

Comparative Genomics and Evolution of Genes Encoding Bacterial (p)ppGpp Synthetases/Hydrolases (the Rel, RelA and SpoT Proteins)

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Abstract

In the gram-negative model organism *Escherichia coli*, the effector molecule of the stringent response, (p)ppGpp, is synthesized by two different enzymes, RelA and SpoT, whereas in the gram-positive model organism *Bacillus subtilis* only one enzyme named Rel is responsible for this activity. Rel and SpoT also possess (p)ppGpp hydrolase activity. BLAST searches were used to identify orthologous genes in databases. The construction and bootstrapping of phylogenetic trees allowed classification of these orthologs. Four groups could be distinguished: With the exception of *Neisseria* and *Bordetella* (β subdivision), the RelA and SpoT groups are exclusively found in the γ subdivision of proteobacteria. Two Rel groups representing the actinobacterial and the *Bacillus/Clostridium* group were also identified. The SpoT proteins are related to the gram positive Rel proteins. RelA proteins carry substitutions in the HD domain (Aravind and Koonin, 1998, TIBS 23: 469-472) responsible for ppGpp degradation. A theory for the evolution of the specialized, paralogous *relA* and *spoT* genes is presented: After gene duplication of an ancestral *rel*-like gene, the *spoT* and *relA* genes evolved from the duplicated genes. The distribution pattern of the paralogous RelA and SpoT proteins supports a new model of linear bacterial evolution (Gupta, 2000, FEMS Microbiol. Rev. 24: 367-402). This model postulates that the γ subdivision of proteobacteria represents the most recently evolved bacterial lineage. However, two paralogous, closely related genes of *Porphyromonas gingivalis* (*Cytophaga-Flavobacterium-Bacteroides* phylum) encoding proteins with functions probably identical to the RelA and SpoT proteins do not fit in this model. Completely sequenced genomes of several obligately parasitic organisms (*Treponema pallidum*, *Chlamydia* species, *Rickettsia prowazekii*) and the obligate aphid symbiont *Buchnera* sp. APS as well as archaea do not contain *rel*-like genes but they are

present in the *Arabidopsis* genome. In crosslinking experiments using different analogs of ppGpp as crosslinking reagents and RNA polymerase preparations of *Escherichia coli*, binding of ppGpp to distinct regions at the C-terminus of the β subunit (the RpoB gene product) and/or at the N-terminus of the β' subunit (the RpoC gene product) was observed previously. RpoB and RpoC sequences of the species which do not possess a *rel* like gene do not exhibit specific insertions or deletions in the ppGpp binding regions.

Introduction

The stringent response is an adaptive global mechanism for control of gene expression in bacterial cells subjected to starvation for amino acids or carbon sources (for review, see Cashel, 1996). Initially discovered forty years ago in the gram-negative model organism *Escherichia coli* (Stent and Brenner, 1961), this response is characterized by profound changes in the transcriptome of starved cells (*e.g.* cessation of rRNA biosynthesis and activation of genes encoding amino acid biosynthesis genes, activation of the stationary sigma factor σ^S (Gentry *et al.*, 1993) and transcription at σ^S -dependent promoters (Kvint *et al.*, 2000). The stringent response is mediated through the synthesis of the effector molecule (p)ppGpp (Cashel and Gallant, 1969).

In *E. coli*, two different proteins are involved in (p)ppGpp synthesis: RelA is bound to ribosomes, senses the amount of uncharged tRNA's and synthesizes (p)ppGpp following amino acid limitation (Haseltine and Block, 1973). The SpoT is a cytosolic protein (Gentry and Cashel, 1995) and functions as (p)ppGpp synthetase after carbon (Hernandez and Bremer, 1991; Murray and Bremer, 1996) and fatty acid (Seyfzadeh *et al.*, 1993) starvation. SpoT is also a (p)ppGpp hydrolase (Hernandez and Bremer, 1991; Murray and Bremer, 1996). The *relA* and *spoT* genes of *E. coli* are related resulting in the hypothesis that these genes evolved by gene duplication (Metzger *et al.*, 1989). Residual (p)ppGpp synthesis in a *relA* mutant is abolished in a *relA/spoT* double mutant (Xiao *et al.*, 1991). *relA/spoT* double mutants show a complex phenotype (morphological alterations; loss in the ability to grow on amino acid-free minimal media) (Xiao *et al.*, 1991; Hernandez and Cashel, 1995). The *relA* and *spoT* genes are listed as paralogous in the COG (cluster of orthologous groups) database (Tatusov *et al.*, 1997; Tatusov *et al.*, 2000) as COG0317.

Several gram-positive bacterial species (*i.e.* *Bacillus subtilis* (Wendrich and Marahiel, 1997); *Corynebacterium glutamicum* (Wehmeier *et al.*, 1998); *Mycobacterium tuberculosis* (Avarbock *et al.*, 1999); *Staphylococcus*

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Table 1. Alphabetical list of species analyzed for enzymes involved in (p)ppGpp metabolism and information about the phylogenetic status. Abbreviations of names are used in the phylogenetic trees and the Tables.

Organism	Phylogenetic position	Abbreviation
<i>Actinobacillus actinomycetemcomitans</i>	Proteobacteria; γ subdivision; Pasteurellaceae	Aac
<i>Arabidopsis thaliana</i>	Eukaryota; Viridiplantae; Embryophyta; Tracheophyta; Magnoliophyta; Rosidae; eurosids II; Brassicales; Brassicaceae	Ath
<i>Aquifex aeolicus</i>	Aquificales; Aquificaceae	Aae
<i>Bacillus anthracis</i>	Firmicutes; Bacillus/Clostridium group; Bacillus/Staphylococcus group; Bacillus cereus group	Ban
<i>Bacillus halodurans</i>	Firmicutes; Bacillus/Clostridium group; Bacillus/Staphylococcus group	Bna
<i>Bacillus stearothermophilus</i>	Firmicutes; Bacillus/Clostridium group; Bacillus/Staphylococcus group	Bst
<i>Bacillus subtilis</i>	Firmicutes; Bacillus/Clostridium group; Bacillus/Staphylococcus group	Bsu
<i>Bordetella bronchiseptica</i>	Proteobacteria; β subdivision; Alcaligenaceae	Bbr
<i>Bordetella pertussis</i>	Proteobacteria; β subdivision; Alcaligenaceae	Bpe
<i>Borrelia burgdorferi</i>	Spirochaetales; Spirochaetaceae	Bbu
<i>Bradyrhizobium japonicum</i>	Proteobacteria; α subdivision; Bradyrhizobium group	Bja
<i>Buchnera</i> sp. APS	Proteobacteria; γ subdivision	Bsp
<i>Campylobacter jejuni</i>	Proteobacteria; ε subdivision; Campylobacter group	Cje
<i>Caulobacter crescentus</i>	Proteobacteria; α subdivision; Caulobacter group	Ccr
<i>Chlamydia trachomatis</i>	Bacteria; Chlamydiales; Chlamydiaceae	Clr
<i>Chlamydia pneumoniae</i>	Bacteria; Chlamydiales; Chlamydiaceae	Cpn
<i>Chlorobium tepidum</i>	Green sulfur bacteria	Cte
<i>Clostridium acetobutylicum</i>	Bacteria; Firmicutes; Bacillus/Clostridium group; Clostridiaceae	Cac
<i>Clostridium difficile</i>	Bacteria; Firmicutes; Bacillus/Clostridium group; Clostridiaceae	Ccd
<i>Corynebacterium diptheriae</i>	Firmicutes; Actinobacteria; Actinobacteridae; Actinomycetales; Corynebacterineae; Corynebacteriaceae	Cdi
<i>Corynebacterium glutamicum</i>	Firmicutes; Actinobacteria; Actinobacteridae; Actinomycetales; Corynebacterineae; Corynebacteriaceae	Cgl
<i>Deinococcus ethenogenes</i>	Green non-sulfur bacteria; Deinococcoides group	Det
<i>Deinococcus radiodurans</i>	Thermus/Deinococcus group; Deinococcales	Dra
<i>Desulfovibrio vulgaris</i>	Proteobacteria; δ subdivision	Dvu
<i>Enterococcus faecalis</i>	Firmicutes; Bacillus/Clostridium group; Enterococcaceae	Efa
<i>Escherichia coli</i>	Proteobacteria; γ subdivision; Enterobacteriaceae	Eco
<i>Geobacter sulfurreducens</i>	Proteobacteria; δ subdivision; Geobacteriaceae	Gsu
<i>Haemophilus ducreyi</i>	Proteobacteria; γ subdivision; Pasteurellaceae	Hdu
<i>Haemophilus influenzae</i>	Proteobacteria; γ subdivision; Pasteurellaceae	Hhi
<i>Helicobacter pylori</i>	Proteobacteria; ε subdivision; Helicobacter group	Hpy
<i>Legionella pneumophila</i>	Proteobacteria; γ subdivision; Legionellaceae	Lpn
<i>Mycobacterium avium</i>	Firmicutes; Actinobacteria; Actinobacteridae; Actinomycetales; Corynebacterineae; Mycobacteriaceae; Mycobacterium avium complex (MAC)	Mav
<i>Mycobacterium bovis</i>	Firmicutes; Actinobacteria; Actinobacteridae; Actinomycetales; Corynebacterineae; Mycobacteriaceae; Mycobacterium tuberculosis complex	Mbo
<i>Mycobacterium leprae</i>	Firmicutes; Actinobacteria; Actinobacteridae; Actinomycetales; Corynebacterineae; Mycobacteriaceae	Mle
<i>Mycobacterium tuberculosis</i>	Firmicutes; Actinobacteria; Actinobacteridae; Actinomycetales; Corynebacterineae; Mycobacteriaceae	Mtu
<i>Mycoplasma genitalium</i>	Firmicutes; Bacillus/Clostridium group; Mollicutes	Mge
<i>Mycoplasma pneumoniae</i>	Firmicutes; Bacillus/Clostridium group; Mollicutes	Mpn
<i>Mycobacterium xanthus</i>	Proteobacteria; δ subdivision; Mycobacteriaceae; Myxococcales; Cystobacterinae	Mxa
<i>Neisseria gonorrhoeae</i>	Proteobacteria; β subdivision; Neisseriaceae	Ngo
<i>Neisseria meningitidis</i>	Proteobacteria; β subdivision; Neisseriaceae	Nme
<i>Pasteurella multocida</i>	Proteobacteria; γ subdivision; Pasteurellaceae	Pmu
<i>Porphyromonas gingivalis</i>	CFB group; Bacteroidaceae	Pgi
<i>Pseudomonas aeruginosa</i>	Proteobacteria; γ subdivision; Pseudomonas group	Paе
<i>Pseudomonas putida</i>	Proteobacteria; γ subdivision; Pseudomonas group	Ppu
<i>Salmonella typhi</i>	Proteobacteria; γ subdivision; Enterobacteriaceae	Sty
<i>Shewanella putrefaciens</i>	Proteobacteria; γ subdivision; Alteromonadaceae	Spu
<i>Spiroplasma citri</i>	Firmicutes; Bacillus/Clostridium group; Mollicutes; Spiroplasmataceae	Sci
<i>Staphylococcus aureus</i>	Firmicutes; Bacillus/Clostridium group; Bacillus/Staphylococcus group	Sau
<i>Streptococcus equi</i>	Firmicutes; Bacillus/Clostridium group; Streptococcaceae	Seq
<i>Streptococcus equisimilis</i>	Firmicutes; Bacillus/Clostridium group; Streptococcaceae	Seq
<i>Streptococcus mutans</i>	Firmicutes; Bacillus/Clostridium group; Streptococcaceae	Smu
<i>Streptococcus pneumoniae</i>	Firmicutes; Bacillus/Clostridium group; Streptococcaceae	Spn
<i>Streptococcus pyogenes</i>	Firmicutes; Bacillus/Clostridium group; Streptococcaceae	Spy
<i>Streptomyces antibioticus</i>	Firmicutes; Actinobacteria; Actinobacteridae; Actinomycetales; Streptomycetales; Streptomycetaceae	San
<i>Streptomyces coelicolor</i>	Firmicutes; Actinobacteria; Actinobacteridae; Actinomycetales; Streptomycetales; Streptomycetaceae	Sco
<i>Synechocystis</i> sp. PCC6803	Cyanobacteria; Chroococcales	Ssp
<i>Thermotoga maritima</i>	Thermotogales	Tma
<i>Thiobacillus ferrooxidans</i>	Proteobacteria; γ subdivision	Tfe
<i>Treponema pallidum</i>	Spirochaetales; Spirochaetaceae	Tpa
<i>Ureaplasma urealyticum</i>	Firmicutes; Bacillus/Clostridium group; Mollicutes	Uur
<i>Vibrio cholerae</i>	Proteobacteria; γ subdivision; Vibrionaceae	Vch
<i>Vibrio</i> sp.	Proteobacteria; γ subdivision; Vibrionaceae	Vsp
<i>Xylella fastidiosa</i>	Proteobacteria; γ subdivision; Xanthomonas group	Xfa
<i>Yersinia pestis</i>	Proteobacteria; γ subdivision; Enterobacteriaceae	Ype

Table 2. RelA orthologs. The sequence of the RelA protein of *E. coli* was used as query sequence in the BLAST search. Individual proteins are sorted according to ascending E-values. For designations of species, see Table 1. The asterisk marks sequence fragments which were not included in the alignments because of their shortness. Designations of proteins: SpoT Pfr: SpoT of *Phlomobacter fragariae* (Foissac *et al.*, 2000); SpoT?: SpoT sequence from an unidentified bacterium (Foissac *et al.*, 2000); CsrS Vsp: CsrS of *Vibrio* sp. (Östling *et al.*, 1995).

Designation	Length	Accession number	E-value
RelA Eco	744	J04039	0.0
RelA Vch	738	Q9S3S3	1.1e-263
RelA Vsp	744	P55133	6.7e-262
RelA Hin	743	P44644	3.5e-256
RelA Pae	747	AAG04323	3.6e-183
RelA Bha	728	BAB04961	1.1e-140
RelA Bsu	734	U86377	2.4e-138
RelA Xfa (XF1316)	718	AE003964	1.2e-136
RelA Nme	769	CAB85211	7.6e-135
Rel Ssp	760	P74007	8e-131
Rel Sau	736	O32419	1.9e-129
Rel Sco	847	X92520	7.5e-128
Rel Cgl	760	O87331	4.1e-127
Rel Mtu	790	Q50638	6.0e-126
Rel Mle	787	Q49640	3.3e-125
Rel Mxa	757	O52177	4.2e-125
Rel Seq	739	Q54089	8.8e-125
Rel San	841	O85709	2.3e-124
Rel Aae	696	O67012	1.2e-123
Rel Dra (DR1838)	787	Q9RTC7	2.6e-111
SpoT Vch (VC2710)	705	AAF95850	3e-108
SpoT Eco	702	P17580	1.7e-101
SpoT Pae	701	AAG08723	1.7e-101
SpoT Nme	725	CAB85138	2.3e-94
SpoT Hin	677	P43811	8.7e-94
Rel Tma (TM0729)	751	Q9WZ18	5.4e-93
Rel Bja	779	Q9RH69	7.2e-84
Rel Bbu	667	O51216	1.1e-80
SpoT Xfa (XF0352)	735	AE003887	8.7e-77
Rel Cje	731	AL139077	5.9e-74
Rel Sci	749	O34098	6.2e-69
RshA Sco (SC4H2.15)	725	O69970	8.6e-67
Rel HpyJ99	776	Q9ZL68	7.6e-66
F1511.23 Ath	715	Q9SYH1	3.3e-59
RSH2 Ath	710	AAF37282	3.4e-52
SpoT Pfr *	388 (fragment)	AAG00076	3.0e-45
T2H3.9 Ath	615	O81418	2.1e-40
RSH1 Ath	883	AAF37281	2.4e-38
Rel Uur	718	AE002125	3.3e-37
Rel Mpn	733	P75386	7.1e-32
Rel Mge	720	P47520	2.0e-29
SpoT ? *	283 (fragment)	AAF73865	1.3e-27
CsrS Vsp *	119 (fragment)	Q56730	1e-20

aureus (Gentry *et al.*, 2000); *Streptococcus equisimilis* (Mechold *et al.*, 1996; Mechold and Malke, 1997); *Streptomyces coelicolor* (Chakraborty *et al.*, 1996; Martinez-Costa *et al.*, 1996, 1998), *Streptomyces antibioticus* (Hoyt and Jones, 1999)) however, possess only one *relA/spoT* paralog which exhibits both (p)ppGpp synthetase and hydrolase activity. In these different species, ppGpp fulfills a variety of regulatory functions: In *B. subtilis*, mutant strains defective in the *relA/spoT* gene exhibit also pleiotropic phenotypes (amino acid auxotrophies (Wendrich and Marahiel (1997)); defects in the expression of several genes controlled by the sporulation sigma factor σ^H and reduced sporulation efficiency (Eymann *et al.*, 2001)). A *rel* mutant of *Mycobacterium tuberculosis* displays slower aerobic growth rate and is less able to survive extended anaerobic incubation (Primm *et al.*, 2000). The *relA/spoT* gene is even

essential in *S. aureus* (Gentry *et al.*, 2000). ppGpp plays a role in antibiotic and pigment production (Martinez-Costa *et al.*, 1996; Chakraborty and Bibb, 1997; Hoyt and Jones, 1999; Hesketh *et al.*, 2001) and morphological differentiation (Chakraborty and Bibb, 1997) in *Streptomyces* species.

In the genomes of the gram-negative species *Myxococcus xanthus* (Harris *et al.*, 1998) and *Porphyromonas gingivalis* (Sen *et al.*, 2000) both *relA* and *spoT* genes seem to be present: Harris *et al.* (1998) cloned and sequenced the *relA* gene of *M. xanthus* and demonstrated that its gene product is essential for fruiting body formation. The other paralogous gene is not yet cloned. Sen *et al.* (2000) performed a search of the "unfinished microbial genome" sequence database of *P. gingivalis* at TIGR, identified two homologues ORF's, and showed that ORF1 encodes RelA of *P. gingivalis*. The other ORF is not yet characterized experimentally.

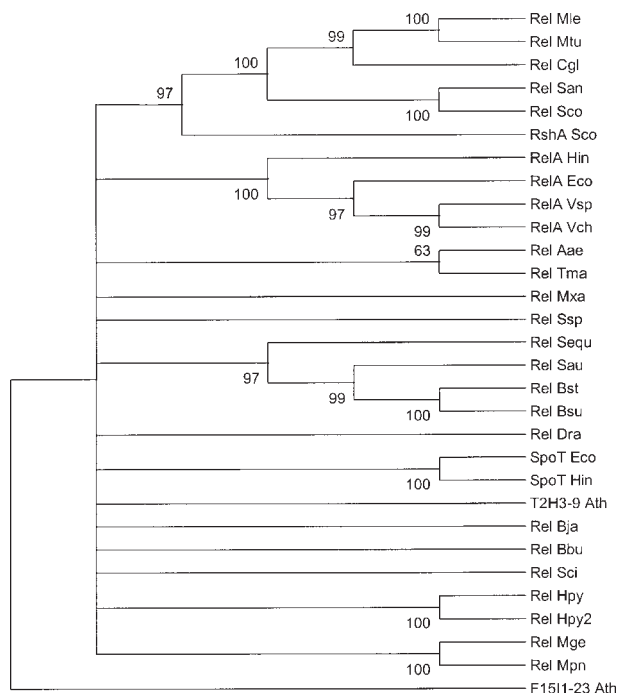


Figure 1. Rectangular cladogram of 30 RelA, SpoT and Rel proteins. Numbers on the branches are percentages of 1,000 bootstrap samples that support the branch; only values >50% are shown. Bootstrapping was performed by the neighbor-joining (NJ) method (Saitou and Nei, 1987) on the basis of the Poisson-corrected amino acid distance (d_{aa}). The sequence F1511-23 of *Arabidopsis* was defined as outgroup.

RelA/SpoT homologs have also been discovered in the model plant *Arabidopsis thaliana* (van der Biezen *et al.*, 2000). These authors used the yeast two-hybrid assay in order to identify proteins that interact with RPP5. This *Arabidopsis* protein which possesses a protein-protein interaction module called "NB-ARC" (van der Biezen and Jones, 1998; Aravind *et al.*, 1999) is part of a signal transduction cascade involved in sensing the fungal pathogen *Peronospora parasitica* (Noel *et al.*, 1999). In this two-hybrid assay, the RSH1 (RelA/SpoT homolog) protein was identified, a predicted plasma membrane-anchored cytoplasmic molecule with significant homology to RelA/SpoT. In a BLAST search two other unlinked *rel*-like genes were detected in the *Arabidopsis* genome, the *RSH2* and *RSH3* genes (van der Biezen *et al.*, 2000). Functional complementation of appropriate *E. coli* and *S. coelicolor* mutants by the *RSH1* gene was also achieved in this study.

Very recently, the cloning and characterization of a gene encoding a *relA/spoT* homolog in *Bacillus stearothermophilus* was reported (Wendrich *et al.*, 2000). The authors include a phylogenetic tree of 32 proteins homologous to RelA/SpoT of *B. subtilis*; van der Biezen *et al.* (2000) also present a phylogenetic tree. The RelA and SpoT proteins of *E. coli* and closely related bacteria form distinct clusters. A cluster of actinobacterial RelA/SpoT sequences and a cluster of sequences representing the *Bacillus/Clostridium* group can be observed. Wendrich *et al.*

(2000) also propose a nomenclature for enzymes involved in (p)ppGpp metabolism: (p)ppGpp synthetase I (represented by RelA of *E. coli*); (p)ppGpp synthetase II (represented by SpoT of *E. coli*) and (p)ppGpp synthetase III (represented by synthetases of gram-positive origin; the name Rel is proposed for this type of enzyme). This nomenclature is adopted in this paper, although biochemical evidence has not yet been reported for several of the proteins named "Rel".

However, Wendrich *et al.* (2000) and van der Biezen *et al.* (2000) do not address the interesting question of evolutionary relationships between these three classes of enzymes to some detail. In this paper, these relationships are investigated using a larger number of RelA, SpoT and Rel sequences for construction of phylogenetic trees (for a list of species analyzed, see Table 1) and by performing statistical tests of tree topologies (*i.e.* bootstrapping (Felsenstein, 1985)). During these studies, we noted that genes encoding Rel-like proteins are absent in completely sequenced genomes of bacteria adopted to intracellular lifestyles. Therefore, we also investigated whether the target regions for binding of (p)ppGpp to the β and β' subunits of RNA polymerase (RNAP) in these species exhibit insertions or deletions (indels).

Results

Four General Groups of ppGpp Synthetase/Hydrolase Proteins

Our initial phylogenetic analysis of the dataset of Wendrich *et al.* (2000) showed a different tree topology than the one reported by the authors (data not shown). The reliability of the branching pattern of these tree data was then tested using bootstrapping (Felsenstein, 1985). Bootstrapping involves a randomization process in a multiple sequence alignment where all residues in a given column of the alignment are preserved in that column. Pseudosamples are created by random selection of sites from the data set with replacement. This is repeated a large number of times (*e.g.* 1000) and a phylogenetic tree is derived for each pseudosample. The percentage of pseudosamples which supports a unique clustering topology provides an estimate of the support for the pattern examined. Two short sequence fragments (*i.e.* CsrS from *Vibrio* sp. (GenBank accession number AAA85717; length: 119 amino acids; Östling *et al.*, 1995) and a fragment derived from a gene of *Pseudomonas denitrificans* (GenBank acc. No. P29941; length 175 aa; Crouzet *et al.*, 1991) which were included in the dataset of Wendrich *et al.* (2000) were excluded from the dataset used for creation of the alignment, because bootstrapping of a dataset containing short sequence fragments is not informative. A phylogenetic tree was constructed by the neighbor-joining (NJ) method (Saitou and Nei, 1987) on the basis of the Poisson-corrected amino acid distance (d_{aa}). Bootstrapping was performed with 1000 replicates (Figure 1). This tree is mainly multifurcating (or "condensed") because many of the interior branches exhibit a bootstrap value lower than 50%. Only branches separating representatives of closely related species as well as the RelA and SpoT proteins are supported by high bootstrap values.

Table 3. SpoT orthologs. The sequence of the SpoT protein of *E. coli* was used as query sequence in the BLAST search. Individual proteins are sorted according to ascending E-values. For designations of species, see Table 1. For explanation of the asterisk, see Figure 2. Designation of the protein: SpoT Pde: SpoT of *Pseudomonas denitrificans* (Crouzet *et al.*, 1991).

Designation	Length	Accession number	E-value
SpoT Eco	702	P17580	0.0
SpoT Vch (VC2710)	705	AAF95850	1.9e-280
SpoT Pae	701	AAG08723	6.6e-207
SpoT Hin	677	P43811	2.3e-204
SpoT Pfr *	388 (fragment)	AAG00076	1.1e-165
SpoT Xfa (XF0352)	735	AE003887	3.7e-165
SpoT Nme	725	CAB85138	1.3e-164
Rel Bha	728	BAB04961	5.8e-148
Rel Bsu	734	U86377	2.5e-147
Rel Seq	739	Q54089	4.2e-143
Rel Ssp	760	P74007	3.8e-142
Rel Sau	736	O32419	8e-140
Rel Mxa	757	O52177	1.9e-138
Rel Aae	696	O67012	1.6e-134
Rel San	841	O85709	1.7e-133
Rel Mtu	790	Q50638	2e-132
Rel Cgl	760	O87331	2.3e-131
Rel Mle	787	Q49640	1.6e-130
Rel Sco	847	X92520	3.8e-127
Rel Dra (DR1838)	787	Q9RTC7	5.5e-126
Rel Tma (TM0729)	751	Q9WZ18	1.3e-125
SpoT ? *	283 (fragment)	AAF73865	9.1e-123
Rel Bja	779	Q9RH69	4.7e-116
RelA Pae	747	AAG04323	9.5e-112
RelA Nme	769	CAB85211	1e-105
RelA Hin	743	P44644	9.4e-103
Rel Bbu	667	O51216	5.1e-102
RelA Eco	744	J04039	5.2e-100
RelA Vsp	744	P55133	7.6e-97
RelA Xfa (XF1316)	718	AE003964	2.9e-94
Rel Cje	731	AL139077	3.4e-91
Rel Sci	749	O34098	1.6e-90
SC4H2.15 Sco	725	O69970	1e-88
RelA Vch (VC249-51)	738	Q9S3S3	2.5e-88
Rel Hpy99	776	Q9ZL68	1.5e-85
RSH1 Ath	883	AAF37281	1.3e-73
F1511.23 Ath	715	Q9SYH1	9.6e-72
Rel Uur	718	AE002125	1.2e-67
RSH2 Ath	710	AAF37282	1.5e-67
T2H3.9 Ath	615	O81418	8.1e-62
Rel Mge	720	P47520	4.1e-56
Rel Mpn	733	P75386	6e-55
CsrS Vsp *	119 (fragment)	Q56730	1.1e-45
SpoT Pde *	175 (fragment)	P29941	2.4e-10

We assumed that a larger set of sequences used as input file for the alignment might result in a more informative tree. Therefore BLAST searches using the RelA (Table 2) and SpoT (Table 3) sequences of *E. coli* and the Rel sequence of *B. subtilis* (Table 4) as query sequences were performed. The Rel sequence of *B. subtilis* was also used to search the "unfinished microbial genomes" database at TIGR (see Table 5). A dataset comprising 80 related sequences was obtained in this way. An alignment of these sequences was performed and an unrooted tree was obtained (Figure 2). Several designations have been replaced by numbers in this tree. A clear separation of the RelA and SpoT families on one hand and of two Rel families of gram-positive origin (*Bacillus/Clostridium* group and an actinobacterial group) can be observed. Bootstrapping of this dataset also resulted in a largely condensed tree with internally supported branches (Figure 3). Interestingly, as

also indicated by Wendrich *et al.* (2000) and as depicted in Figure 1, mycoplasmal Rel proteins form a clear outgroup.

It seemed likely that exclusion of certain sequences from the dataset used to produce the alignment might increase the resolution of the condensed part of the tree. In a first attempt, the outgroup sequences of *Arabidopsis* (RSH2 Ath, F1511-23 Ath) and of the *Mycoplasma* (Rel Mpn, Rel Mge, Rel Uur) group as well as the *Arabidopsis* sequences T2H3-9 and RSH1 which are located in the condensed part of the tree were removed from the dataset (total: 73 sequences) subjected to the alignment. The resulting bootstrapped tree did not exhibit increased resolution in the condensed part of the tree (data not shown). In a second attempt, internal sequences of the condensed region of the tree in Figure 3 (*i.e.* Rel Dra, Rel Ssp, Rel Det, Rel Aae, Rel Tma, Rel Mxa, Rel Gsu, Rel

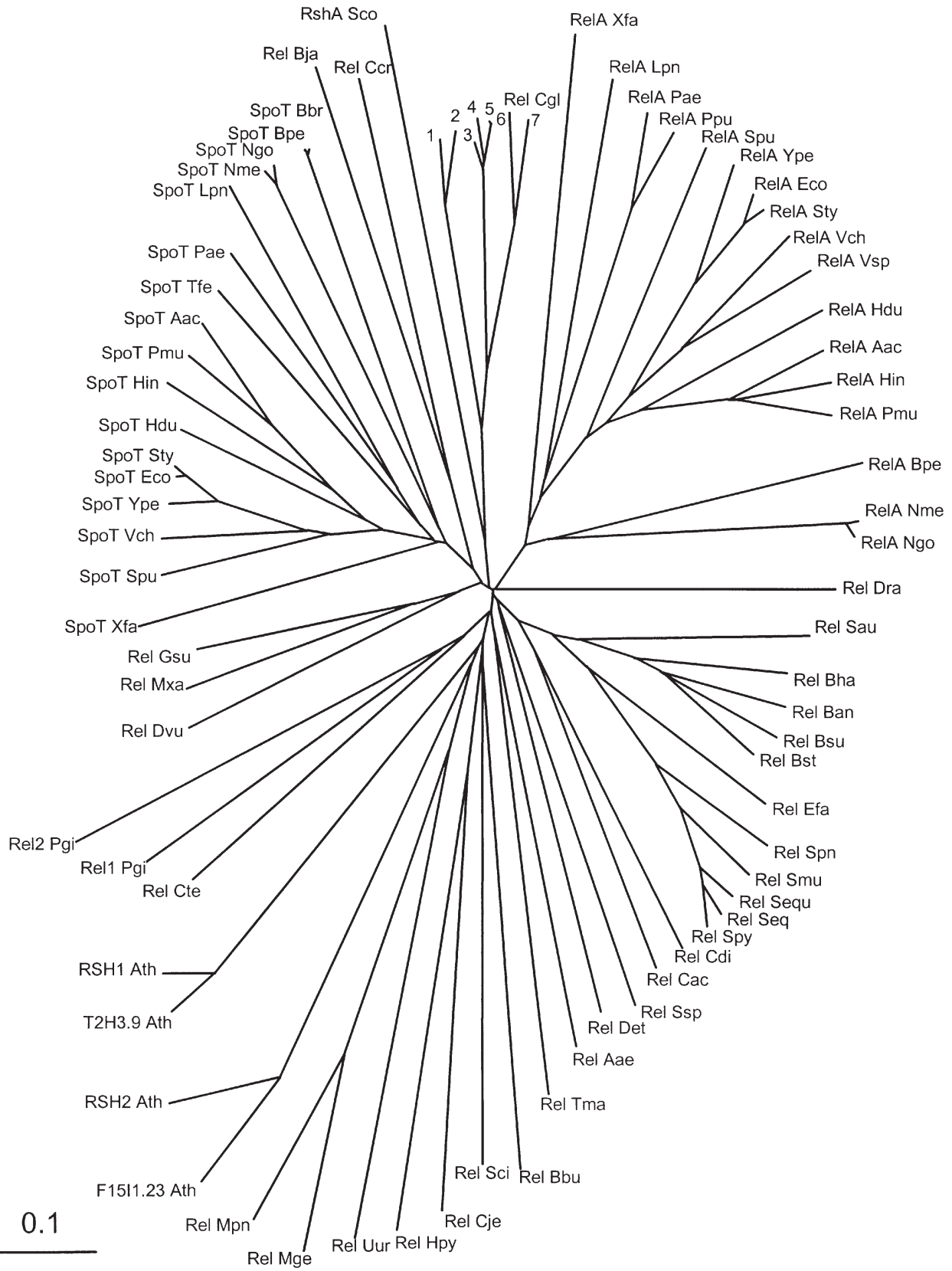


Figure 2. Radial phylogenetic tree of 80 RelA, SpoT and Rel proteins. Due to space limitations, the designation of individual Rel proteins was replaced by numbers. 1 refers to Rel Sco, 2 to Rel San, 3 to Rel Mav, 4 to Rel Mle, 5 to Rel Mtu, 6 to Rel Mbo and 7 to Rel Cdiph.

Table 4. Rel orthologs. The sequence of the Rel protein of *B. subtilis* was used as query sequence in the BLAST search. Individual proteins are sorted according to ascending E-values. For designations of species, see Table 1. For explanation of the asterisk, see Figure 2.

Designation	Length	Accession number	E-Value
Rel Bsu	734	U86377	0.0
Rel Bha	728	BAB04961	2e-278
Rel Sau	736	O32419	1.2e-241
Rel Seq	739	Q54089	9.9e-197
Rel Ssp	760	P74007	7.5e-167
Rel Mxa	757	O52177	7.5e-167
Rel Sco	847	X92520	2.6e-164
Rel San	841	O85709	3.1e-161
Rel Mtu	790	Q50638	2.9e-158
Rel Mle	787	Q49640	2.3e-156
Rel Cgl	760	O87331	2.4e-154
Rel Aae	760	O67012	9.5e-148
SpoT Eco	702	P17580	3e-140
SpoT Pae	701	AAG08723	3.1e-138
RelA Pae	747	AAG04323	3.5e-137
Rel Tma (TM0729)	751	Q9WZ18	9.3e-137
SpoT Vch (VC2710)	705	AAF95850	4.5e-135
RelA Eco	744	J04039	6.8e-134
RelA Vch	738	Q9S3S3	1.8e-133
RelA Vsp	744	P55133	1.1e-131
RelA Hin	743	P44644	5.7e-130
SpoT Nme	725	CAB85138	3.7e-129
SpoT Hin	677	P43811	2.2e-122
RelA Nme	769	CAB85211	1.3e-121
SpoT Xfa (XF0352)	735	AE003887	1.7e-120
Rel Bbu	667	O51216	1.2e-104
Rel Sci	749	O34098	1.4e-103
RelA Xfa (XF1316)	718	AE003964	6e-103
Rel Bja	779	Q9RH69	6.8e-102
Rel Cje	731	AL139077	4.8e-93
Rel Dra (DR1838)	787	Q9RTC7	2.7e-91
RshA Sco (SC4H2.15)	725	O69970	9.8e-88
Rel Hpy	776	Q9ZL68	1.3e-82
Rel Uur	718	AE002125	2.8e-78
T2H3.9 Ath	615	AAC28176	1.9e-71
Rel Mge	720	P47520	5.3e-69
Rel Mpn	733	P75386	2.3e-68
F1511.23 Ath (identical to RSH3)	715	AAD25787	1.2e-67
RSH3	712	AAF37283	e-67
SpoT Pfr *	388 (fragment)	AAG00076	3.3e-65
RSH1 Ath	883	AAF37281	8e-64
RSH2 Ath	710	AAF37282	1.1e-62
SpoT ? *	283 (fragment)	AAF73865	3.9e-43
CsrS Vsp *	119 (fragment)	Q56730	1.3e-25
SpoT Pde *	175 (fragment)	P29941	1.8e-8

Dvu) were removed from the dataset (total: 65 sequences) used for production of the alignment. However, use this dataset in the construction of a bootstrapped tree also did not result in increased resolution of the consensed part on the tree (data not shown). In a third attempt, seven other outgroup sequences (*i.e.* Rel Cte, Rel1 Pgi, Rel2 Pgi, Rel Bbu, Rel Sci, Rel Cje and Rel Hpy) were removed from the dataset. The alignment of this dataset (58 sequences) finally resulted in improved resolution of the resulting bootstrapped tree (Figure 4).

A statistically insignificant branching (53%) separates the RelA proteins from the SpoT proteins and two groups of Rel proteins from gram-positive species. A statistically significant branching (99%) separates the Rel proteins of the *Bacillus/Clostridium* group from the SpoT proteins and the actinobacterial Rel proteins. The SpoT proteins are separated from the actinobacterial Rel proteins by a statistically less significant branching (78%). An unrooted,

radial tree originating from this alignment is presented in Figure 5. It should be noted that on the other hand, when another dataset comprising only the sequences representing the condensed part of the tree of Figure 2 was used as input file for an alignment, this procedure also did not increase the resolution in the condensed part of the resulting bootstrapped tree (data not shown).

Substitutions in the HD Domain of RelA Proteins

It was recognized by Aravind and Koonin (1998) that the HD domain defines a new superfamily of metal-dependent phosphohydrolases. Aravind and Koonin (1998) define five motifs within the HD domain. Three of them (motifs I, II, and V) contain highly conserved histidine or aspartate residues. Site-directed mutagenesis experiments of genes encoding cGMP-phosphodiesterases have shown that mutations of the histidine residue in the HD signature in motif II or of the aspartate residue in motif V resulted in the

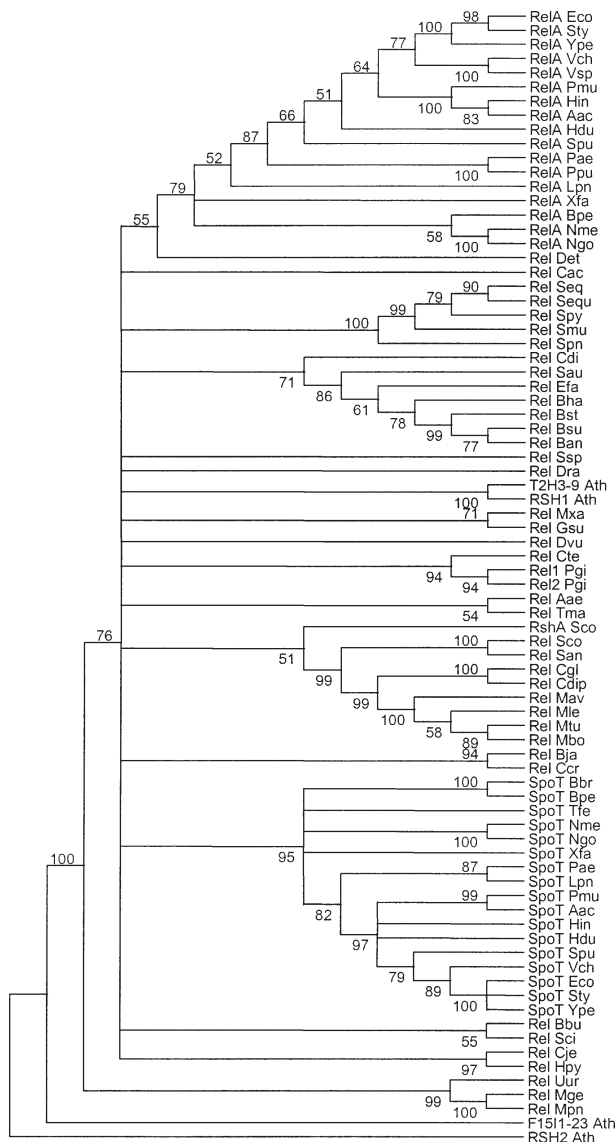


Figure 3. Rectangular cladogram of 80 RelA, SpoT and Rel proteins. Numbers on the branches indicated are bootstrap values as described in the legend to Figure 1.

most severe effect on the catalytic activity (Turko *et al.*, 1996). Aravind and Koonin (1998) showed that the RelA and SpoT proteins are also members of this superfamily. But the RelA protein of *E. coli* contains substitutions in the predicted catalytic residues of this domain. Aravind and Koonin (1998) suggest that the HD domain of RelA is inactivated but still retains its native structure. An inactive phosphohydrolase domain helps to explain why RelA proteins are not able to degrade ppGpp. On the other hand, the N-terminus of SpoT encompassing the HD domain is sufficient for ppGpp hydrolase activity (Gentry and Cashel, 1996). An alignment of the N-termini of the SpoT, Rel and RelA proteins is presented in Supplementary Material (<http://jmmb.net/supplementary>). This alignment shows the region around conserved motifs I and II and the region around conserved motif V. Importantly, all the proteins

identified in this study as putative RelA proteins carry substitutions in all conserved regions whereas Rel and SpoT proteins and the *Arabidopsis* paralogs possess a functional HD domain (<http://jmmb.net/supplementary>). Interestingly, in most substitutions of the HD signature sequence, insertion of a proline residue took place, an amino acid known to influence protein folding (*e.g.* Eyles and Gierasch, 2000). Nevertheless, the SpoT, Rel and RelA proteins share some overall similarity in this region. This analysis also suggests that the protein designated RelA of *M. xanthus* by Harris *et al.* (1998) is in fact a Rel or SpoT protein.

Comparison of Putative ppGpp Binding Sites

During this analysis, we noted that genes encoding (p)ppGpp synthetases/hydrolases are absent in the completely sequenced genomes of the obligate parasites *Chlamydia pneumoniae* and *C. trachomatis* (Kalman *et al.*, 1999), *Rickettsia prowazekii* (Andersson *et al.*, 1998), and *Treponema pallidum* (Fraser *et al.*, 1998). Similarly, the genome of an intracellular symbiont of aphids, *Buchnera* sp. APS (Shigenobu *et al.*, 2000) does not contain a RelA or SpoT gene, although this extremely adapted bacterium is a close relative of *E. coli*. It seems likely that the *rel* genes were deleted during the adaptation to the intracellular environment and the establishment of specialized lifestyles in a process called "reductive evolution" (Andersson and Kurland, 1998).

Genetic studies (Glass *et al.*, 1986) using strains carrying amber mutations in *rpo* genes and biochemical investigations (Reddy *et al.*, 1995; Chatterji *et al.*, 1998; Touloukhonov *et al.*, 2001) using various ppGpp analogs as crosslinking agents identified RNAP of *E. coli* as target for ppGpp. Glass *et al.* (1986) concluded that ppGpp binds to the β subunit encoded by *rpoB* and also identified a common motif in purine-nucleotide binding proteins (GXXXXGK; la Cour *et al.*, 1985) in the amino acid sequence of RpoB at residues 880 to 886. By a combination of a variety of methods (SDS-PAGE analysis of the [32 P] N_3 ppGpp (azido-ppGpp)-bound enzyme, tryptic digestion and Western blots), Chatterji *et al.* (1998) identified a 45 kDa fragment of the β subunit as binding site for ppGpp. This fragment is located at the C-terminus and consists of residues 802-1211, 1216 or 1223 (Chatterji *et al.*, 1998). Using the smaller ppGpp analog 6-thio-ppGpp, however, the β' subunit encoded by *rpoC* was very recently identified as target for ppGpp (Touloukhonov *et al.*, 2001). The binding site for ppGpp was determined at the N-terminus (between residues 29-102) of the β' subunit. The authors explain these conflicting results by the properties of the different crosslinking reagents and by the 3D model of RNAP (Zhang *et al.*, 1999). This model shows that the N-terminus of β' and the C-terminus of β are spatially close and constitute an intertwined interface. Touloukhonov *et al.*, (2001) postulate that binding of ppGpp to RNAP is allosteric, that the binding site is modular and is located close to the intersubunit interface of the N-terminal and C-terminal ends of β' and β .

In order to reveal the presence or absence of the different identified ppGpp binding regions in the β' and β sequences, alignments of RpoB (Table 6; <http://jmmb.net/supplementary>) and RpoC (Table 7; <http://jmmb.net/>)

Table 5. Rel orthologs. The sequence of the Rel protein of *B. subtilis* was used as query sequence in the BLAST search of the "unfinished microbial genomes" database at TIGR. For designations of species, see Table 1. Individual proteins are sorted according to ascending E-values.

Designation	Length	Contig designation	E-value
Rel Ban	727	gnl TIGR_1392 banth_1489	0.0
Rel Bst	707 (fragment)	gnl UOKNOR_1422 bstear_Contig408	0.0
Rel Efa	737	gnl TIGR gef_10492	0.0
Rel Cdi	735	gnl Sanger_1496 cdifficile_Contig876	0.0
Rel Spn	740	gnl TIGR S.pneumoniae_3836	0.0
Rel Smu	740	gnl UOKNOR_1309 S.mutans_Contig98	0.0
Rel Spy	739	gnl OUACGT_1315 Spyogenes_Contig1	0.0
Rel Cac	740	gnl GTC C.aceto_gnl	0.0
Rel Sequ	719 (fragment)	gnl Sanger_1336 sequi_Contig474	0.0
Rel Gsu	716	gnl TIGR_35554 gsulf_57	0.0
Rel Det	728	gnl TIGR_61435 deth_1541	0.0
Rel Mbo	738	gnl Sanger_1765 mbovis_Contig403	e-171
Rel Cdiph	759	gnl Sanger_1717 cdiph_Contig2	e-166
Rel Dvu	717	gnl TIGR_881 dvulg_159	e-166
Rel Mav	647	gnl TIGR M.avium_223	e-162
SpoT Tfe	759	gnl TIGR t.ferrooxidans_6160	e-149
SpoT Ype	707 (fragment)	gnl Sanger_632 Y.pesits_Contig1008	e-148
SpoT Sty	709 (fragment)	gnl Sanger_601 S.typhi_CT18	e-148
RelA Sty	744	gnl Sanger_601 S.typhi_CT18	e-142
RelA Ype	744	gnl Sanger_632 Y.pesits_Contig854	e-146
RelA Ppu	746	gnl TIGR pputida_10724	e-145
Rel Cte	731	gnl TIGR C.tepidum_3499	e-143
SpoT Spu	712 (fragment)	gnl TIGR_24 sputre_6408	e-142
RelA Spu	735	gnl TIGR_24 sputre_6426	e-142
RelA Lpn	734	gnl CUCGC_446 lpneumo_MF.18.110397	e-141
RelA Pmu	739	gnl CBCUMN_747 Pmultocida	e-140
SpoT Pmu	711 (fragment)	gnl CBCUMN_747 Pmultocida	e-139
SpoT Aac	712 (fragment)	gnl OUACGT_714 A.actin_Contig253	e-139
SpoT Bbr	759	gnl Sanger_518 bbronchi_Contig2526	e-140
SpoT Bpe	759	gnl Sanger_520 B.pertussis_Contig236	e-140
RelA Aac	743	gnl OUACGT_714 A.actin_Contig233	e-136
SpoT Ngo	718	gnl OUACGT_485 Ngon_Contig1	e-134
RelA Hdu	735	gnl HTSC_730 ducreyi	e-133
RelA Ngo	737	gnl OUACGT_485 Ngon_Contig1	e-129
RelA Bpe	737	gnl Sanger_520 B.pertussis_Contig301	e-120
SpoT Hdu	569 (fragment)	gnl HTSC_730 ducreyi	e-111
Rel1 Pgi	762	gnl TIGR P.gingivalis_GPG.con	e-114
Rel Ccr	743	gnl TIGR C.crescentus_12574	e-113
SpoT Lpn	530 (fragment)	gnl CUCGC_446 lpneumo_MF.91.102397	e-94
Rel2 Pgi	746	gnl TIGR P.gingivalis_GPG.con	e-81

supplementary) amino acid sequences were performed. Our Supplementary Material (<http://jmmmb.net/supplementary>) shows that the common motif in purine nucleotide-binding proteins described above is present also in RpoB sequences of obligately parasitic bacteria and in the symbiont *Buchnera* sp. APS. Large insertions around

residues 1140 to 1260 of the RpoB alignment are present in the proteobacterial and chlamydial sequences and again around residues 1340 to 1420 in the proteobacterial, mycobacterial and mycoplasmal sequences. Most interestingly, the RpoB and RpoC sequences of the species which do not possess a *rel*-like gene do not exhibit specific

Table 6. RpoB sequences compared in this study. For designations of species, see Table 1.

Designation	Length	Accession number
RpoB Eco	1342	RNEBC
RpoB Bsu	1193	P37870
RpoB Tpa	1178	O83269
RpoB Bsp	1342	BAB12761
RpoB Rpr	1374	O52271
RpoB Ctr	1252	O84317
RpoB Cpn	1252	BAA98291
RpoB Hin	1343	P43738
RpoB Sau	1182	P47768
RpoB Mpn	1391	P78013
RpoB Bbu	1155	AAB91501
RpoB Mtu	1172	CAB09390

Table 7. RpoC sequences compared in this study.

Designation	Length	Accession number
RpoC Eco	1407	RNECC
RpoC Bsu	1199	P37871
RpoC Sau	1057	P47770
RpoC Mtu	1316	P47769
RpoC Bsp	1407	BAB12760
RpoC Hin	1415	P43739
RpoC Rpr	1372	Q9ZE20
RpoC Ctr	1396	O84316
RpoC Cpn	1393	Q9Z999
RpoC Tpa	1416	O83270
RpoC Bbu	1377	O51349
RpoC Mpn	1290	P75271

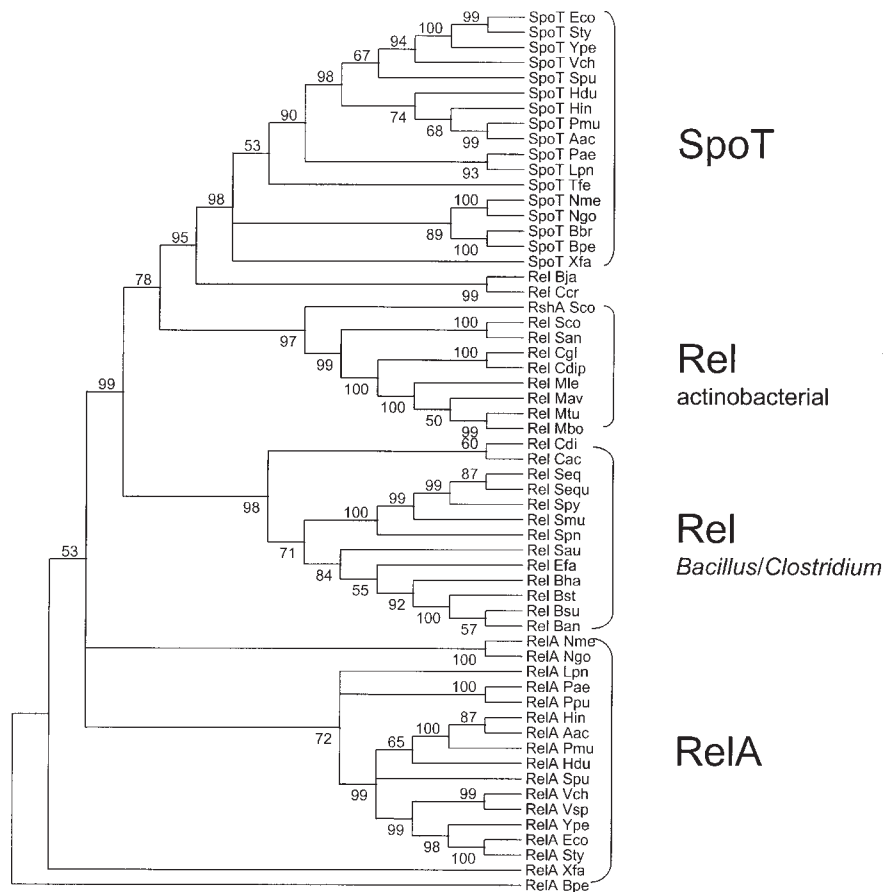


Figure 4. Rectangular cladogram of 58 RelA, SpoT and Rel proteins. Four different groups can be distinguished. Numbers on the branches indicated are bootstrap values as described in the legend to Figure 1.

alterations. This might indicate that the binding sites for ppGpp are still present in the β and β' subunits of these species.

Discussion

Absence of ppGpp in Archaea, Presence of ppGpp in Plants

Genes encoding (p)ppGpp synthetases are absent in all completely sequenced genomes of Archaea. Since eubacterial RNA polymerase is the target for the regulatory action of (p)ppGpp and since archaea possess a transcriptional apparatus similar to that of eukaryotes instead (Langer *et al.*, 1995), this difference between bacteria and archaea might explain this absence. Starvation studies in halobacteria demonstrated stringent control of stable RNA biosynthesis but both growth rate control and stringent control are probably governed by mechanisms that operate in the absence of ppGpp (Cimmino *et al.*, 1993; Scoarughi *et al.*, 1995).

The finding that *Arabidopsis* possesses RelA/SpoT homologs (van der Biezen *et al.*, 2000) might indicate that plants have retained some aspects of the bacterial transcription apparatus. Indeed, in *Arabidopsis* proteins similar to bacterial sigma factors (σ^{70}) have been identified

as products of genes specifically expressed in leaves and destined for chloroplasts (Isono *et al.*, 1997). Also, chloroplast genomes encode subunits of bacterial-type RNA polymerases (Sugiura *et al.*, 1998; Turmel *et al.*, 1999; Allison, 2000). Furthermore, ppGpp was detected in the alga *Chlamydomonas reinhardtii* (Heizmann and Howell, 1978).

Syntenic Considerations

Syntenic studies (comparison of the relative positions of genes in the genome of different organisms) might be a source for auxiliary information to identify orthologous genes in genome sequencing projects (Huynen and Bork, 1998). In a systematic comparison using 256 operons of *E. coli* and 100 operons of *B. subtilis* as query sequences, Itoh *et al.* (1999) tried to identify operons showing a conserved order of orthologous genes in 11 complete genome sequences. These authors found that operon structures are very rarely conserved. The sequences of the *relA* and *spoT* operons of *E. coli* (but not the sequence of the corresponding *rel* gene of *B. subtilis*) were also used as query sequences. This analysis showed that no genome contains operons showing exactly the same organization as the *spoT* and *relA* operons of *E. coli*. However, due to the complexity of the *spoT* operon, the function of this

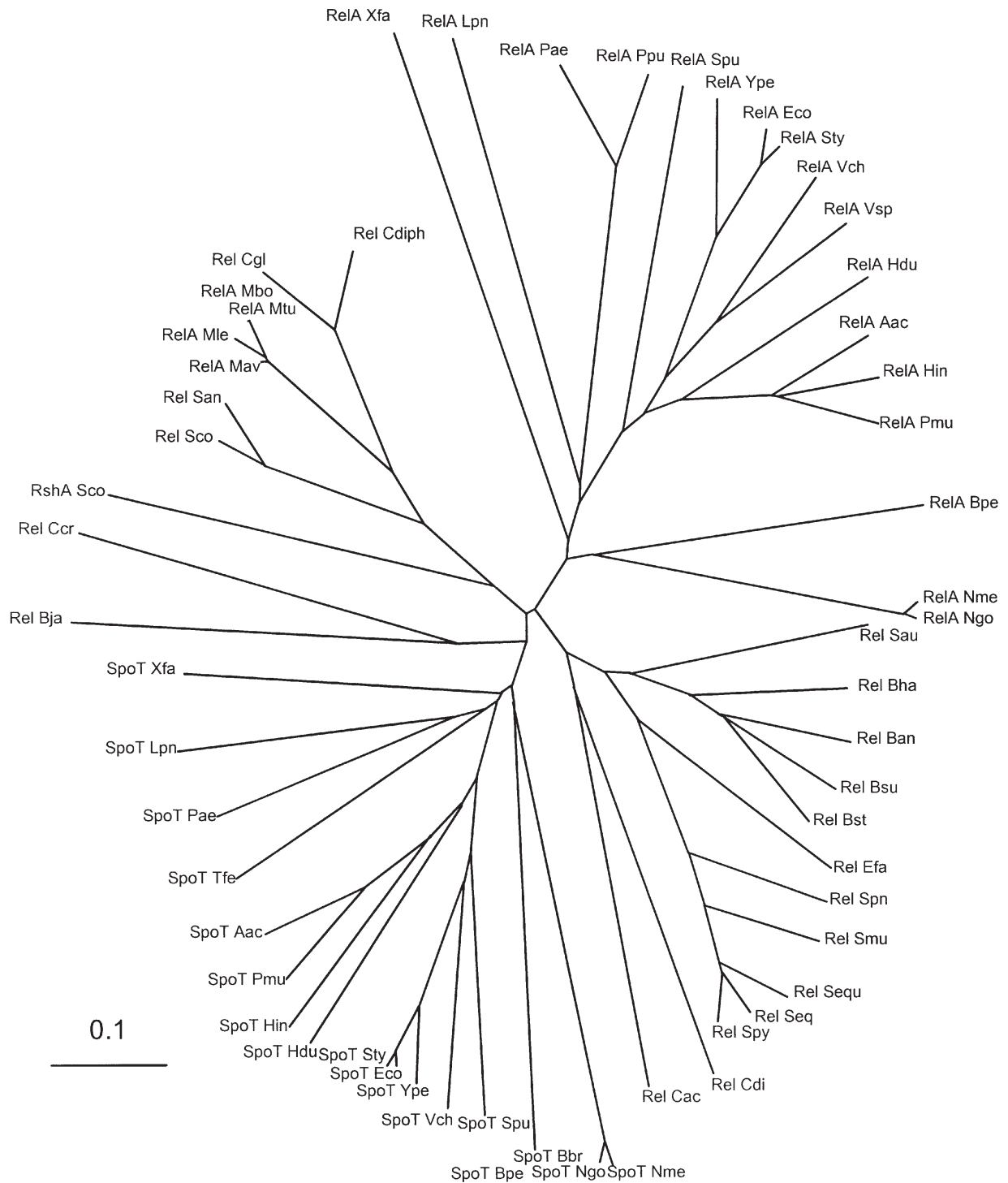


Figure 5. Radial phylogenetic tree of 58 RelA, SpoT and Rel proteins.

operon was misclassified as "DNA/RNA processing" in this study. For this reason, a short description of the *relA* and *spoT* operons of *E. coli* and the *rel* gene region of *B. subtilis* is given.

In *E. coli*, the *relA* gene is the first gene in an operon containing also the genes encoding the MazEF (*ma-ze*, hebrew for "what is it") antitoxin/toxin module (Aizenman *et al.*, 1996). Two promoters of the *mazEF* genes are

negatively autoregulated by MazE and MazF and expression of the module is positively regulated by FIS (Marianovsky *et al.*, 2001). In *B. subtilis* and other gram-positive bacteria, the putative *mazEF* locus is located upstream of the *sigB* operon (Mittenhuber, 1999), encoding the general stress response (Hecker and Völker, 1998) sigma factor σ^B and its regulatory proteins (Wise and Price, 1995). A third gene in the *relA* operon, *mazG*, encodes a

protein with unknown function. A BLAST search revealed that similar genes are present in many bacterial genomes (data not shown). They are designated as similar to a hypothetical 26.1 kDa protein named YBL1 (accession number P33653) of *Streptomyces cacaoi* (Urabe and Ogawara, 1992).

The *spoT* gene of *Escherichia coli* is the third gene in an operon consisting of five genes. The order is *gmk-rpoZ-spoT-trmH-recG*. *Gmk* encodes guanylate kinase (Gentry *et al.*, 1993), *rpoZ* the Ω -subunit of RNAP (Gentry and Burgess, 1989), *trmH* a tRNA modifying enzyme (tRNA (Gm18) 2'-O-methyltransferase) (Persson *et al.*, 1997) and *recG* a DNA helicase (Lloyd and Sharples, 1991; Kalman *et al.*, 1992). Interestingly, the first three genes are linked to guanosine metabolism whereas the *trmH* and *recG* gene products play a role in RNA and DNA processing. Lloyd and Sharples (1991) identified a putative promoter near the end of *spoT*, indicating that *trmH* and *recG* might be transcribed as separate unit.

The organization of the *rel* loci of *B. subtilis*, *M. leprae*, *M. tuberculosis*, *S. equisimilis* and *S. coelicolor* is similar (Wendrich and Marahiel, 1997; Avarbock *et al.*, 1999). Upstream of the *rel* gene, the *apt* genes (encoding adenine phosphoryltransferase catalyzing the formation of adenosine monophosphate from adenine and phosphoribosyl pyrophosphate in a salvage reaction) and genes encoding components of the protein export machinery (*secDF*) are located in the same direction and downstream of *rel*, the *cypH* genes encoding cyclophilin (a peptidyl-prolyl *cis-trans* isomerase) are located in the opposite direction.

Proposal for the Evolution of the Rel, RelA and SpoT Proteins

The Mollicutes (*Mycoplasma* and relatives) which are normally associated with the *Bacillus/Clostridium* group of gram-positive bacteria form a distinct outgroup in the trees. These organisms undergo faster rates of evolution and quite regularly form outgroups in phylogenetic trees of orthologous protein sequences (Eisen, 1998).

With the exception of *Neisseria* and *Bordetella* species both belonging to the β -proteobacteria, two different genes encoding enzymes involved in (p)ppGpp synthesis and degradation, namely RelA and SpoT are only found in the β and γ subdivision of proteobacteria. The SpoT genes of this group are related to genes encoding the actinobacterial group and the *Bacillus/Clostridium* group of Rel proteins (Figures 4 and 5; see also the E-values in Table 3). Since complete protein sequences were compared, the separation of the RelA proteins from the Rel and SpoT proteins is most probably due to the fact that the Rel and SpoT proteins possess ppGpp hydrolase activity, whereas the RelA proteins lack this activity. It is also interesting to note that the paralogous genes of individual species are not closely related to each other. Multiple, parallel gene duplications in individual species as origin of the *relA* and *spoT* genes can therefore be excluded.

Facilitated by the knowledge of the enzymatic activities of the Rel, RelA and SpoT proteins and based on some logical speculation, a model for evolution of the RelA and SpoT proteins within the β and γ subdivision of proteobacteria is proposed:

(1) The common ancestor of the β - and γ -proteobacteria possessed a *rel* gene of actinobacterial origin. The *spoT* gene evolved from this gene under adaptation of the gene product to carbon and fatty acid starvation.

(2) Following gene duplication, the additional copy of this ancestral gene is able to adopt to new functions. In this case, this adoption of new function was most probably inactivation of the HD domain, loss of (p)ppGpp hydrolyzing activity and adaptation to amino acid starvation. This copy evolved to the *relA* gene of β - and γ -proteobacteria.

The Special Case of *P. gingivalis*

P. gingivalis, however represents an interesting exception from the scenario described in the previous paragraph: This organism possesses a *relA* and a *spoT* gene (Sen *et al.*, 2000; <http://jmmb.net/supplementary>) which are however not related to the proteobacterial *relA* and *spoT* genes (Figure 3). Both paralogs which are named Rel1 Pgi and Rel2 Pgi in this paper are closely related to each other and to Rel of *Chlorobium tepidum* (Figure 3). In *P. gingivalis*, the evolution of the paralogous *rel1* (*spoT*-related) and *rel2* (*relA*-related) occurred independently of the proteobacterial *relA* and *spoT* genes. At the moment, it is impossible to decide whether this paralogy represents an example of convergent evolution. Alternatively, lateral gene transfer of short, catalytically active domains of ppGpp synthetases/hydrolases might have been involved in the evolution of the *rel1* and *rel2* genes of *P. gingivalis*. In order to solve this question, more sequences encoding ppGpp synthetases/hydrolases from representatives of the CFB (*Cytophaga-Flavobacterium-Bacteroides*) phylum and green sulfur bacteria should be determined and compared (see also next paragraph). Alternatively, a careful phylogenetic analysis of the individual domains of the Rel, RelA and SpoT proteins (Aravind and Koonin, 1998), which is however beyond the scope of this paper might help to solve this issue. Such an analysis has been performed for the separate domains of the conserved HSP70 (DnaK) family: Striking differences in the relative rate of amino acid replacement in different rates were observed and were interpreted as evidence for functional divergence (Hughes, 1993).

ppGpp Synthetases/Hydrolases and Bacterial Evolution

A drastically different view of bacterial evolution has been recently postulated by Gupta (2000a, b). This hypothesis places the γ subdivision of proteobacteria at the end of an evolutionary linear scheme. According to this theory, the γ subdivision of proteobacteria constitutes the most recently evolved bacterial group. Therefore, specialized, paralogous *spoT* and *relA* genes might be a relatively new invention of bacterial evolution which might explain this limited distribution pattern among the β - and γ -proteobacteria. Similarly, the (maybe more efficient) vitamin B₆ biosynthesis pathway of *E. coli* is mainly restricted to the γ subdivision, whereas another widely conserved, biochemically uncharacterized pathway is presumably operating in other species (Mittenhuber, 2001). Other unique features of the γ -proteobacteria were recently recognized by Margolin (2000) in his study on prokaryotic cell division: Genes encoding the ZipA, FtsL and FtsN proteins are only found

in genomes of completely sequenced γ -proteobacteria. Very recently, Eisen (2000) noted that the currently available datasets of completely sequenced genomes are not representative of evolutionary diversity. It might be predicted that the availability of completely sequenced genomes of proteobacterial species belonging to subdivisions other than β and γ might help to identify the precursor of the proteobacterial *relA* and *spoT* genes.

ppGpp is Not Present in Obligately Intracellular Species

The species lacking a *rel*-like gene require living cells for their replication and growth and cannot be cultivated *in vitro*. The absence of *rel*-like genes in genomes of obligately parasitic organisms was also detected in a comparative genome analysis of *B. burgdorferi* and *T. pallidum* (Subramanian *et al.*, 1999). *R. prowazekii* possesses a few short pieces of a pseudogene which show strong sequence similarity to the *relA/spoT* homologs (Andersson *et al.*, 1998; Zomorodipour and Andersson, 1999), whereas no *rel*-like gene is present in the *C. trachomatis* genome (Stephens *et al.*, 1998). Interestingly, some cultivable intracellular pathogens (*e.g.* *B. burgdorferi* and *Mycoplasma* species among others) possess *rel*-like genes. In most cases, cultivation of bacteria implies that the inoculum is able to form colonies. Investigations on vertical sections of *E. coli* colonies show that the colony is composed of different layers of bacteria, containing also nonviable bacteria (Shapiro, 1994). It is very likely that many cells in a developed colony are starved for nutrients and that the stringent response is switched on in these cells. It might be extremely interesting to investigate whether a functional connection between *in vitro* cultivability of bacterial species and the absence of genes encoding (p)ppGpp synthetases/hydrolases in obligate parasites can be established experimentally.

Experimental Procedures

Using the deduced protein sequences of the *relA* and *spoT* genes of *E. coli* and of the *rel* gene of *B. subtilis* as query sequences, BLAST searches (Altschul *et al.*, 1997) of the non-redundant protein database nrdb95 (Holm and Sander, 1998) were performed at <http://dove.embl-heidelberg.de/Blast2/> using the *blastp* program. BLAST searches of unfinished microbial genomes using the deduced protein sequence of the *rel* gene of *B. subtilis* as query were performed at http://www.ncbi.nlm.nih.gov/Microb_blast/unfinishedgenome.html using the program *tblastn*. Raw DNA sequences were translated using the program "Translate tool" at <http://www.expasy.ch/tools/dna.html>. The COG database can be assessed at <http://www.ncbi.nlm.nih.gov/COG>. Alignments were generated using CLUSTALW (Thompson *et al.*, 1994) at <http://www.ebi.ac.uk/clustalw/>. Radial phylogenetic trees were constructed using the data from the alignments with the help of the program TreeView (<http://taxonomy.zoology.gla.ac.uk/rod/treeview.html>; Page, 1996) and edited using the program Metafile Companion (<http://www.companionsoftware.com>). Bootstrapping of datasets was performed by the neighbor-joining (NJ) method on the basis of the Poisson-corrected amino acid distance (d_{aa}) (Saitou and Nei, 1987; for a discussion of different statistical

methods and their applications see Hughes, 1999; Nei and Kumar, 2000) using MEGA 2.0 (<http://www.megasoftware.net>; Kumar *et al.*, 2001).

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