA protein methyltransferase (O6-methylguanine-DNA alkyltransferase), a wHTH domain fused to a C-terminal biotin ligase domain [23].

In related examples, the HTH domain recruits a catalytic domain that may act on proteins, rather than nucleic acids. A striking example of this is the wHTH domain fused to the N-terminus of the Rio family of protein kinases from archaea and eukaryotes [30]. The Rio family of kinases function in 40S ribosomal subunit maturation [80,81], and the wHTH domain recruits the linked protein kinase domain to an rRNA processing protein complex. The LexA protein, the repressor of several bacterial DNA repair genes, represents another variation on this general functional theme. It contains a protease domain of the signal peptidase fold fused to a wHTH domain. The protease domain catalyzes an autocatalytic cleavage in response to a DNA damage signal and triggers dissociation of its wHTH domain from target sequences, thereby allowing transcription of DNA repair genes [82]. Architectures analogous to LexA are also seen in the repressors typified by the heat-response transcription factor HdiR from the Lactococcus lactis [83], where a LexA-like protease domain is fused to a cl-like HTH instead of the wHTH seen in LexA (Fig. 3). This implies that the mechanism of transcription regulation with a proteolytic processing step was innovated independently on at least two occasions in evolution.

The next major architectural theme involving combinations of HTH and enzymatic domains appears to be related to feedback regulation of metabolic pathways. In such combinations, the HTH domain is fused to an enzymatic domain catalyzing a key step in a biosynthetic pathway, and usually regulates the transcription of genes in that pathway. One of the archetypal representatives of this architectural theme is the biotin operon repressor, BirA, which contains an N-terminal HTH domain fused to a C-terminal biotin ligase domain [23]. In the presence of biotin the enzymatic domain synthesizes the co-repressor, and the HTH domain represses the transcription of the biotin biosynthesis genes. Comparative genomics suggests that architectures involving fusions to a range of enzymes from cofactor, nucleotide, amino acid and carbohydrate metabolism are fairly common in archaea and bacteria [30] (Table 1). Some notable fusions include combination of the HTH with nicotinamide mononucleotide adenyl transferase and a P-loop kinase in NadR, with the pyridoxal-phosphate dependent aminotransferase domain (in bacterial HTH proteins of the GntR family), the orotate phosphoribosyltransferase (in archaea), sugar kinases (Rok family in bacteria), purine phosphoribosyltransferase (in archaea) and the threonine synthase (restricted to the genus Pyrococcus) (Fig. 3). Some of these architectures, like BirA are widely distributed in the prokaryotic genomes and appear to be ancient. Others like the fusion of an OmpR family wHTH with the uroporphyrinogen-III synthase are found only in actinobacteria, while yet others like a fusion to the threonine synthase are restricted to a single genus, and are apparently of more recent prove-
nance. This observation suggests that the combinations of HTHs with enzymatic domains have been repeatedly selected for throughout the span of prokaryotic evolution.

Two other specialized classes of domain architectures arise through fusions of the HTH domains with either of two types of P-loop NTPase domains, namely the NtrC-like AAA+ domains [84] and the STAND (signal transduction ATPases with numerous domains) NTPase domain [85]. Proteins containing the NtrC-like AAA+ domains are found only in those bacteria that contain sigma-54, and they bind a distant enhancer element and activate transcription of sigma-54 bound promoters. The AAA+ ATPase domains of these proteins perform an ATP-dependent chaperone-like activity that converts the "closed" sigma-54-containing transcription complexes to an "open" configuration, which is favorable for transcription initiation [84]. The NtrC-like AAA+ domains are fused to at least two different types of HTH domains. The classical versions like NtrC and TyrR are fused to a C-terminal basic tri-helical HTH domain of the Fis family [86]. The second version typified by the Bacillus levanase operon regulator, LevR, instead contains an N-terminal wHTH domain, suggesting that there have been two independent fusions of the HTH domain with NtrC-like AAA+ ATPases (Fig. 3). The STAND P-loop NTPases are, as a rule, large multimodular proteins that appear to catalyze the ATP-dependent assembly of complexes in variety of signaling contexts [85]. They typically contain repetitive superstructure-forming domains, such as the WD and TPR domains, which may serve as surfaces for the assembly of multi-protein complexes [85]. The archetypical members of the architectural class combining the HTH and STAND domains are the E. coli MalT [87], Bacillus GutR [88] and Streptomyces AfsR proteins [89]. A recent analysis of the STAND superfamilly revealed that the HTH domains have been fused to them on several independent occasions [85]. The fusions involving the OmpR family of wHTH domains (e.g. in AfsR) usually link the HTH to the N-terminus of the STAND NTPase domain. In contrast, fusions involving the LuxR family of HTH link it to the C-terminus of the STAND module, with a set of α-helical repeats occurring between these two modules (e.g. GutR and MalT) (Fig. 3). The STAND-domain-containing transcription regulators are likely to integrate multiple signaling inputs via interactions of their STAND and super-structure forming domains, and are particularly prevalent in the developmentally or organizationally more complex bacteria. Another version of the HTH domain, associated with a restriction endonuclease fold and STAND NTPases domains (Fig. 3), is found in the PH-type ATPases that are expanded in Pyrococci. These domains have been predicted to localize the endonuclease domains of these proteins to their target sequences [30].

3.4. Architectures related to two-component, PTS and serine/threonine kinase signaling

The two component phospho-relay system, involving the histidine kinase and the receiver domain, which is phosphorylated on a conserved aspartate, comprise one of the most common signaling systems in bacteria and certain archaea [90,91]. The fusions of the receiver domain with the HTH are typical of transcriptional regulators responding to histidine kinase-dependent signaling. Two of the most common architectures, seen in the majority of bacteria, involve combinations of a single N-terminal receiver domain to either a LuxR-like tetra-helical HTH domain (e.g. UhpA and NarL) or wHTH domain (e.g. OmpR and PhoB, Fig. 4). Less frequent fusions involving HTH domains of the AraC- and the CitB families are seen in certain bacteria. Other than these simple architectures, several more complicated architectures involving multiple receiver domains or even fusions to additional histidine kinase (e.g. B. cereus protein BC3207) and NtrC-like AAA+ ATPase (e.g. E. coli NtrC) domains are also observed (Fig. 3) [90,91].

The PTS sugar-transport systems [92] use a phosphorelay cascade to transfer a phosphate from phosphoenolpyruvate to a histidine on the PTS regulatory domain (PRD), which often co-occurs in the same polypeptide to catalyze the ATP-dependent assembly of complexes in variety of signaling contexts. They typically contain repetitive superstructure-forming domains, such as the WD and TPR domains, which may serve as surfaces for the assembly of multi-protein complexes [85]. The archetypal members of the architectural class combining the HTH and STAND domains are the E. coli MalT [87], Bacillus GutR [88] and Streptomyces AfsR proteins [89]. A recent analysis of the STAND superfamilly revealed that the HTH domains have been fused to them on several independent occasions [85]. The fusions involving the OmpR family of wHTH domains (e.g. in AfsR) usually link the HTH to the N-terminus of the STAND NTPase domain. In contrast, fusions involving the LuxR family of HTH link it to the C-terminus of the STAND module, with a set of α-helical repeats occurring between these two modules (e.g. GutR and MalT) (Fig. 3). The STAND-domain-containing transcription regulators are likely to integrate multiple signaling inputs via interactions of their STAND and super-structure forming domains, and are particularly prevalent in the developmentally or organizationally more complex bacteria. Another version of the HTH domain, associated with a restriction endonuclease fold and STAND NTPases domains (Fig. 3), is found in the PH-type ATPases that are expanded in Pyrococci. These domains have been predicted to localize the endonuclease domains of these proteins to their target sequences [30].

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with the upstream protein kinase in regulating the biogenesis of the mycobacterial cell wall [99]. Taken together, HTH domains fused to the receiver, PRD, 3H and FHA domains represent a distinct class of architectures that are typical of proteins responding to environmental and physiological stimuli downstream of signaling cascades.

3.5. Architectures related to single-component signaling

In contrast to the above-discussed signaling cascades, the single-component systems are defined as those signaling systems in which the transcription regulatory domain and the stimulus sensor domain are combined in a single protein. These architectures, which are functionally analogous to the fusions of the HTHs with the metabolic enzymes, are by far the most prevalent architectural category in prokaryotes (Table 1 and Figs. 3 and 4). In their simplest form they combine a HTH domain with a small molecule binding domain (SMBDs) [97]. More complex architectures may involve multiple SMBDs or even additional domains such as the NtrC-like AAA+. The same SMBDs found in the single component systems may also occasionally be found fused to two-component regulators, where they may supply secondary allosteric inputs (Fig. 3).

The most common SMBDs fused to HTHs in the single component systems are drawn from a relative small set of ancient protein folds (Table 1): (1) The PAS-like fold, with representatives such as the PAS domain, the GAF domain, and the ligand binding domains of the IclR-type transcription factors [66,100]. (2) The periplasmic binding protein types I and II domains, which include the ligand-binding domains of the LysR family [101–103]. (3) The ferredoxin-like fold, which includes the ACT domain and related ligand-sensing domains
of the Lrp-like transcription factors and the classic ferredoxins, which are fused to HTH domains in archaebal and cyanobacterial proteins [104–106]. (4) The double-stranded β-helix (cupin) fold, which contains the AraC-type ligand-binding domains, as well as the cNMP binding domains [97,107]. (5) The CBS domain that occurs as an obligate dyad [108]. (6) The GyrI domain, which contains two copies of the SH2S structural module, appears to be one of the principal ligand-binding domains of the MerR family [109].

Some other SMBDs share a common fold with enzymatic domains, but appear to be catalytically inactive versions that merely bind low-molecular weight substrates. Examples of these are: (1) the UTRA domain, which is found in the HutC/FarR group of GntR family transcription factors and possesses the same fold as chorismate lyase [110] and (2) the DeoR ligand-binding domain, which shares a common α/β fold, which is also present in the enzymes of the phosphosugar isomerase family such as ribose phosphate isomerase [111]. The enormous genomic information has resulted in the availability of proteins from numerous prokaryotes, displaying several new domain architectures. In many of these proteins, uncharacterized globular domains are fused with the HTH and other signaling domains, in architectures analogous to those of known sensory domains. Thus, these analogous architectures enable the prediction of novel sensory domains of one-component systems. By this procedure several new candidate sensory domains, with somewhat lower abundance than the previously described domains, were uncovered (Table 1).

An example of such a domain is suggested by the PocR protein, from Salmonella, with an AraC-like HTH domain, which binds the effector 1,2-propanediol, and regulates the propanediol regulon [112]. It contains a distinct globular N-terminal domain that is also found fused to histidine kinases, chemotaxis receptors, and diguanylate phosphodiesterases of the HD-GYP family in other bacterial proteins (VA and LA unpublished). These domain combinations suggest that it is an evolutionarily mobile, small-molecule-sensing domain, which probably initiates responses through the domains with which it is linked.

We also found a conserved domain in the Salmonella typhi invasion regulator IagA [113], which occurs C-terminal to the OmpR-like wHTH domain (Fig. 3). Additionally, it occurs independently in other bacterial proteins fused to adenyl cyclase and histidine kinase domains (data not shown). Iterative sequence profile searches suggest that this domain shares a common fold with the TolB N-terminal domain [114,115], and is typically found at the N-termini of super-structure forming repeats such as TPR and WD40 repeats (Fig. 3). These architectures, and the interactions of the TolB protein [116], suggest that this domain probably acts in unison with the super-structure forming proteins as a potential sensor for the assembly state of certain multi-protein complexes. Hence, transcription factors like IagA with the TolB-N-related domains might regulate transcription in response to the dynamics of multi-protein complexes.

3.6. Unusual functional adaptations of the HTH domain

Beyond its usual DNA binding role the HTH domain appears to have been exapted for a variety of functions, where it is utilized as a molecular adaptor. For example the permuted version of the wHTH in the N-termini of the methionine aminopeptidases appears to represent an ancient recruitment to a protein–protein interaction function [60]. Several such instances of recruitment of the HTH domain to protein–protein interactions are seen in the eukaryotes. One such example is the PINT domain, which forms the structural scaffold of the pro teaseomal lid, the signalosome and the eukaryotic initiation factor eIF3 [117,118]. It appears to have been derived from a prokaryotic wHTH precursor which secondarily lost its DNA-binding properties. The Snf8 family of proteins in eukaryotes contains two tandem copies of a wHTH domain related to the PINT domain. This family includes the Vps22, Vps25 and Vps36 proteins, which are required for sorting of transmembrane proteins and lipids into the multivesicular-bodies in the eukaryotic vesicular transport system [119–121]. Just as in the case of the PINT domain, the duplicate wHTH domains of the Snf8 family proteins provide a scaffold for the formation of multi-protein ESCRT complexes required for vesicular trafficking. Additionally, the same complex of Snf8 family proteins are also implicated in transcriptional elongation [119], suggesting that they might have been secondarily recruited for a eukaryote-specific role in vesicular transport from an original role in transcriptional regulation. The cyclins and Retinoblastoma are derivatives of the ancestral TFIIB protein which were utilized for specific protein–protein interactions, respectively involved in regulating the eukaryotic cell-cycle controlling kinases and the E2F/DP1 proteins [122]. Likewise, the DEP domain, which is found in several signaling proteins [49], the cullin C-terminal domain (found in cullins, which are the adaptors for the anaphase-promoting ubiquitin ligases) [50], and the C-terminal domain of Esa1-like histone acetylases [123] are other notable examples in which the wHTH domain has been recruited to mediate specific protein–protein interactions in diverse eukaryote-specific signaling contexts. In most of these proteins the HTH domain sticks out as a distinct domain in a complex modular architectural context.

The plant isoflavone O-methyltransferases contain a N-terminal wHTH domain which is related to that in bacterial transcription factors, and appears to function as a dimerization domain [124]. The closest relatives of
these plant methyltransferases are seen in bacteria (e.g. McmR from Streptomyces lavendulae) suggesting that the plant lineage probably acquired these enzymes through lateral transfer from a bacterial source. Hence, it is possible that the wHTH domain in the bacterial precursor originally functioned as a transcription regulator that was fused to the methyltransferase domain (see above for discussion on analogous fusions) but was subsequently reused in the plants in a structural role. A similar case is presented by the carbamoylphosphate synthetase (CPS), which contains a tandem duplication of HTH domains between the carboxyphosphate and carbamoyl phosphate synthetic modules of the enzyme (Fig. 3). These HTH domains are of the simple trihelical versions, like those encountered in bacterial transcription factors such as Fis and LuxR. However, in CPS, rather than binding DNA, they apparently function as protein–protein interaction domains that convert the enzyme to its oligomeric form in the presence of uridine [125].

4. The evolutionary classification of HTH domains

As the HTH is a small domain, which exhibits extreme sequence divergence, reconstruction of its higher-order natural classification is fraught with problems arising from the erosion of evolutionary signal. Nevertheless, the availability of numerous high-resolution structures and extensive sequence information allows us at least to reconstruct the major evolutionary radiations of this domain. This reconstruction is based on three distinct sources of information: (1) Structural features (see above for discussion) help in establishing the relationships at the highest level. (2) Sequence information can be used for clustering based on similarity scores, conventional phylogenetic analysis and cladistic analysis with discrete sequence characters. These sequence-based procedures help in resolving the relationships at a lower level, such as defining the principal sequence families, the relationships within them, and some of the higher-level groupings between sequence families. (3) Phyletic patterns (Fig. 4) help in reconstructing the temporal aspects of the evolutionary history of these domains and also help in constraining the directions of derivation of particular versions from others. The combined scenario gleaned from these directions is presented schematically in Fig. 5 (for details of the combined approach see [85]).

This higher-order evolutionary scheme is characterized by the presence of several basal lineages that retain the primitive basic tri-helical version HTH fold, but have no other shared-derived character that groups them together. These basal lineages are followed by the two great monophyletic lineages, namely the tetrahelical superclass and its derivatives and the wHTH and its derivatives. Beyond these, there are few other highly derived versions whose provenance is hard to establish on account of their extreme sequence and structure divergence. Below, we briefly describe the major evolutionary lineages of the HTH along with their phyletic patterns and functional diversification.

4.1. Lineages of basic tri-helical HTH domains

Several distinct families that retain the primitive simple tri-helical HTH domain are represented in one or more of the major divisions of life. The duplicate HTH domains found in the carbamoyl phosphate synthetase represent a distinctive lineage of simple trihelical domains present in all the 3 super-kingsdoms of life. Phylogenetic trees show that these proteins follow the “standard model topology”, with a distinct archaeo-eukaryotic branch and a bacterial branch [126]. This topology and their phyletic pattern suggest that this lineage most probably, goes back to the LUCA. The HTH domains bearing the catalytic tyrosine of the topoisomerase I family that is found in all the 3 super-kingsdoms and the archaeal topoisomerase VI are also distinctive lineages of tri-helical HTH domains with no specific relationship to any other HTH domains. The phyletic pattern of the former lineage suggests that it was probably present in the common ancestor of the 3 super-kingsdoms [127].

Rbp10 family, which is defined by the eponymous RNA polymerase core subunit [54,128], is universal in both archaea and eukaryotes and appears to have been part of the shared vertical inheritance of the archaeo-eukaryotic lineage. Likewise the sigma factor family [129] is conserved throughout the bacteria but bona fide representatives of this group are absent in the archaea and eukaryotes. Members of this group are characterized by a tandem duplication of the HTH domain, sigma-70, which is the basal transcription factor of the bacteria, is usually present in single copy in all bacterial lineages and shows a noticeable phylogenetic signal suggestive of a largely vertical inheritance since the last common ancestor of all the extant bacteria [129,130]. In contrast other sigma factor subfamilies show evidence for lateral transfer, gene loss and lineage specific expansions (Fig. 4). In particular, the ECF subfamily appears to have been widely expanded in numerous bacterial lineages, especially in those with complex metabolic and developmental capabilities [129]. The lineage specific diversification of the ECF subfamily of sigma appears to have been a major contributor to the evolution of niche-specific adaptations in bacteria by allowing diverse patterns of differential gene expression (see below). The sigma-54 family appears to have been derived independently from the remaining sigma factors and is another distinct lineage of tri-helical HTHs that is sporadically distributed in bacteria.
Fig. 5. Higher order evolutionary relationships of HTH domains. The horizontal lines show temporal epochs corresponding to three major transitions in evolution, the last Universal common ancestor, the diversification of archaea and bacteria and the evolution of the eukaryotes. Solid lines reflect the maximum depth of time to which a particular family can be traced. Broken lines indicate an uncertainty with respect to the exact point of origin of a lineage. Colored circles at the termini of the lines represent broad functional classes: where yellow represents DNA binding, pink represents RNA binding and blue, interaction with proteins. The ellipses encompass groups of lineages from which a new lineage with relatively limited distribution could have potentially emerged. Lineages of archaeal origin are colored blue, those of bacterial origin are colored orange and those present in archaea and bacteria are colored black. Lineages only detected in the eukaryotes are colored green. The yellow triangle reflects the origin of the L11 family of proteins, the blue triangle reflects the origin of the winged HTHs and the red triangle reflects the origin of the tetra-helical version. The phyletic distribution of the lineages are also shown in brackets, where A: Archaea; B: bacteria; E: eukaryotes; proteo: Proteobacteria and Crown: Crown group eukaryotes. The ‘>’ reflects lateral transfer with the arrow head pointing to the potential direction of transfer.
The *Fis family* of basic tri-helical HTH domains is also found solely in the bacteria and typically appears at the C-termini of the NtrC-like AAA+ domains [131]. It appears to have emerged early in bacterial evolution and spread widely via lateral gene transfer along with the spread of sigma factors of the sigma-54 family. The Fis protein itself appears to have been secondarily derived in the proteobacteria through the gene fission of an NtrC-like regulator [131]. Likewise, the trihelical HTH domains of the *Rok family* are restricted to the bacteria and always found in combination with sugar kinase domains of Hsp70/actin fold. The archaeal *Boa family* of tri-helical HTH domains contains a unique all β-strand domain at the N-terminus that is likely to bind its effectors (Table 1, Fig. 4). The *Myb family* is a pan-eukaryotic family of simple tri-helical domains that appears to have diversified into multiple members prior to the diversification of all extant eukaryotic lineages (Fig. 5). Some versions of the Myb family, the SANT subfamily, appear to have been recruited secondarily for protein–protein interactions in the eukaryotic chromatin [132]. In bacteria, the Myb domain is only seen in the RsfA-related pre-spore transcription factors of the low-GC Gram-positive bacteria [133], suggesting that it was acquired relatively late through lateral transfer from the eukaryotes. In addition to the Myb domain, the *homeodomain and POU domain families* [41], which are also basic tri-helical HTH domain, are respectively widespread in the crown-group eukaryotes and metazoans. The HTH domain of the eukaryotic tumor suppressor *BRCA2* [72] is another simple trihelical version that was derived early in eukaryotes. However, it shows no specific relationships to any other HTH domains with a similar structural configuration.

Beyond these classical families there are several trihelical HTH domains associated with diverse transposases and resolvases, such as the gamma–delta resolvase. The exact point of origin of the HTH domains associated with these mobile elements is difficult to ascertain, particularly with several independent lateral transfers to archaea (Fig. 4). The first four of these families are nearly pan-eukaryotic, while the fifth strictly bacterial in their distribution suggesting that these HTH families had probably diverged from each other even in the common ancestor of all bacteria (Fig. 5). The latter two lineages are more limited in their distribution, but are found in most proteobacteria, and low GC Gram positive bacteria (Fig. 4). Within some of these families several distinct lineages, often defined by specific architectural themes can be identified. However, most of these families contain a dominant architectural theme suggesting these might have been the earliest versions of these families. The *MetI-Arc (RHH)* family of transcription factors appears to have been derived from the basic tri-helical bundle [61]. They are most frequently found as transcriptional regulators of the mobile toxin–antitoxin operons [140]. Hence, it is possible that they were originally derived in such toxin–antitoxin systems, through rapid divergence from a conventional HTH. This appears to have happened early in the evolution of one of the prokaryotic lineages, after which they were widely disseminated across the prokaryotes through horizontal mobility.

### 4.2. The tetra-helical HTH superclass and its derivatives

The first major monophyletic clade of HTH domains is defined by the unifying structural feature of the tetra-helical bundle. Sequence similarities help in identifying several major lineages within this group. The *cI-like family*, typified by the phage lambda cI protein, contains representatives from across the 3 super-kingsdoms of life (Fig. 4). Several distinct subfamilies can be recognized within this family. The largest of these is the bacterial repressor subfamily typified by the protein PbsX (Xre) from the *B. subtilis* prophage 168 [141]. Another notable subfamily is the *MBF1 subfamily*, which is nearly universally conserved in the archaeo-eukaryotic lineage functions, and is an adaptor that appears to bridge the specific transcriptional regulators to the basal transcription machinery MBF1 [142].

The next major assemblage within the tetra-helical superclass comprises of 6 major families that are exclusively prokaryotic in their distribution (Figs. 4 and 5). This assemblage includes the *AraC, LuxR, LacI, DnaA, TrpR and TetR families*, which are predominantly bacterial with several independent lateral transfers to archaea (Fig. 4). The first four of these families are nearly pan-bacterial in their distribution suggesting that these HTH families had probably diverged from each other even in the common ancestor of all bacteria (Fig. 5).

The latter two lineages are more limited in their distribution, but are found in most proteobacteria, and low GC Gram positive bacteria (Fig. 4). Within some of these families several distinct lineages, often defined by specific architectural themes can be identified. However, most of these families contain a dominant architectural theme suggesting these might have been the earliest versions of these families. The *AraC family* contains a duplication of the tetra-helical version of the HTH domain [143], and typically occurs fused to a sugar binding domain.
domain of the DSBH fold [97,107] suggesting that they predominantly function as sugar-sensing transcription factors. However, in complex cyanobacteria, like *Nostoc*, the majority of AraC family HTH domains occur fused to a novel sensor domain that is also found in the siderophore alcalgin sensing transcriptional activator, AlcR, from *Bordetella* [144] (Fig. 6 and Table 1). The most common subfamily of the LuxR family contains fusions to the receiver domain, as seen in the case of the *E. coli* protein NarL. In the case of the LacI family the dominant architecture features a fusion of the HTH domain with a small-molecule binding domain of the periplasmic solute-binding protein fold. DnaA is usually found only in a single copy in all bacterial genomes, with the HTH occurring at the C-terminus of the AAA+ domain [145]. The DnaA protein functions in replication initiation, and also as a transcription factor [146]. Additionally, sporadic versions of the tetrahelical HTH superclass are also seen in several phage transposases related to the Mu transposase.

TFIIB, a basal transcription factor in the archaeo-eukaryotic lineage, defines the *TFIIB family*, a derivative of the tetra-helical class. In the archaea there is typically a single version of this family (Fig. 4). However, in the euukaryotes not only did TFIIB undergo duplication, but it also spawned two more divergent families, namely the cyclins and the Rb proteins [122]. Structural comparisons suggest that the euukaryote-specific *Bright domain family* [55,147], which includes DNA-binding proteins involved in chromatin dynamics was also derived from a TFIIB-like precursor prior to the radiation of euukaryotes from their common ancestor. The provenance of the only bacterial relatives of this version, namely the SpoOA protein from endospore-forming bacteria [56] remains unclear.

### 4.3. The wHTH superclass

The second major monophyletic clade of HTH domains are unified by the presence of a striking derived feature, the “wing”. Several major assemblages with varying distributions can be identified within the wHTH superclass. The wHTH superclass includes the majority of prokaryotic transcription factors. Thirteen major families of prokaryotic wHTH domains, namely the *BirA, ArsR, GntR, DtxR–FurR, CitB, LysR, ModE, MarR, PadR, YicD, Rrf2, ScpB* and *HrcA–RuvB* families, are unified by the presence of a characteristic helix after the wing, and comprise the largest monophyletic assemblage within the wHTH superclass (Fig. 5). Of these, representatives of the ArsR, MarR, YicD, GntR, HrcA–RuvB are seen in both archaea and bacteria (Fig. 4) with phylogenetic trees suggesting distinct pan-archaeal and pan-bacterial branches within them (data not shown). This would imply that these families possibly even go back to the LUCA (Fig. 5). The members of these families, barring some striking exceptions (see below) function as transcription factors suggesting that they could have descended from an ancestral protein that had some role in transcriptional regulation. The MarR family has vastly proliferated in the archaea to give rise to several archaeal subfamilies and includes most of the major archaea-specific wHTH transcription factors. In bacteria the RuvB sub-family, which is derived from the ancestral HrcA lineage, appears to have secondarily acquired a role in DNA recombination after a fusion with the AAA+ domain in the ATPase subunit of the Holliday junction resolvase [148]. Another subfamily of the HrcA–RuvB family, S19AE is a small subunit ribosomal protein in the archaeo-eukaryotic lineage, with a potential RNA-binding function (Fig. 5). A similar case is observed in the GntR family, which has vastly proliferated in bacteria giving rise to many of the major bacterial one-component transcription factors. However, in the archaeo-eukaryotic clade it is represented by a single lineage, the small subunit ribosomal protein S25AE (Figs. 4 and 5).

In contrast to the above families, the BirA, ModE, PadR, ScpB and DtxR–Fur families appear to have had a bacterial origin followed by sporadic lateral transfers to the archaea (Fig. 4). The DtxR–Fur family appears to have specialized early on in regulating metal-dependent transcription of genes. The ScpB family is typically represented in most bacteria by a single protein, which is encoded in the same operon as a kleisin and a SMC-type ABC ATPase [149,150]. The ScpB protein contains a tandem duplication of the wHTH with a C-terminal helix after the wing (Fig. 3), and has been shown to regulate the activity of the chromosome organizing SMC ATPases [149,150]. The archaeal representatives of this family appear to have been acquired through a lateral transfer from the cyanobacteria (LMI and LA unpublished). The pan-bacterial families, namely CitB, LysR and Rrf2, are largely absent in the archaea (Fig. 4). Interestingly, a single member of the Rrf2 family (GLP_14_27362_29578) is seen in the early branching euukaryote *Giardia lamblia*. A number of wHTH domains with distinct sequence conservation profiles, and occurring in large multi-domain proteins also belong to this assemblage of wHTH domains with a C-terminal helix after the wing. These include (1) two HTH domains of the topoisoaseme II family, (2) the HTH domain of the enigmatic topoisoaseme V, which is currently found only in *Methanopyrus kandleri* [78], (3) The ESA1 family of euukaryotic histone acetyltransferases and (4) the N-terminal HTH domains of the Rio kinases from archaea and euukaryotes. Two related wHTH domains from *Giardia* (Genbank Gis: 29245940, 29247865) also appear to have been derived from within the above-discussed assemblage of wHTH domains, but their precise affinities are obscured due to rapid sequence divergence.
Fig. 6. A multiple alignment of the AlcR N-terminal module. Multiple sequence alignment the AlcrN domain was constructed using T-Coffee after parsing high-scoring pairs from PSI-BLAST search results. The Jpred secondary structure is shown above the alignment with H representing an α-helix and E representing a β-strand. The 85% consensus shown below the alignment was derived using the following amino acid classes: hydrophobic (h: ALICVMYFW, yellow shading); small (s: ACDGNPSTV, green) and polar (p: CDEHKNQRST, blue). The conserved $C_212$ is shaded red. The limits of the domains are indicated by the residue positions, on each end of the sequence. The sequences are denoted by their gene name followed by the species abbreviation and GenBank Identifier (gi). The species abbreviations are: Ana: Nostoc sp., Bbro: Bordetella bronchiseptica, Bpar: Bordetella parapertussis, Ctet: Clostridium tetani, Ecar: Erwinia carotovora, Gvio: Gloeobacter violaceus, Pae: Pseudomonas aeruginosa, Plum: Photorhabdus luminescens, Psyr: Pseudomonas syringae, Rpal: Rhodopseudomonas palustris, Rsol: Ralstonia solanacearum, Smel: Sinorhizobium meliloti, Spne: Streptococcus pneumoniae, Ssp: Synechocystis sp., Styp: Salmonella typhimurium, Tden: Treponema denticola, Wsuc: Wolinella succinogenes.
The next major monophyletic assemblage of wHTH superclass includes the DeoR, ArgR, LexR, YitL, Lrp-AsnC, ZBD (Z-DNA binding domain), and RNase R families. These families are unified by overall sequence similarity, and a conserved pattern with a conserved glutamine or arginine residue between helix-1 and helix-2 of the HTH domain. Of these the Lrp-AsnC family is widely conserved in both bacteria and archaea (Figs. 4 and 5) and in phylogenetic trees displays distinct branches separating the majority of archaeal and bacterial members. Hence, it is possible that the Lrp-AsnC family goes back to the LUCA. The ArgR and DeoR are predominantly bacterial families, whereas the LexR group is sporadically found, mainly in low GC Gram positive bacteria. The RNase R family is a limited group that is represented by just a single pan-bacterial orthologous lineage in the form of the wHTH domain at the extreme N-terminus of the Rnase R protein (Fig. 3). Its widespread distribution in the bacteria suggests that it emerged early in the evolution of this lineage. The ZBD family is restricted to the crown group eukaryotes and in animals it is fused to the deaminase domain involved in hyper-mutation of the immunoglobulin genes [151,152]. The restricted phyletic pattern of the ZBD family suggests that it may have evolved from one of the prokaryotic families after lateral transfer to the crown group eukaryotes (Fig. 5).

The wHTH domains found in the archaeo-eukaryotic proteins involved in replication initiation, namely the MCM proteins, the CDC6-Orcl proteins (C-terminal to the AAA+ ATPase domains in both these cases) and the RP-A protein comprise the replication initiation family of wHTH domains. Both the MCM and the CDC6 versions appear to have been present right from the base of the archaeo-eukaryotic lineage [30,153,154]. The version associated with RP-A occurs as a stand-alone protein in the archaea, while it appears to have been fused to the single strand DNA binding OB fold domains in the eukaryotes. Also belonging to this family are the C-terminal wHTH domains from the replicative helicase-primase enzymes of various viruses such as P4, plasmid Rep proteins and the eukaryotic RFX-type DNA binding domain seen in transcription factors like the MHC class II transcription factor HRFX1 [75,155,156]. This family might have originally evolved to recognize specific DNA features associated with the replication initiation sites, and recruiting the catalytic activities involved in pre-initiation and initiation to these sites [75]. The RFX domains are specifically related to certain phage helicase-primase wHTH domains belonging to this family. This suggests that the crown group eukaryotes may have acquired the RFX domains from such a viral replication protein and reused them as a transcription regulator (Fig. 5).

The wHTH domains of the archaeal phenylalanyl tRNA synthetase α-subunits, the eukaryotic ribosomal protein S10 and the bacterial selenocysteine-specific elongation factor SelB appear to comprise a family principally associated with translation and RNA metabolism proteins (Figs. 3 and 5). Their phyletic patterns suggest that their recruitment to RNA-specific functions appears to have occurred after the separation of the major superkingdoms of life, though it is possible that a standalone precursor of this family was already present in the LUCA. Another distinct family of wHTH domains with an exclusive single-stranded RNA-binding function is the La domain family [47,48,157]. The La domain has previously only been reported from eukaryotes; however, using sequence profile analysis we show that it is homologous to the N-terminal domain of the NAD-dependent RNA 2’-phosphotransferase [158], which removes the phosphate from the 2’ ends of RNA. In the RNA 2’-phosphotransferase the La domain bears two of the four absolutely conserved catalytic histidines (LA unpublished), suggesting that it is another case of recruitment of the HTH domain for a catalytic role. The RNA 2’-phosphotransferases are highly conserved in the archaeo-eukaryotic and sporadically observed in the bacteria. This suggests that the La family of HTH domains emerged early in the archaeo-eukaryotic lineage and were subsequently laterally transferred to bacteria. In the eukaryotes the non-catalytic versions (the classical La domains) were recruited for binding 3’ poly(U)-rich elements of nascent RNA polymerase III transcripts and translation regulation [47,48]. The fungal protein frequency [157] contains an as yet functionally uncharacterized version of the La domain, which may regulate the circadian clock via RNA metabolism.

There are other distinct families of wHTH transcription factors in prokaryotes with related 2- or 3-stranded wHTH domains, but they do not appear to belong to any of the aforementioned assemblages. These include the LexA, OmpR, and IclR families that appear to be pan-bacterial families, with at best a rare presence in the archaea (Figs. 4 and 5). The classical representatives of the LexA family appear to be involved in regulating responses to DNA damage in diverse bacteria [82]. However, a highly divergent, potential offshoot of the LexA family is seen fused to the C-termini of large DNA helicases (lhr) that are found sporadically in several bacteria (Fig. 3). The eukaryotes display their own families of specific transcription factors, such as the Forkhead-histone H1 [25,26] and E2F-DP1 families [159], and the basal transcription factors, such as the TFIIIE-Rap30)-TFIIC-63K family [160] that show structurally similar wHTH domains, but lack specific sequence relationships with any of the prokaryotic families. Within the TFIIIF-TFIIC-63K family, the RNA polymerase III transcription factor, TFIIC-63K, is conserved throughout eukaryotes, but TFIIIE appears to be restricted to the crown group and the apicomplexans.
The wHTH domains of the TFIIF and TFIIC-63K are functionally similar to those of the Forkhead-histone H1 family suggesting that the latter were probably derived from more widespread family of basal transcription factors [160]. The remaining eukaryotic families appear to be chiefly represented in the eukaryotic crown group, implying that they arose relatively late from pre-existing eukaryotic wHTH domains found in the basal transcription machinery or via rapid divergence from laterally transferred prokaryotic transcription factors. A similar scenario appears to be applicable for the four eukaryotic wHTH families that are not involved in DNA-binding, namely the PINT domain, the SNF8 (with two tandem copies of the wHTH domain), culin and the DEP domain families.

Beyond these multi-member families there are highly conserved lineages of 2- or 3-stranded wHTH domains that are typically found in single orthologous groups of proteins, and cannot be linked to any of the larger assemblages. Such lineages include wHTHs found in the bacterial proteins such as FtsK, DprA and assemblages. Such lineages include wHTHs found in the bacterial proteins, and cannot be linked to any of the larger phylogenetic groups that are typically found in single orthologous groups of proteins.

Distinct from the 2- and the 3-stranded HTH domains are the 4-stranded HTH domains that appear to form a separate monophyletic assemblage within the wHTH clade (Fig. 5). The main prokaryotic family in this assemblage is the Crp family [36] that has a pan-bacterial distribution and sporadic presence in few archaea. Thus, it seems to represent a bacterial innovation that was disseminated to the archaea via lateral transfer. Members of this family are typically fused to a C-terminal cNMP-binding domain, and appear to have specialized early on as cyclic nucleotide dependent regulators. The eukaryotes contain a single major family of this assemblage, the HSF family (heat shock transcription factor), which is present only in the crown group eukaryotes. In the animals this protein appears to have spawned two distinct sub-families that are prototyped by the ETS domain and the IRF domain (interferon regulator factors) [162,163]. Given their relatively restricted presence in eukaryotes, it is possible that they have originated through rapid divergence from laterally transferred prokaryotic versions. A similar scenario could be envisaged for the origin of the orphan initiator binding protein from Trichomonas, which also contains a 4-stranded version of the HTH domain.

The MerR-like assemblage of truncated wHTH domains are derivatives of 3-stranded wHTH domains (Fig. 2). The MerR family proper [37] and the related wHTH domains present in the DNA-binding region of the bacterial phenylalanyl tRNA synthetase β-subunit [164] show a pan-bacterial distribution. In bacteria the MerR family has vastly proliferated into several distinct subfamilies, like the SoxR and CueR subfamilies [37]. However, most other versions show a more restricted distribution. A potential RNA-binding version of this domain is the N-terminal domain of translation initiation factor IF2 from certain bacteria [165]. The remaining versions, observed in the phage lambda excisionase and terminase proteins, the phage Mu-repressor family and the eukaryotic DNA repair protein XP-A and animal transcription factors of the Dachshund family, appear to be DNA binding [166-168]. Eukaryotic Xp-A family is involved in nucleotide excision repair [169], and appears to have been derived at some point in eukaryotic evolution from the functionally similar phage excisionase. The principal diversification of this assemblage appears to have happened early in bacterial evolution resulting in the two ancient families, MerR and the FTRS β-subunit N-terminal domain. Given that regular 3-stranded wHTHs are also found in association with other translation proteins, like the archaeal FTRS-α subunit and SelB, it possible that the prototype of the MerR-like version was derived from such a form through loss of the initial helix.

4.4. Other miscellaneous families of HTH domains

In addition to the above-described major assemblages, there are a few ancient HTH families with uncertain affinities. The chief amongst these are the related ribosomal proteins L11 and S18 (Fig. 2). The former is conserved in all three super-kingsdoms of life and binds RNA in the L11-stalk structure, which appears to go back to the shared ancestral core of the ribosome [170]. S18 appears to have been derived from L11 in the bacteria. Despite its apparent ancient origins L11 appears to be a highly derived version of the HTH. Hence, notwithstanding certain general similarities with the 4-stranded wHTHs, it is more likely that L11 has convergently acquired these features early in evolution.

5. Proteome-wide demographic trends of HTH domains

The availability of a large number and phylectic diversity of complete genome sequences allows robust estimation of the general trends in the proteome-wide distribution of HTH domains. In order to detect the occurrences of HTH domains in proteomes, position-specific score matrices or sequence profiles were constructed for the various distinct families of these domains using seed alignments with diverse representatives. These sequence profiles were then used to iteratively search the target proteomes with the PSI-BLAST program [42]. Alternatively, the alignments were used to generate hidden Markov models, which were similarly used to search the proteomes with the HMMER program [42]. Using a combination of these procedures we determined the total number of proteins containing HTH domains encoded by all completely sequenced organisms that were available at
the time of the analysis. For prokaryotes, the plot of the total number of HTH domains against gene number per genome is best fitted by the power equation of the form \( y = k \times x^c \) (where \( k \) and \( c \) are constants; \( r^2 = 0.76 \); see Fig. 7(a)). The \( r^2 \) of 0.89 for this fit suggests that this tendency is indeed strongly maintained across a wide diversity of genomes. This non-linear scaling of HTH domain numbers with gene number is consistent with recent studies that have suggested that the transcription factor counts follow a power equation with respect to gene number [171]. Given that the HTH domains are the main transcription factors in most of the prokaryotic genomes it is clear that the trend observed for transcription factors is principally a reflection of the distribution of HTH domains (Fig. 7). This distribution function suggests that as gene number increases, a greater than linear number of HTH domain regulators are required per gene.

Examination of major architectural classes of HTH indicates that there is an interesting differential class-wise partitioning of the trends. HTH proteins belong to two-component, serine/threonine kinase and PTS signaling cascades are almost entirely missing in archaea. In the bacteria, where they are abundantly present, their numbers show a linear scaling with respect to gene count per genome (\( r^2 = 0.8 \)) (Fig. 7(c)) [42]. This suggests that each HTH in two-component and related phosphorylation cascades regulates a fixed number of target genes, and as the gene numbers grow larger, the regulators increase in direct proportion to their increase in targets. This is consistent with a model of evolution in which the two-component systems and their target genes undergo duplication with approximately the same probability as the size of a bacterial genome increases in evolution. In contrast to the two-component systems, the one component systems, which chiefly comprise of those HTH domains fused to specific SMDBs or metabolic enzymes, scaled non-linearly with increase in gene count per genome. The best fit for the one component systems was obtained with the power equation of the form \( y = k \times x^c \) (\( r^2 = 0.76 \), Fig. 7(b)). This equation suggests that the HTH domains belonging to the one component system are likely to be a significant contributor to the over-all power equation-type distribution of the HTH domains. This observation implies that as genome size increases a greater than proportional increase in the numbers of one-component transcription factors is required for controlling the newly added genes. This tendency might correlate with the need to regulate specialized groups of their genes, by combining the distinct inputs sensed by the effector-binding domains of multiple sets of one-component transcription factors in the metabolically or organizationally complex bacteria with large genomes.

The counts of sigma factors, too on an average, are positively correlated with gene count per genome (Fig. 7(d)), and scale non-linearly with it (the best fit being provided by a quadratic curve of the form \( y = a \times x^2 + b \times x \); where ‘a’ and ‘b’ are constants; \( r^2 = 0.76 \)). The non-linear scaling of the sigma factors suggests that in the more complex genomes the additional genes are distributed amongst several functionally specialized gene batteries, which are under the regulation of devoted sigma factors responding to specific situations. Interestingly, a few genomes show a significantly greater than expected number of sigma factors (Fig. 7(d)). The most striking example is seen in the case of *Phytoplasma asteris*, which, like other Mycoplasmas, has a highly reduced genome with just over 700 genes [172]. Whereas, the other Mycoplasmas have only a basal sigma-54, *P. asteris* has, in addition to sigma-54, a recent lineage-specific expansion of 11 sigma factors that are related to the *Bacillus* sigma F. Likewise, *Bacteroides thetaiotaomicron* and *Nitrosomonas* show recent lineage-specific expansions of ECF-type sigma factors that have given rise to at least 10 closely related paralogous members in their proteomes. In the case of *P. asteris* there is evidence that the sigma factors may constitute a novel transposon [173]. While this possibility also exists in the case of the other bacteria that show a greater than expected number of sigma factors, it is likely that some of them might have been utilized as transcriptional regulators. This potential link between sigma factors and transposon is consistent with the repeated recruitment of HTH domains from transposases as specific transcriptional regulators.

The genomic data from eukaryotes is still not adequate to discuss general genome-wide trends. However, the available data suggests that eukaryotes display different tendencies from the prokaryotes. The HTH domains of the homeo and Myb proteins are amongst the most prominent DNA-binding domains of transcriptional regulators in the plant and animal lineages. However, their numbers are relatively low in the other eukaryotic lineages. This suggests that the rise in prominence of HTH transcription factors may have been a relatively late phenomenon that occurred on multiple occasions in the eukaryotic crown group. The parasitic apicomplexa, including those forms that have genome sizes comparable to some free-living fungi, have far fewer transcription factors in general [174]. The early branching eukaryote *Giardia lamblia* has at least 13 Myb domains and 2 Bright domains, but apparently no other representatives of the HTH domains found in the eukaryotic specific transcription factors (LA unpublished). Furthermore, the scaling of the total number of transcription factors in eukaryotes with gene counts per genome is not equivalent with what is observed in the prokaryotes. This difference may arise due to two main reasons: (1) the emergence of a complex apparatus for chromatin-structure regulation might have changed the nature of transcriptional control in eukaryotes. (2) Most eukaryotic transcription factors are effectively
down-stream of signaling cascades that communicate from the cell membrane, or the cytoplasm to the nucleus. Hence, there are few equivalents of the genuine prokaryote-type two-component systems in the eukaryotes. Additionally, the eukaryotes might also extensively employ post-transcriptional control mechanisms, involving regulatory RNAs, resulting in a lower dependence on transcriptional regulators \[175,176\]. A combination of these factors might account for the difference in the average number of genes controlled by per transcription factor in eukaryotes and prokaryotes.

6. General considerations on the natural history of the HTH fold and implications for the evolution of transcription

With the exception of the ribosomal protein L11 no bona fide HTH domains with an ancestral RNA-binding role can be confidently traced back to the LUCA. In the HrcA and GntR families the representatives from the archaeo-eukaryotic clade are ribosomal proteins (S19AE and S25AE); whereas all the known bacterial representatives of this family are specific transcription factors (Fig. 5). All other versions of the HTH associated with translation and RNA metabolism, such as those found in La/RNA 2' phosphatase, SelB and ribosomal protein S10E, appear to have been derived after the separation of the archaeo-eukaryotic and bacterial clades. The simplest interpretation of these observations is that the majority of HTH domains associated with RNA metabolism settled into their extant functional niches only after the divergence of the major lineages from LUCA. Hence, excluding L11, it is likely that other HTH domains associated with RNA metabolism in the LUCA performed more generic functions compared to their extant counterparts. The DNA-binding property is strongly preserved across diverse lineages and structural variants of the HTH fold, and involves helix-3, despite the several variations in the details of the interactions of individual versions of the domain. This observation, taken together with the more sporadic
distribution of the versions of the domain associated with translation and RNA metabolism, suggests that the ancestor of most of the extant versions of the HTH, excluding L11, was a DNA-binding protein. Furthermore, the diversification of this domain was potentially associated with the emergence of DNA as genetic material [75,153].

While there are HTH domains in the basal transcription factors of both the bacterial (sigma factors) and archaeo-eukaryotic (TFIIB, TFIIE, and MBF1) lineages, none of these can be considered as being truly orthologous [30,32,33]. In contrast, several families of the HTH domains in specific transcription factors appear to be extensively shared by the bacteria and archaea (Fig. 4). Though several of the prokaryotic families shared by bacteria and archaea can be easily explained as arising from relatively recent lateral transfer between the prokaryotic super-kingsdoms, some others like the Mar R, Ars R, Yct D, Lrp, HrcA and Gnt R families appear to show distinct pan-archaeal and pan-bacterial groups suggesting that they were present in the earliest representatives of each of the super-kingsdoms, hence potentially go back to the LUCA (Fig. 5). Despite the basal transcription machinery shared with the archaea, the specific transcription factors that are traceable to the last eukaryotic common ancestor do not belong to the same families as the specific transcription factors seen in the archaea [30]. These patterns raise a profound conundrum regarding the origin of the phyletic patterns of HTH domains in the basal and specific transcriptional regulators of extant organisms.

Even though several scenarios could in principle explain these patterns, there are only a few parsimonious alternatives that account for the currently available data. Given that both the basal transcription machinery and the domain architecture of the RNA polymerase catalytic subunits are very different in the bacterial and the archaeo-eukaryotic lineages [130], one could extrapolate that the basal transcription factors arose only after the two great lineages had separated from the LUCA. In this situation, the sharing of the specific transcription factors by the archaea and the bacteria could be explained in two possible ways: (1) The LUCA had several specific transcription factors but no basal transcription factors. (2) Alternatively, there were neither specific nor basal transcription factors in the LUCA and both types emerged after the lineages separated. However, multiple very early lateral transfer events resulted in prokaryotic lineages sharing a common set of specific transcription factors. This latter scenario is the consistent with the evidence for extensive lateral transfer between the two prokaryotic super-kingsdoms throughout their evolution [32,177]. At least in the case of the HrcA and GntR families, the striking difference in functions of the extant bacterial and archaeo-eukaryotic representatives suggests that the ancestral versions of these families in LUCA performed a generic nucleic-acid-binding function. They appear to have been secondarily recruited as specific transcription factors only in the bacteria, thereby supporting the second scenario. The basal transcription factors and ribosomal proteins tend to show a stronger signal of vertical inheritance as compared to specific transcription factors which are prone to rampant gene loss and lateral transfer [32,177]. This is not unexpected, given the functional constraints acting on the basal transcription factors, and might confound the reconstruction of the actual evolutionary scenario for the specific transcription factors.

Although the first scenario of basal transcription factors emerging after the specific transcription factors might appear counter-intuitive, detailed analysis of the reconstructed house-keeping functions in the LUCA suggest that it is hardly implausible. Earlier studies on the DNA replication and chromosome partitioning systems suggest that the central enzymes of the DNA replication apparatus appear to have emerged only after the split of the archaeo-eukaryotic and bacterial lineages [32,127,178]. Thus, the DNA replication system resembles the basal transcription apparatus in its origins. More specifically, the absence of an ancestral DNA polymerase and associated replication enzymes in the LUCA suggest that it probably had a system of replication involving reverse transcription [127]. However, simpler but completely functional DNA-dependent RNAPolymerase (DdRP) subunits have been reconstructed for the LUCA [130,179]. Hence, it is possible that the LUCA did not use basal transcription factors and the DdRPs chiefly functioned as enzymes that supplied the RNA template for the replication process involving a reverse transcription step. The precursors of the specific transcription factors might have still functioned under these circumstances, primarily acting as general repressors that regulated the synthesis of the RNA template. The basal transcription factors probably arose only when the genome got organized into multiple tandem operons, each needing its own transcription initiation signal. This suggestion is consistent with the fact that the prokaryotic specific transcription factors can function equally well with both archaeo-eukaryotic- and bacterial-type basal transcription machinery and RNA polymerases, despite the numerous differences [180].

Irrespective of the scenarios, the HTH domains of both the simple tri-helical type (e.g. carbamoylphosphate synthetase and topoisomerase I) and the wHTH type (e.g. topoisomerase II family) are present in proteins that can be confidently traced back to the LUCA. Depending on the scenario, there were at least 6–11 different HTH domains in the cellular genomes, suggesting that the fold had undergone structural and functional differentiation even before the period of the LUCA. Subsequently, in course of prokaryotic evolution, the
HTH domains rapidly expanded in several prokaryotic genomes to rank amongst the folds with highest representation \[32,35\]. The currently available evidence suggests that the common ancestor of extant eukaryotes arose later, via an endosymbiotic event involving an archaeal precursor for the nucleus, translation and secretion apparatus and an \(\alpha\)-proteobacterial precursor for the mitochondrion. However, neither the mitochondrial, nor the nuclear genome, appears to have retained the specific transcription factors might have been inherited from the respective prokaryotic precursors. Instead, the principal pan-eukaryotic HTH transcription factor families, like Bright and MYB domains, are only distantly related to the prokaryotic counterparts. The origin of the eukaryotes saw the emergence of a distinct nuclear compartment, extensive RNA-dependent post-transcriptional gene regulation, a complex chromatin structure and the proliferation of enzymatic complexes involved in chromatin dynamics, such as the Swi2 ATPases, acetylases and deacetylases \[181\]. The compartmentalization of the cell probably rendered the prokaryote-type one-component systems ineffective in the eukaryotes. Furthermore, the regulators of chromatin dynamics, such as the histone deacetylases and associated Swi2 ATPases took up the role of transcriptional repressors \[181\], and probably resulted in the ancestral prokaryote-type repressors becoming superfluous. Hence, the origin of the eukaryotes was probably accompanied by a massive loss of the prokaryote-type transcription factors, along with the innovation through rapid sequence divergence of new versions that suited the eukaryotic milieu. Some of the HTH domains inherited from the ancestral prokaryotic genomes were also reused by the eukaryotes as adaptor modules in signaling systems un-related to DNA-binding or transcription regulation. However, many versions inherited from the archaea appear to have persisted in the basal transcription machinery, where they were indispensable for transcription of the nuclear genome.

Finally, the rise of organizational complexity in plant, animal and fungal lineages went hand-in-hand with the emergence of new specific transcription regulators. Some were drawn from pre-existing HTH families, like the Myb domain, while others like the HSF, Homeo, Pou, Pipsqueak and Paired families arose through rapid divergence from different sources. The HTH domains of transposons provided the source material for some of these domains, whereas other might have diverged rapidly from laterally transferred prokaryotic transcription factors. Thus, on one hand prokaryotes appear to share a sizeable common set of transcription factors, whose phyletic patterns are chiefly governed by the lateral transfers and gene losses acting over and above a basic signal of vertical inheritance. On the other hand, the eukaryotes share only a few unique ancient DNA-binding domains, and their transcription factors have chiefly emerged through multiple lineage-specific expansions.

7. General conclusions

The HTH domain, one of the best-studied of the double-stranded DNA-binding domains, is one of the key protein domains in the transcriptional apparatus of all extant organisms. With the “hindsight” of over two decades of investigations since the discovery of the domain we attempt to provide a synthetic overview of the natural history of the HTH domain from the viewpoint of comparative genomics. Despite the HTH being a rather simple structural scaffold, it is observed to be capable of considerable structural variety and functional versatility, while still preserving a core set of correlated structure-function features. Most HTH domains, despite their structural diversity, participate in a variety of functions that depend on their DNA-binding properties. These include their central role in mediating the substrate interactions of various enzymes that operate on DNA, and their role as both basal and specific transcription regulators. Thus, the HTH domains are the predominant transcription factors in all prokaryotic organisms and the more complex eukaryotes, such as the plants, animals and fungi. Beyond these DNA-binding functions, the HTH domains have been recruited on multiple occasions in a RNA-binding capacity and as mediators of protein–protein interactions. The last universal common ancestor already had anywhere between 6 and 11 distinct versions of the HTH domain, which covered much of the structural diversity, and at least some of the functional diversity seen in the extant versions of the domain. Though several families of specific transcription factors are shared by the two prokaryotic kingdoms and may even go back to their common ancestor, the HTH proteins in the basal transcription factors do not appear to be orthologous. This presents an interesting evolutionary conundrum, whose solution might emerge from new data on alternative transcription and replication systems, like those in viruses and other selfish elements \[75,156\].

The HTH domain occurs frequently in modular proteins, whose domain architectures are often correlated with the general functional properties of the protein. In prokaryotes the dominant domain architecture is the one-component system that combines the HTH with a sensor domain. While many of the sensor domains of commonly known one-component systems have been characterized previously, several others remain structurally and functionally unexplored, and suggest a new direction for exploring the intricacies of biological sensors in prokaryotic systems. The extensive use of divergent HTH domains in protein–protein interactions, especially in eukaryotes, is another area that might
develop further in the future as the actual mechanistic details of such interactions become clearer. In prokaryotic systems the wealth of sequence and structure data might finally allow us to investigate some of the more difficult problems such as, the overall transcriptional regulatory network of organisms, and the details of how target DNA sequence and ligand specificity are achieved by transcriptional regulators. We hope that the overview presented by us will provide a framework for such future investigations.

8. Supplementary material

A complete list of gis of the HTH domains detected in 183 completely sequenced prokaryotic genomes, and alignments of major families will be available by ftp: ftp.ncbi.nih.gov/pub/aravind/.

References


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