

Research

## Interkingdom gene fusions

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### Abstract

**Background:** Genome comparisons have revealed major lateral gene transfer between the three primary kingdoms of life - Bacteria, Archaea, and Eukarya. Another important evolutionary phenomenon involves the evolutionary mobility of protein domains that form versatile multidomain architectures. We were interested in investigating the possibility of a combination of these phenomena, with an invading gene merging with a pre-existing gene in the recipient genome.

**Results:** Complete genomes of fifteen bacteria, four archaea and one eukaryote were searched for interkingdom gene fusions (IKFs); that is, genes coding for proteins that apparently consist of domains originating from different primary kingdoms. Phylogenetic analysis supported 37 cases of IKF, each of which includes a 'native' domain and a horizontally acquired 'alien' domain. IKFs could have evolved via lateral transfer of a gene coding for the alien domain (or a larger protein containing this domain) followed by recombination with a native gene. For several IKFs, this scenario is supported by the presence of a gene coding for a second, stand-alone version of the alien domain in the recipient genome. Among the genomes investigated, the greatest number of IKFs has been detected in *Mycobacterium tuberculosis*, where they are almost always accompanied by a stand-alone alien domain. For most of the IKF cases detected in other genomes, the stand-alone counterpart is missing.

**Conclusions:** The results of comparative genome analysis show that IKF formation is a real, but relatively rare, evolutionary phenomenon. We hypothesize that IKFs are formed primarily via the proposed two-stage mechanism, but other than in the Actinomycetes, in which IKF generation seems to be an active, ongoing process, most of the stand-alone intermediates have been eliminated, perhaps because of functional redundancy.

### Background

Comparative genome analysis has revealed major lateral gene transfer between the three primary kingdoms of life, Bacteria, Archaea, and Eukarya [1-4]. The best recognized form of lateral gene flux is the transfer of numerous genes from mitochondria and chloroplasts to eukaryotic nuclear genomes [5]. Far beyond that, however, the role of lateral gene exchange, along with lineage-specific gene loss, as one of the principal factors of evolution, at least among prokaryotes, is obvious

from the fact that the great majority of conserved families of orthologous genes show a 'patchy' phyletic distribution [6,7]. In many cases, such families are shared by phylogenetically distant species (for example, bacteria and archaea), while they are missing in some of the more closely related species (for example, bacteria from the same lineage). Correlations have been noticed between the preferred routes of gene transfer and the lifestyles of the organisms involved. Thus, massive gene exchange seems to have occurred between

archaeal and bacterial hyperthermophiles [8,9], whereas certain parasitic bacteria, for example, chlamydia and spirochetes, appear to have acquired significantly more eukaryotic genes than free-living bacteria [10-12].

Another evolutionary trend that is predominant in eukaryotes, but is important also in bacteria and archaea, involves the evolutionary mobility of protein domains that combine to form variable multidomain architectures [13-18]. Domain fusion is one of the foundations of most forms of regulation and signal transduction in the cell. Examples include prokaryotic transcriptional regulators, most of which consist of the DNA-binding helix-turn-helix domain fused to a variety of small-molecule-binding domains [19], the two-component signal transduction system that is based on fusions of histidine kinases with sensor domains and of receiver domains with DNA-binding domains [20], and the sugar phosphotransferase (PTS) systems that include complex fusions of several enzymes [21]. In the evolution of eukaryotes, domain fusion takes the form of domain accretion, whereby proteins from complex organisms (such as animals) that are involved in various forms of regulation and signal transduction tend to accrue multiple domains that facilitate the formation of complex networks of interactions [22].

We were interested in exploring the possibility of a meeting between these two major evolutionary phenomena - lateral gene exchange and gene fusion - which would result in the formation of multidomain proteins in which different domains display distinct evolutionary provenance. In particular, we sought to identify fusions between domains originating from different primary kingdoms - Bacteria, Archaea and Eukarya - which we term interkingdom gene (domain) fusions (IKFs), and obtain clues to the pathways of IKF origin through comparative genome analysis. We show that, although IKF in general is a rare phenomenon, one bacterial lineage, the Actinomycetes, displays a significantly increased frequency of such events; we also propose a probable mechanism for IKF formation.

## Results and discussion

To identify IKFs, all protein sequences encoded in the analyzed genomes were compared to the non-redundant protein

database, and those proteins in which distinct parts showed the greatest similarity to homologs from different primary kingdoms were identified (see the Materials and methods section). In most cases, the reported alignments were highly statistically significant, leaving no doubt that true homologs were detected (Table 1). On the few occasions when the database search statistics in themselves were not fully convincing (for example, the OB-fold nucleic acid-binding domain in the *Bacillus subtilis* protein YhcN and the methyltransferase domain in the YabN protein, also from *B. subtilis*), the homologous relationship was validated by detection of the salient sequence motifs known to be involved in the corresponding protein functions (data not shown). Such motif analysis was performed for all analyzed domains in order not only to validate homology, but also to distinguish between active and inactivated forms of enzymes. Figure 1 shows multiple alignments of two domains involved in an IKF, illustrating the conservation of the characteristic functional motifs and the specific similarity between each of the domains of the IKF protein (in this case from *Aquifex aeolicus*) and their archaeal and bacterial homologs, respectively.

In several cases, the chimeric origin of a gene was obvious at a qualitative level because no homolog of the 'alien' domain with comparable sequence similarity was detected in the recipient superkingdom (Table 1, Figure 2a,b). For the rest of the candidate IKFs, phylogenetic tree analysis was performed to corroborate the origin of the invading domain by horizontal transfer; statistically significant grouping of a candidate IKF domain with homologs from the donor superkingdom provides such evidence (Figure 2c,d). The overall number of confirmed IKFs is relatively small - 37 in 21 compared genomes (about 0.1% of the genes) - compared to the total number of likely interkingdom gene transfers. For completely sequenced bacterial genomes this has been conservatively estimated as 1-2% of the genes, with a greater fraction (2-10%) detected in archaea and hyperthermophilic bacteria ([23], and K.S. Makarova, L. Aravind and E.V.K., unpublished observations). Examination of the clusters of orthologous groups (COGs) of proteins from complete genomes [6], in which multidomain proteins are split into the constituent domains if the orthologs of the latter are present as stand-alone forms in some of the genomes, shows that IKFs constitute only a small fraction of all fusions of

### Figure 1

Multiple alignments of two domains comprising an interkingdom domain fusion. Alignments of (a) the PHP-hydrolase domain [4] and (b) the pyruvate formate lyase activating enzyme domain of the IKF protein aq\_2060 from *A. aeolicus*. The sequences of the aq\_2060 domains are placed with the most similar sequences of the corresponding stand-alone enzymes, bacterial ones in the case of PHP-hydrolase and archaeal ones in the case of the pyruvate formate lyase activating enzyme. The phylogenetic trees produced from these alignments are shown in Figure 2c. The numbers in parentheses show the lengths of regions between the aligned blocks that are not shown. The consensus includes amino acid residues and residue classes that are conserved in 75% of the aligned sequences; the residue classes are as follows: h, hydrophobic; l, aliphatic; a, aromatic; s, small; u, tiny; p, polar; b, big; t, residues with high turn-forming propensity. Asterisks show the predicted active site residues; note the replacements in some of the sequences that are predicted to be inactivated versions of the respective enzymes (see text). The alignments were colored using the BOXSHADE program [30]; individual residues conserved in at least 50% of the aligned sequences are in red; residues similar to the conserved ones and groups of conserved similar residues are in blue.

(a)

PHP-hydrolase

Multiple sequence alignment of PHP-hydrolase sequences. The alignment shows conserved regions across various species including Hs1573, Hsp1481, BS\_yadB, Rv1008, YefH, HI0454, Rp682, Bb0194, Tp0979, Tm0667, sl11786, Tp0994, yigW, M3009, Mpl145, Ct594, Cpn0787, yjjv, HI0081, Ybl0556, Mj1582, Mth233, Pml208, Ymr262w, and Consensus757. Conserved motifs are highlighted in bold.

Multiple sequence alignment of Pyruvate formate-lyase activating enzyme sequences. The alignment shows conserved regions across various species including Hs1573, Hsp1481, BS\_yadB, Rv1008, YefH, HI0454, Rp682, Bb0194, Tp0979, Tm0667, sl11786, Tp0994, yigW, M3009, Mpl145, Ct594, Cpn0787, yjjv, HI0081, H11664, Ybl0556, Mj1582, Mth233, Pml208, Ymr262w, and Consensus757. Conserved motifs are highlighted in bold.

(b)

Pyruvate formate-lyase activating enzyme

Multiple sequence alignment of Pyruvate formate-lyase activating enzyme sequences (continued). The alignment shows conserved regions across various species including Hs1573, Mj1632, Mth62, Mpl1643, Af2104, Tm0948, Mpl017, Af1615, Mj1093, Mj1871, Mj0804, Af0918, Ph1966, Hsp0768, jhp0705, aq\_2183, Rv0869c, Rv1309, sl10901, moaA, HI1676, HsaMOC51A, CelF49H6.5, and Consensus757. Conserved motifs are highlighted in bold.

Multiple sequence alignment of Pyruvate formate-lyase activating enzyme sequences (continued). The alignment shows conserved regions across various species including aq\_2060, Mj1632, Mth62, Mpl1643, Af2104, Tm0948, Ph1017, Af1615, Mj1093, Mj1871, Mj0804, Af0918, Ph1966, Hsp0768, jhp0705, aq\_2183, BS\_narA, Rv0869c, Rv1309, sl10901, moaA, HI1676, HsaMOC51A, CelF49H6.5, and Consensus757. Conserved motifs are highlighted in bold.

vertical navigation bar containing the text: content, reviews, reports, deposited research, refereed research, interactions, information

**Table 1****Interkingdom domain fusions and their probable origins**

IKF gene (GI number and gene name) and origin of domains	Best 'native' hit (E-value, amino acid residue range, species)/domain function	Best 'alien' hit (E-value, amino acid residue range, species)/domain function	Protein function	Stand-alone paralog of the alien domain	Comment
<b>Archaea</b>					
<i>Aeropyrum pernix</i> 5106104_ APE2400 Archaeal-bacterial	2621953_Mth 5e-27; 282-445; uncharacterized domain conserved among archaea (homolog of the amino-terminal domain of sialic acid synthase)	2633525_Bs 4e-54; 16-272; hydroxymethyl-pyrimidine phosphate kinase	Hydroxymethyl-pyrimidine phosphate kinase involved in thiamine biosynthesis (additional function?)	None	Pyrococci encode proteins with the same domain organization and closest similarity to <i>A. pernix</i> ; <i>M. jannaschii</i> encodes a protein with the same domain organization but low similarity; Mt encodes a HMP-kinase with moderate similarity
<i>Methanococcus jannaschii</i> 1591138_ MJ0434 Archaeal-bacterial-eukaryotic	2128140_Mj; 1e-19; 2-94; uncharacterized domain	7270033_At; 0.003; 120-222; AlG2-like stress-related protein	Unknown; possible role in stress response	None	The amino-terminal domain is present in several stand-alone copies in <i>M. jannaschii</i> , but otherwise, is seen mostly in bacteria; the possibility of acquisition of a bacterial gene by the <i>Methanococcus</i> lineage is conceivable
<i>Methanobacterium thermoautotrophicum</i> 2621249_ MTH204 Archaeal-eukaryotic/bacterial	5103547_Ap; 1e-34; 137-326; 5-formyl-tetrahydrofolate cyclo-ligase	1651798_Ssp; 0.002; 8-139; uncharacterized membrane-associated domain	Membrane-associated 5-formyl-tetrahydrofolate cyclo-ligase(?); exact function unknown	None	In <i>Ssp</i> , the amino-terminal domain is fused to another uncharacterized domain. An ortholog with conserved domain organization is seen in <i>Mycobacterium</i> , but many other bacteria encode stand-alone versions of this domain, which could be the actual sources of horizontal gene transfer
2621673_ MTH594 Archaeal-bacterial	3256572_Ph; 3e-10; 5-137; inactivated RecA domain	2984130_Aa; 6e-19; 233-390; GTPase	GTPase, possible role in signal transduction	2621855	
2622642_ MTH1523 Archaeal-bacterial	5105992_Ap; 3e-36; 5-226; glucose-1-phosphate thymidyl transferase	2569943_Axy; 2e-05; 226-334; mannose-6-phosphate isomerase	Glucose-1-phosphate thymidyl transferase/ glucose-6-phosphate isomerase	None	
<b>Bacteria</b>					
<i>Aquifex aeolicus</i> 2983622_ aq_1151 Bacterial-archaeal	2633696_Bs; 5e-65; 325-795; c-di-GMP phosphodiesterase	2650176_Af; 0.005; 116-279; PAS/PAC domain	Signal transduction c-di-GMP phosphodiesterase	None	
2984285_ aq_2060 Bacterial-archaeal	586875_Bs; 4e-63 1-252; PHP superfamily hydrolase	3915955_Mj; 3e-09; 270-441; pyruvate formate-lyase activating enzyme (Fe-S cluster oxidoreductase)	Molybdenum cofactor bisynthesis enzyme(?)	None	

**Table 1 (continued)**

IKF gene (GI number and gene name) and origin of domains	Best 'native' hit (E-value, amino acid residue range, species)/domain function	Best 'alien' hit (E-value, amino acid residue range, species)/domain function	Protein function	Stand-alone paralog of the alien domain	Comment
<i>Bacillus subtilis</i> 2632283_yaaH, 1945087_ydhD Bacterial-eukaryotic	4980914_Tm 1e-06 2-92; LysM repeat domain	399377_Rn 2e-11 221-402; chitinase	Chitinase	2635915	<i>B. subtilis</i> encodes two paralogous proteins with the same domain architecture
2633242_yhcR Bacterial-archaeal	645819_Dr; 1e-64; 584-1068; 5'-nucleotidase; 1175987_ ECR100; 2e-09; 377-521; thermonuclease	2622704_Mth; 0.008 151-257; nucleic acid-binding domain (OB-fold)	Nuclease-nucleotidase (probable repair enzyme)	None	
2632325_yabN Bacterial-eukaryotic	4981449_Tm; 2e-62; 223-483; MazG (predicted pyro- phosphatase)	3873806_Ce; 0.003; 7-125; SAM-dependent methyl-transferase	Methyl-transferase/ pyro-phosphatase (metabolic enzyme of an unknown pathway?)	None	Other than in chlamydiae, the SWI domain is seen in eukaryotic chromatin- associated proteins, leading to the suggestion that chlamydial topoisomerase is involved in chromosome condensation
<i>Chlamydomonas reinhardtii</i> 4377077_ CPn0769 Bacterial-eukaryotic	730965_Bs; e-148; 1-727; DNA topoisomerase I	3581917_Sp; 3e-10; 792-866; SWI domain	DNA topoisomerase I, possibly involved in chromatin condensation	7189103	SWI is a typical eukaryotic domain not found in prokaryotes other than chlamydia (the ortholog in <i>Chlamydia trachomatis</i> has the same domain architecture)
<i>Deinococcus radiodurans</i> 6459294_ DR1533 Bacterial-eukaryotic	7248325_Sco; 0.001; 171-265; McrA family endonuclease	6754878_Mm; 9e-28; 4-148; G9a domain (DNA- binding?)	DNase	None	The G9a domain is not detectable in other prokaryotes. In eukaryotes, this domain so far has been found only as part of multidomain nuclear proteins, including transcription factors
<i>Escherichia coli</i> 1787179_ b0947 Bacterial-eukaryotic	94933_Ppu; 3e-10; 287-367; ferredoxin	3747107_Rn; 3e-32; 4-261; uncharacterized domain (thiol oxidoreductase?)	Oxidoreductase	None	The eukaryotic domain is present (as a partial sequence) also in the beta-proteobacterium <i>Vogesella</i> . This domain contains a conserved pair of cysteines, which together with the ferredoxin fusion, may suggest a thiol oxidoreductase activity. Most of the eukaryotic proteins containing this domain appear to be mitochondrial, suggesting the possibility of an alternative evolutionary scenario
1787678_ b1410 Bacterial-archaeal	487713_Sli; 3e-05; 408-522; SAM-dependent methyl-transferase	5459012_Pab; 1e-17; 33-274; lyso-phospholipase	Methyl-transferase/ Lipase (exact function unclear)	None	
1787679_ynbD Archaeal-eukaryotic	1591375_Mj; 4e-04; 50-218; membrane-associated acid phosphatase	7160233_Sp; 1e-06; 346-415; tyrosine phosphatase	Membrane-associated bifunctional phosphatase	None	An unusual case of fusion between an apparently archaeal and a typical eukaryotic domain in a bacterium

comment

reviews

reports

deposited research

refereed research

interactions

information

**Table 1 (continued)**

IKF gene (GI number and gene name) and origin of domains	Best 'native' hit (E-value, amino acid residue range, species)/domain function	Best 'alien' hit (E-value, amino acid residue range, species)/domain function	Protein function	Stand-alone paralog of the alien domain	Comment
1788589_ b2255 Bacterial-eukaryotic	5763950_Sco; 4e-35; 1-259; methionyl-tRNA formyl-transferase	3860247_At; 1e-55; 318-652; dTDP-glucose 4-6- dehydratase	Bifunctional enzyme; exact function unclear	None	
1788938_yfiQ bacterial-Archaeal/ eukaryotic	929735_Nsp; 8e-32; 637-874; acetyl-transferase	2649370_Af; 4e-85; 6-689; acetyl-CoA synthetase	acetyl-CoA synthetase/ acetyl-transferase; exact function unclear	None	
<i>Mycobacterium tuberculosis</i>					
2909507_ Rv2488c, 2791528_Rv1358, 1419061_ Rv1358 Bacterial-eukaryotic	6469244_Sco; 5e-64; 19-603; 4726088_Rer; 2e-12; 818-1073	4151109_Tbr; 6e-04; 6-167; adenylate cyclase	Adenylate cyclase/ ATPase; probable transcription regulator	7476546, 7476738	<i>M. tuberculosis</i> encodes three paralogous proteins that consist of three domains, the eukaryotic- type adenylate cyclase, AP (apoptotic) ATPase and DNA- binding response regulator, and two stand-alone versions of adenylate cyclase, which show the closest similarity to the cyclase domain of the multidomain proteins
1314025_ Rv0886 Bacterial-eukaryotic	120037_Tt; 1e-11; 2-79; ferredoxin	178213_Hs; 4e-65; 93-543; ferredoxin reductase	Ferredoxin/ ferredoxin reductase	2076681	<i>D. radiodurans</i> also encodes the eukaryotic-type ferredoxin reductase, but the ferredoxin fusion is unique to mycobacteria
3261732_ Rv0998 Bacterial-eukaryotic	2661695_Sco; 3e-13; 148-328; acetyl-transferase	279520_Dd; 7e-07; 30-105; cAMP-binding domain	cAMP-dependent acetyl-transferase(?)	4455714 ( <i>M. leprae</i> )	
2326726_ Rv1683 Bacterial-eukaryotic	421331_Cvi; 1e-24; 23-359; poly (3-hydroxy- butyrate) synthase	2645721_Mm; 6e-26; 456-972; very-long-chain acyl-CoA synthetase	Bifunctional enzyme of poly (3-hydroxy-butyrate) synthesis	1929080	
1403447_ Rv2006 Bacterial-eukaryotic	6752338_Sco; 2e-27; 23-240; phosphatase; 6448751_Sco; 0.0; 534-1320; trehalose hydrolase	3892714_At; 8e-27; 264-521; trehalose-6-phosphate phosphatase	Polyfunctional enzyme of trehalose metabolism	2661651	In this protein, the domain of apparent eukaryotic origin is flanked by bacterial domains from both sides
2896788_ Rv2051c Bacterial-eukaryotic	117648_Ec; 1e-16; 94-514; apolipoprotein N-acyltransferase	3073773_Mm; 4e-31; 588-829; dolichol-phosphate- mannose synthase	Polyfunctional enzyme of lipid metabolism	2337823 ( <i>M. leprae</i> ); 6468712 ( <i>Streptomyces coelicolor</i> )	The presence of the stand-alone version of the eukaryotic domain in <i>Streptomyces</i> suggests an ancient horizontal transfer
2791523_ Rv2483c Bacterial-eukaryotic	6225563_Scy; 7e-16; 36-253; phosphoserine phosphatase	1098605_Cnu; 5e-22; 289-492; 1-acyl-sn- glycerol-3-phosphate acyltransferase	Multifunctional enzyme of phospholipid metabolism	None	
2894233_ Rv3323c Bacterial-eukaryotic	2633801_Bs; 3e-19; 89-208; molybdopterin synthase large subunit (MoAe)	4538974_At; 7e-06; 2-82; molybdopterin synthase small subunit (MoAd)	Molybdopterin synthase	2076687	The same domain organization is seen in <i>D. radiodurans</i> , but in this case, both components appear to be of bacterial origin

**Table 1 (continued)**

IKF gene (GI number and gene name) and origin of domains	Best 'native' hit (E-value, amino acid residue range, species)/domain function	Best 'alien' hit (E-value, amino acid residue range, species)/domain function	Protein function	Stand-alone paralog of the alien domain	Comment
2960152_ Rv3728, 7477551_ Rv3239c Bacterial-eukaryotic	4753872_Sco; 1e-35; 56-428; transmembrane efflux protein	466119_Ce; 7e-20; 549-964; cAMP-binding domain- phosphoesterase	cAMP-regulated efflux pump(?)	2501688	<i>M. tuberculosis</i> encodes two strongly similar paralogs with the same domain architecture
2960153_ Rv3729 Bacterial-archaeal	4731342_Sl; 3e-14; 510-776; C5-O-methyl- Transferase (mitomycin biosynthesis)	1591330_Mj; 3e-58; molybdenum cofactor biosynthesis protein MoaA (Fe-S oxidoreductase)	Bifunctional enzyme of molybdenum cofactor biosynthesis	1806159	The amino-terminal domain stand-alone paralog is more similar to archaeal homologs than to the stand-alone paralog, but nevertheless, the latter appears to be of archaeal origin
3261806_ Rv3811 Bacterial-eukaryotic	40487_Cg; 3e-12; 404-494; major secreted protein	7304009_Dm; 2e-12; 198-384; peptidoglycan recognition protein	Secreted protein	7649504 ( <i>S. coelicolor</i> )	The stand-alone version of the eukaryotic domain is present only in <i>Streptomyces</i>
<i>Treponema pallidum</i> 3322964_ TP0667 Bacterial-eukaryotic	7225946_Nm; 9e-04; 10-154; threonyl-tRNA synthetase (TGS and H3H domains)	320868_Sc; 2e-13; 290-488; uridine kinase	Uridine kinase	None	A co-linear ortholog is present in <i>Thermotoga</i>
<i>Thermotoga maritima</i> 4981276_ TM0751 Bacterial-eukaryotic	68516_Bs; 3e-07; 11-200; threonyl-tRNA synthetase (TGS and H3H domains)	3218401_Sp; 2e-11; 288-475; uridine kinase	Uridine kinase	None	A co-linear ortholog is present in <i>Treponema</i>
Eukaryotes <i>Saccharomyces cerevisiae</i>					
536367_ Ybr094w Eukaryotic/ Bacterial-archaeal	586134_Bt; 9e-10; tubulin-tyrosine ligase	7450047_Aa; 8e-09; acid phosphatase (SurE)	Bifunctional signal- transduction protein	5249 ( <i>Yarrowia lipolytica</i> )	SurE homologs are not detectable in eukaryotes other than yeasts
1431219_ YDL141w Eukaryotic- bacterial	577625_Hs; 1e-39 Biotin-[propionyl- CoA-carboxylase(ATP- hydrolysing)] ligase	3328426_Ct 5e-27; biotin protein ligase	Bifunctional biotin- protein ligase	None	An ortholog with an identical domain architecture is present in <i>S. pombe</i>
458922_ YHR206W Eukaryotic-bacterial	477096_Gg; 8e-18; 78-216 heat shock transcription factor domain	1653075_Ssp; 7e-17; 375-503; CheY domain	heat shock transcription factor	None	An ortholog with an identical domain architecture is present in <i>S. pombe</i> (3327019)
486539_ YKR069w Eukaryotic-bacterial	1146165_At; 3e-34; 249-556; uroporphyrin III methylase	2983676_Aa; 1e-04; 22-188; precorrin-2 oxidase	Siroheme synthase	2330809 ( <i>S. pombe</i> )	<i>S. pombe</i> also encodes a co-linear ortholog (3581882); apparent displacement of the bacterial precorrin-2 oxidase by a distinct Rossmann fold domain
1302305_ YNL256w Eukaryotic-bacterial	4938476_At; 5e-65; 324-861 7,8-dihydro-6- hydroxymethylpterin- pyro-phosphokinase+ Dihydro-pterate synthase	3212189_Hi; 5e-05; 62-148; 187-297; dihydro-neopterin aldolase	Multifunctional enzyme of folate biosynthesis	None	Co-linear orthologs in <i>S. pombe</i> (7490442) and <i>Pneumocystis carinii</i> (283062)

Table 1 (continued)

IKF gene (GI number and gene name) and origin of domains	Best 'native' hit (E-value, amino acid residue range, species)/domain function	Best 'alien' hit (E-value, amino acid residue range, species)/domain function	Protein function	Stand-alone paralog of the alien domain	Comment
I419887_ YOL066c Eukaryotic-bacterial	7297709_Dm; 2e-72; 42-408; large ribosomal subunit pseudoU synthase	5918510_Sco; 2e-10; 436-574; pyrimidine deaminase	Bifunctional RNA modification enzyme	2213559 ( <i>S. pombe</i> )	The known bacterial homologs have a two-domain organization; the evolutionary scenario could have included domain rearrangements
I419865_ YOL055c, 2132251_ YPL258c, 2132289_ YPR121w Eukaryotic-bacterial	2462827_At; 1e-39; 22-390; phosphomethyl pyrimidinekinase (thiamine biosynthesis)	1075360_Hi; 6e-24; 342-549; transcriptional activator	Transcriptional regulator of thiamine biosynthesis genes(?)	None	Yeast encodes three strongly similar paralogs with identical domain organization; co-linear orthologs are present in other ascomycetes
I370444_YPL214c Eukaryotic-archaeal/ Bacterial	2746079_Bn; 1e-27; 9-233; thiamin-phosphate pyro-phosphorylase	2648451_Af; 9e-27; 251-531; hydroxyethyl-thiazole kinase	Bifunctional thiamine biosynthesis enzyme	None	Except for the one from <i>A. fulgidus</i> , all highly conserved homologs of the kinase domain of this protein are bacterial; it appears likely that the <i>A. fulgidus</i> gene is the result of horizontal transfer

The following complete genomes were analyzed. Archaea: *Aeropyrum pernix* (Ap); *Archaeoglobus fulgidus* (Af); *Methanococcus jannaschii* (Mj); *Methanobacterium thermoautotrophicum* (Mth); *Pyrococcus horikoshii* (Ph); Bacteria: *Aquifex aeolicus* (Aa); *Borrelia burgdorferi* (Bb); *Bacillus subtilis* (Bs); *Chlamydomonas pneumoniae* (Cp); *Deinococcus radiodurans* (Dr); *Escherichia coli* (Ec); *Haemophilus influenzae* (Hi); *Helicobacter pylori* (Hp); *Mycobacterium tuberculosis* (Mt); *Mycoplasma pneumoniae* (Mp); *Rickettsia prowazekii* (Rp); *Synechocystis* sp. (Ssp); *Thermotoga maritima* (Tm); *Treponema pallidum* (Tp). No IKFs were detected in the genomes that are not shown in the table. Additional species name abbreviations: At, *Arabidopsis thaliana*; Axy, *Acetobacter xylinus*; Bn, *Brassica napus*; Ce, *Caenorhabditis elegans*; Cvi, *Chromatium vinosum*; Gg, *Gallus gallus*; Hs, *Homo sapiens*; Mm, *Mus musculus*; Rn, *Rattus norvegicus*; Sco, *Streptomyces coelicolor*; Sl, *Streptomyces lavendulae*.

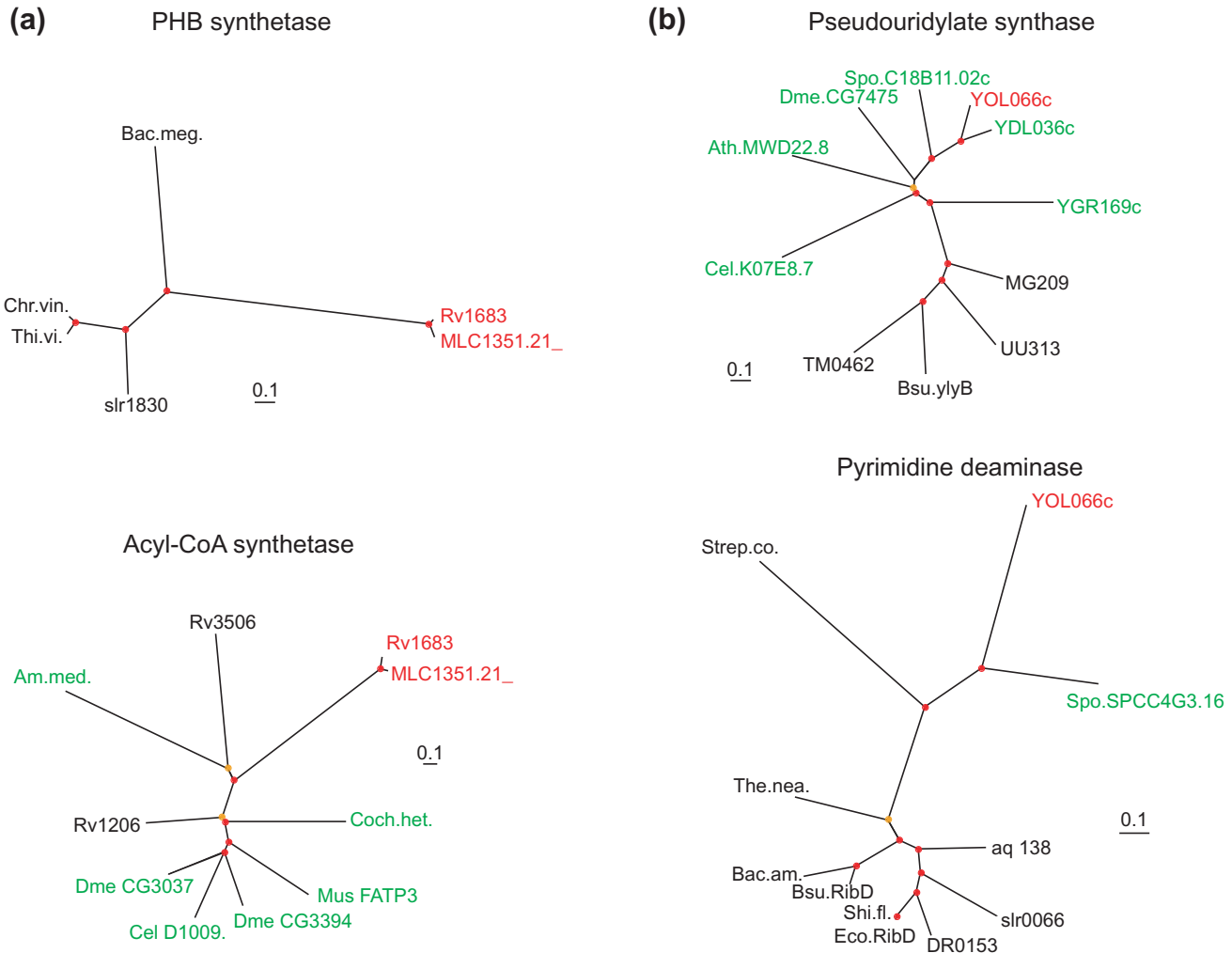
evolutionarily mobile domains (Figure 3). Generally, the small number of identified IKFs compared to the total number of inferred horizontal transfer events and the total number of domain fusions could be compatible with a random model of domain fusion subsequent to lateral gene transfer.

However, the distribution of IKFs among genomes is distinctly non-random, suggesting that such a simple model may be incorrect. Specifically, 12 IKFs were detected in *Mycobacterium tuberculosis* and 10 were found in the yeast *Saccharomyces cerevisiae*, but only a small number or none was identified in each of the other bacterial and archaeal genomes (Figure 2, Table 1). The excess of IKFs in *Mycobacterium* is particularly notable, given that the fraction of genes horizontally transferred from archaea and eukaryotes in the mycobacterial genome is only slightly greater than that in most of the other bacteria, and considerably lower than that in the hyperthermophilic bacteria *Aquifex* and *Thermotoga* (K.S. Makarova, L. Aravind and E.V.K., unpublished observations). Similarly, whereas the overall number of domain fusions in *M. tuberculosis* is greater than in most other bacteria, the difference is insufficient to account for the over-representation of IKFs; furthermore, the cyanobacterium *Synechocystis* sp. has an even greater overall number

of fusions but does not have any detectable IKFs (Figure 3). At present, we cannot provide a defensible biological explanation for the comparatively high frequency of IKF in *Mycobacterium*. It is tempting to interpret this trend in terms of adaptation of this bacterium to its relatively recently occupied parasitic niche, but examination of the individual IKF cases does not offer immediate clues in mycobacterial biology. The yeast IKFs clearly represent relatively recent horizontal transfers distinct from the gene influx from the mitochondria following the establishment of endosymbiosis because, under the protocol of IKF detection used here, only those alien domains were identified that have no counterparts in other eukaryotes.

Most of the IKFs are unique, but *B. subtilis*, *M. tuberculosis* and yeast each also encode families of two to three paralogous IKFs, which apparently have evolved by duplication subsequent to the respective fusion events (Table 1). Strikingly, the same IKF, the three-domain uridine kinase, is shared by *Treponema pallidum* and *Thermotoga maritima* (Table 1). Given that these two bacteria are not specifically related and that *Borrelia burgdorferi*, the second spirochete whose genome has been sequenced, encodes a typical bacterial uridine kinase, the presence of a common IKF in *Treponema* and *Thermotoga* cannot be realistically attributed



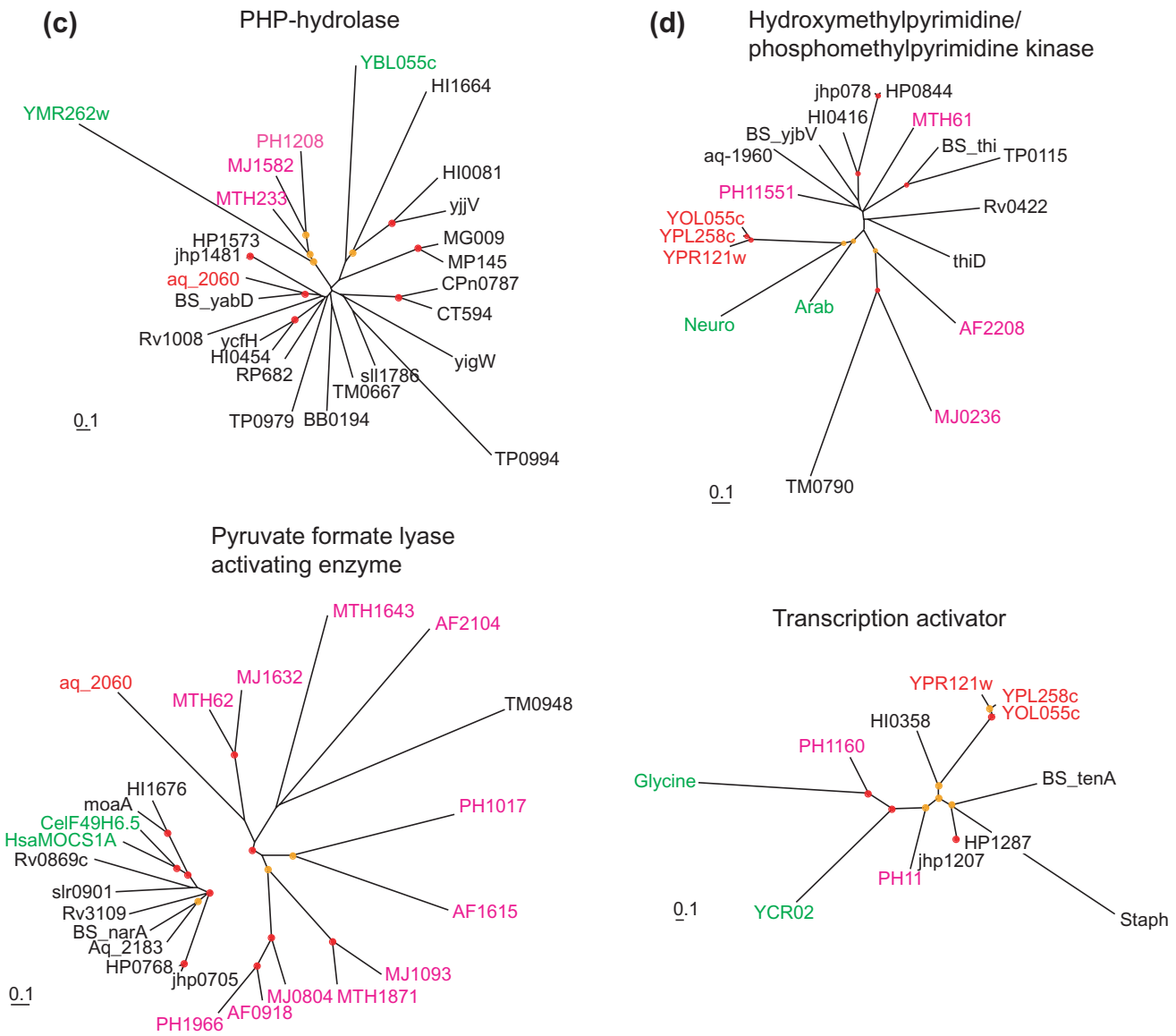


**Figure 2 (and following page)**

Examples of phylogenetic trees supporting the contribution of interkingdom horizontal gene transfer to the emergence of interkingdom domain fusions. The names of proteins from different primary kingdoms are color-coded: black, bacterial; pink, archaeal; green, eukaryotic; the domains involved in the apparent IKF are shown in red. Red circles show nodes with bootstrap support >70%, and yellow circles show nodes with 50-70% support. The bar unit corresponds to 0.1 substitutions per site (10 PAM). **(a)** IKF: Rv1683 (gi| 7476858) from *M. tuberculosis*. Fusion of a bacterial poly(3-hydroxy-butyrates) (PHB) synthase and eukaryotic very long chain acyl-CoA synthetase. Note the absence of eukaryotic homologs in the PHB synthase tree and of bacterial homologs other than the two from *M. leprae* in the acyl-CoA synthetase tree. **(b)** IKF: yeast YOL066c (gi|6324506). Fusion of a eukaryotic pseudouridylate synthetase with a bacterial pyrimidine deaminase. Note the absence of eukaryotic homologs, other than that from *S. pombe*, in the pyrimidine deaminase tree. **(c)** IKF: aq\_2060 (gi|2984285) from *Aquifex aeolicus*. This protein is a fusion of a PHP superfamily hydrolase of apparent bacterial origin and a pyruvate formate-lyase activating enzyme of archaeal origin. **(d)** IKF: yeast YOL055c (gi|1419865), YPL258c (gi|2132251) and YPR121w (gi|2132289) from *S. cerevisiae*. Fusion of a eukaryotic phosphomethylpyrimidine kinase and a bacterial transcriptional activator. Species abbreviations: Bac.meg., *Bacillus megaterium*; Chr.vin., *Chromatium vinosum*; Thi.vi., *Thiocystis violacea*; Am.med., *Amycolatopsis mediterranei*; Coch.het., *Cochliobolus heterostrophus*; Dme, *Drosophila melanogaster*; Cel, *Caenorhabditis elegans*; Mus, *Mus musculus*; Spo, *Schizosaccharomyces pombe*; Ath, *Arabidopsis thaliana*; Strep.co., *Streptomyces coelicolor*; The.nea., *Thermotoga neapolitana*; Bac. am., *Bacillus amyloliquefaciens*; Shi.fl., *Shigella flexneri*; Hsa, *Homo sapiens*.

to vertical inheritance of this gene from a common ancestor. It thus probably reflects horizontal transfer of the gene encoding the three-domain protein subsequent to its emergence in either the spirochetes or the Thermotogales.

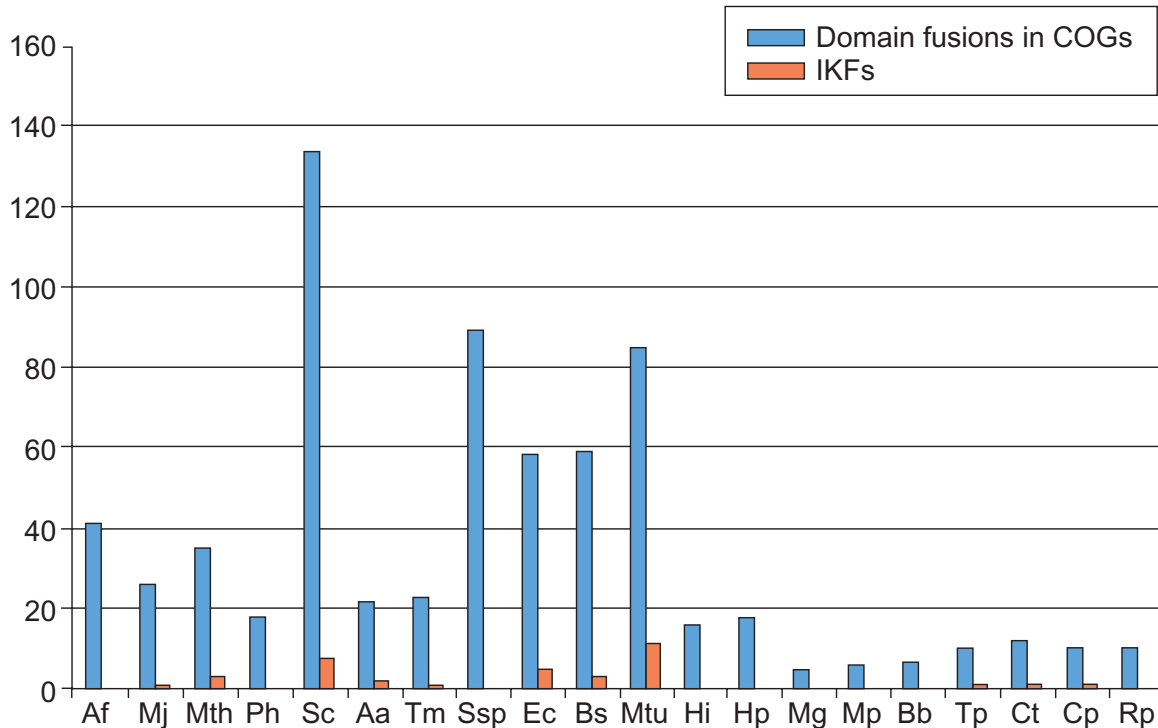
Two evolutionary issues pertaining to IKFs need to be addressed, namely the mechanism(s) of their origin and the selective forces responsible for their preservation. From general considerations, it seems likely that IKFs have



**Figure 2 (continued)**

evolved via a two-step process, which involves lateral transfer of the complete gene coding for the IKF's alien portion, followed by domain fusion. This scenario rests on the assumption that the acquired foreign gene is selectively advantageous, because otherwise it would have been inactivated by mutations before recombination could take place. Under this mechanism, the alien portion of an IKF is likely to be present in the recipient genome also as a stand-alone gene. A clear-cut case of such a duplication of a horizontally transferred domain has been noticed in *Chlamydia*, whose genomes encode the SWI domain, implicated in chromatin condensation, both as a stand-alone protein and as the carboxy-terminal portion of topoisomerase I [10]. Apart

from this case, the IKFs fall into two readily discernible classes, namely those from *Mycobacterium* and all the rest. *M. tuberculosis* (the only complete genome of an actinomyete available) possesses considerably more IKFs than any other bacterial or archaeal species (see above), and typically, the alien portions of these proteins show high level of similarity to the homologs from the donor superkingdom (eukaryotes). Most significantly, there is also, with a single exception, a stand-alone counterpart in the mycobacterial genome; in some cases, such a counterpart is seen only in a closely related species, *M. leprae*, and in one case, it is found in *Streptomyces*, a distantly related actinomyete (Table 1). In the other genomes, the IKFs are generally less similar to



**Figure 3**

Overall numbers of domain fusions estimated using the COGs and interkingdom domain fusions encoded in completely sequenced genomes. The data for estimating the overall number of domain fusions were from the current COG release [6], which does not include several bacterial and archaeal species (for example, *Aeropyrum pernix* and *Deinococcus radiodurans*) that have been analyzed in the present work (Table 1). Accordingly, the data for these genomes are not shown in the figure. Species name abbreviations: Af, *Archaeoglobus fulgidus*; Mj, *Methanococcus jannaschii*; Mth, *Methanobacterium thermoautotrophicum*; Ph, *Pyrococcus horikoshii*; Sc, *Saccharomyces cerevisiae*; Aa, *Aquifex aeolicus*; Tm, *Thermotoga maritima*; Ssp, *Synechocystis* sp.; Ec, *Escherichia coli*; Bs, *Bacillus subtilis*; Mtu, *Mycobacterium tuberculosis*; Hi, *Haemophilus influenzae*; Hp, *Helicobacter pylori*; Mg, *Mycoplasma genitalium*; Mp, *Mycoplasma pneumoniae*; Bb, *Borrelia burgdorferi*; Tp, *Treponema pallidum*; Ct, *Chlamydia trachomatis*; Cp, *Chlamydia pneumoniae*; Rp, *Rickettsia prowazekii*.

the apparent donor and, with a few exceptions, stand-alone versions of the alien domains are missing (Table 1). The hypothesis that seems to be most compatible with these observations is that IKFs indeed evolve via a stand-alone, horizontally transferred intermediate, but in the case of ancient IKFs, these intermediates are typically eliminated during evolution, perhaps because their function becomes redundant with the formation of the IKF. The IKFs identified in actinomycetes appear to result from relatively recent gene fusion events so that the original, stand-alone transferred genes are still present in the genome.

The IKFs include a variety of protein functions. Only some of these are well understood such as, for example, those of the bifunctional nucleotide and coenzyme metabolism enzymes that are particularly abundant in yeast (Table 1). In other cases, the function of an IKF-encoded protein could be predicted only tentatively on the basis of the functions of its constituent domains (Table 1). The selective advantage of the

formation of multidomain proteins, at least as far as enzymes are involved, lies in the possibility of effective coupling of the reactions catalyzed by the different domains [16]; this may be generalized also for functional coordination of non-enzymatic domains. Fusion may result in the addition of a regulatory function to an enzymatic one. For example, it appears most likely that the RNA-binding TGS domain [24] in the uridine kinases of *Treponema pallidum* and *Thermotoga maritima* is involved in autoregulation of translation. The unusual aspect of the IKFs appears to be the compatibility of evolutionarily distant domains.

Examination of the phyletic distribution of the multidomain architectures of IKFs may help in pinpointing the evolutionary stage at which the fusion (but not necessarily the preceding horizontal gene transfer) has occurred. For example, the fusion of the SWI domain with topoisomerase belongs after the radiation of *Chlamydia* from other bacterial lineages, but before the radiation of *Chlamydia pneumoniae* and *Chlamydia*

*trachomatis* (Table 1). The majority of IKFs detected in the yeast *S. cerevisiae* are also present in *Schizosaccharomyces pombe* and/or other ascomycetes (Table 1, and data not shown), but not in any other eukaryotes, and accordingly, they should have evolved at a relatively early stage of fungal evolution, but not before the fungal clade diverged from the rest of the eukaryotic crown group.

Finally, it should be noted that formation of some of the IKFs might have required more complex rearrangements of the contributing proteins than simple domain fusion. Figure 4 shows the domain architectures of proteins that contribute domains to two IKFs. In each case, a simple fusion between genes encoding the respective individual domains is insufficient to explain the emergence of the IKF. For example, the uridine kinase example mentioned above (Figure 4a) should have involved isolation of the TGS-HxxxH domains of threonyl-tRNA synthetase before or concomitantly with their fusion with the uridine kinase. The specific molecular mechanism could have involved selective duplication of the upstream portion of the threonyl-tRNA synthetase gene. Similarly, the sialic acid synthase homologous domain, which is fused to hydroxymethylpyrimidine phosphate kinase in *A. pernix* and pyrococci, appears to have been derived from two-domain proteins that additionally contain a helix-turn-helix DNA-binding domain (Figure 4b). These hypotheses of a complex mechanism of gene fusion involved in the emergence of IKFs are based on a limited sample of sequenced genomes. An alternative possibility is that, before the postulated horizontal transfer event, the recipient domain(s) has been encoded by a stand-alone gene; such genes that do not contain the fused alien domain may yet be discovered in newly sequenced genomes. In fact, a stand-alone version of the sialic acid synthase homologous domain is seen in *Methanobacterium*, although it is considerably less similar to the IKF than the version fused to the HTH domain (Figure 4b).

The identification of IKFs underscores the complexity of the evolutionary process as revealed by comparison of multiple genomes. In and by itself, this phenomenon may not have a unique biological significance, but it reveals the overlap between two major evolutionary trends, horizontal gene transfer and protein domain rearrangement, and shows that domains, rather than entire proteins (genes), should be considered fundamental units of genetic material exchange.

## Materials and methods

Protein sequences encoded in 21 complete genomes of archaea, bacteria and the yeast *Saccharomyces cerevisiae* were extracted from the Genome division of the Entrez retrieval system [25]. Each protein encoded in these genomes was used as the query in a comparison against the non-redundant protein sequence database (National Center for Biotechnology Information, NIH, Bethesda, USA) using the BLASTP program [26]. For each query, the set of local

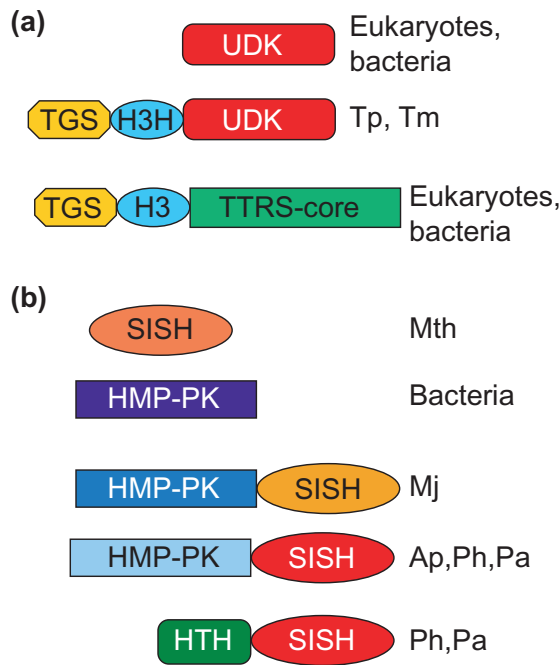
similarities detected by BLASTP was automatically (using a Perl script written for this purpose) screened for putative IKFs, that is situations in which the query did not have full-size homologs outside its immediate taxonomic group (for example, the Proteobacteria for *Escherichia coli*) and in which different regions of the query showed the greatest similarity to proteins from different primary kingdoms. The pseudocode for the script follows:

```

Let H be {h1,h2,... hN} <- hits for the query Q
Lq <- query length
TSq <- query superkingdom
TFq <- query family
for each p < Lq {
  No <- 0
  TSbestL <- ""
  TSbestR <- ""
  for each h in H by decreasing score {
    TSh <- hit superkingdom
    TFh <- hit family
    if(TFh == TFq){ next h }
    Ph(p) <- position of h relative to p
    if(Ph(p) == overlap) {
      if(TSh != TSq){ next h }
      No <- No + 1
      if(No >= maximum allowed No){ exit }
    }elseif(Ph(p) == left) {
      if(TSbestL not empty){ next h }
      TSbestL <- Tsh
    }elseif(Ph(p) == right) {
      if(TSbestR not empty){ next h }
      TSbestR <- Tsh
    }
  }
  if(TSbestL != TSbestR){
    report Q as a candidate to IKF
    exit
  }
}
exit

```

The script itself is available as an additional data file with the online version of this paper. The candidate IKF cases were further examined to detect situations where one or more distinct regions of the query could be classified as 'native' or 'alien' either on the basis of the lack of close homologs from the respective primary kingdom or using phylogenetic analysis. Multiple sequence alignments were generated using the ClustalW program [27], and when necessary, manually corrected to ensure the proper alignment of conserved motifs typical of the respective domains. Phylogenetic trees were constructed using the PROTDIST and FITCH programs of the PHYLIP package [28]. Trees were made separately for each domain of a putative IKF, and its mixed ancestry was considered confirmed if the affinities of the domains with different primary kingdoms were supported by bootstrap values of at



**Figure 4**

Multidomain architectures of interkingdom fusion proteins and their homologs (examples). **(a)** The three-domain uridine kinase; **(b)** the sialic acid synthase homologous domain fused to hydroxymethylpyrimidine phosphate kinase. Domain name abbreviations: TTRS, threonyl-tRNA synthetase; UDK, uridine kinase; TGS and H3H, amino-terminal domains of TTRS; HMP-PK, hydroxymethylpyrimidine phosphate kinase; SISH, sialic acid synthase homologous domain; HTH, helix-turn-helix DNA-binding domain. Different shades represent distinct sequence families of each domain. Species name abbreviations: Tp, *Treponema pallidum*; Tm, *Thermotoga maritima*; Mth, *Methanobacterium thermoautotrophicum*; Mj, *Methanococcus jannaschii*; Ap, *Aeropyrum pernix*; Ph, *Pyrococcus horikoshii*; Pa, *Pyrococcus abyssi*.

least 50%. Additional iterative database searches were performed using the PSI-BLAST program [26,29] in order to predict functions of the individual domains of the identified IKFs in cases when these were not immediately clear.

### Additional data

The following additional data are included with the online version of this paper: the Perl script used to screen local similarities for putative IKFs.

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