Common and specific amino acid residues in the prokaryotic polypeptide release factors RF1 and RF2: possible functional implications

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ABSTRACT
Termination of protein synthesis is promoted in ribosomes by proper stop codon discrimination by class 1 polypeptide release factors (RFs). A large set of prokaryotic RFs differing in stop codon specificity, RF1 for UAG and UAA, and RF2 for UGA and UAA, was analyzed by means of a recently developed computational method allowing identification of the specificity-determining positions (SDPs) in families composed of proteins with similar but not identical function. Fifteen SDPs were identified within the RF1/2 superdomain II/IV known to be implicated in stop codon decoding. Three of these SDPs had particularly high scores. Five residues invariant for RF1 and RF2 [invariant amino acid residues (IRs)] were spatially clustered with the highest-scoring SDPs that in turn were located in two zones within the SDP/IR area. Zone 1 (domain II) included PxT and SPF motifs identified earlier by others as ‘discriminator tripeptides’. We suggest that IRs in this zone take part in the recognition of U, the first base of all stop codons. Zone 2 (domain IV) possessed two SDPs with the highest scores not identified earlier. Presumably, they also take part in stop codon binding and discrimination. Elucidation of potential functional role(s) of the newly identified SDP/IR zones requires further experiments.

INTRODUCTION
Three triplets, UAA, UAG and UGA, located at the end of coding mRNA sequences, are signals for terminating polypeptide synthesis on ribosomes. When one of these stop codons encounters the ribosomal A site while the P site is occupied by peptidyl-tRNA, a protein, called class 1 release factor (RF), binds to the A site [for recent reviews see (1–4)]. In prokaryotes RF1 decodes UAA and UAG stop codons, whereas RF2 is specific for UAA and UGA. The genetic code of mitochondria and mycoplasmas is modified and does not contain the UGA stop codon. Consequently, they possess only one RF that corresponds to bacterial RF1 (5). In eukaryotes, only one factor, named eRF1, decodes all three stop codons (6). Although it has been proved that eRF1 rather than the ribosome itself determines the specificity of stop codon decoding (7,8), the decoding mechanism remains enigmatic despite the fact that some amino acids essential for stop codon discrimination by eRF1 have already been identified (9–11).

In contrast to eukaryotes, for prokaryotic factors two ‘protein anticodons’ have been identified, PxT in RF1 and SPF in RF2 (12,13). Site-specific mutagenesis of these residues both in vivo and in vitro led to a switch in the stop codon specificity of the mutant Escherichia coli RFs. Presumably, these residues are involved in discrimination of the second and third purines of stop codons (12–14). However, it remains unknown whether the decoding specificity of RF1/RF2 is determined solely by these tripeptides or other amino acids also contribute to the discrimination process.

In an attempt to get better insight into stop codon recognition by bacterial RFs, we applied a newly developed algorithm SDPpred that automatically selects amino acid residues accounting for differences in functional specificity among homologous proteins (15,16). The prerequisite for applying this method is a common biochemical function for the selected family of proteins and different specificity in subfamilies. The RF1/RF2 case meets this requirement because of their well-known functional and structural similarity. This method…
predicts specificity-determining positions (SDPs) that by definition are positions of multiple alignment which are conserved within subfamilies consisting of the proteins with the same specificity, but differ between these subfamilies. The major advantage of this approach stems from the fact that in contrast to some other previously suggested techniques, SDPpred directly takes into account the non-uniformity of amino acid substitution frequencies, automatically selects the fraction of alignment positions with the highest correlations with the specificity and finds differences that are least probable to arise by chance during evolution. Furthermore, SDPpred does not require information about the 3D protein structure, which instead can be used at the post-processing step to validate the predictions. There are also other approaches for comparative analysis of protein subfamilies. One of these methods, called DIVERGE (17) takes into account the differences in mutation rates in different parts of the phylogenetic tree of the protein family and identifies sites that had experienced an RS during their evolution.

In this work we compared the results of RF1/RF2 family analysis by means of SDPpred and DIVERGE. In addition we considered invariant amino acid residues (IRs) common for both subfamilies. All identified SDPs and IRs were mapped onto the 3D structure of both RFs. The bioinformatics approach predicts new functionally essential amino acid residues in RF family.

**MATERIALS AND METHODS**

**Preparation of datasets**

The RF1 and RF2 amino acid sequences of *E.coli* were used as the seeds for the BLAST (18) search. RF-like proteins (1334) were found in the current protein sequence databases (GenBank translations, PIR and others). We used the ClustalX software (ftp://ftp-igm.c.u-strasbg.fr/pub/ClustalX/) to align all these sequences. A maximum-likelihood species tree was reconstructed with the PROTML with the JTT amino acid replacement model from the PHYLIP package (http://evolution.genetics.washington.edu/phylip.html) (20). RF sequences from mycoplasmal or mitochondrial genomes were not included in the training set, as they are deficient in RF2 and could therefore distort the pattern of RF1 specificity. Based on the multiple alignment and the phylogenetic tree we discarded all weak homologs, and all RFs containing large N- or C-terminal deletions. Then, among the remaining organisms only those with both RF1 and RF2 sequences were used for further analysis. Thus we obtained the multiple alignment of 234 RF1/RF2 orthologs from the genomes of various prokaryotes, consisting of 117 RF1s and 117 RF2s. The final alignment was manually controlled and edited using the BIOEDIT multiple alignment analysis package (http://www.mbio.ncsu.edu/BioEdit/page2.html). Selected sequences and alignments are available as Supplementary Data.

**Selection of SDPs and RSPs**

The multiple sequence alignment (MSA) of 234 RF sequences was used as the training set for SDP selection. SDPpred compares amino acid residues in paralogous protein subfamilies: it computes mutual information for each position of the MSA; then, using random shuffling of each column, calculates the statistical significance of the observed values (Z-scores); and finally, using the Bernoulli estimator procedure, finds those columns of the MSA that have the highest Z-scores (SDPs) and have the minimal probability to arise by chance. SDPpred is available at http://math.genebee.msu.su/~psn. The set of obtained SDPs was tested as the profile for attribution of RF protein sequences not included into the training set.

The DIVERGE (17) algorithm utilizes two models of mutation rate variation among sites: the ‘homogeneous gamma model’, which assumes that the mutation rate at each site remains constant over the whole history of a protein family, and the ‘non-homogeneous gamma model’, which allows sites to mutate at different rates in various branches of the family tree (RS). DIVERGE calculates posterior probabilities of each position to follow the non-homogeneous model of functional divergence. If this probability equals 1 or is close to 1, this indicates positions, which experienced the RS. We refer to these positions as RSP positions. We used the cutoff of 0.99 for selection of RSPs.

**Selection of IRs clustered with SDPs**

Amino acid residues invariant in both RF1 and RF2 (IRs) were selected in the same MSA that was used for SDP and RSP prediction. For filtering of potential sequence errors we compared IRs list with ‘highly conserved residues’ (in this case we allowed 98% identity). The spatial localization of the selected IRs was further analyzed. We calculated the centroids (centres of mass of their side chains) for all SDPs and IRs using the SYBYL software (http://www.tripos.com). The crystal structures for both RF1 (21) and RF2 (22) were used for SDP and IR centroid identification. We calculated pairwise distances between these centroids (data not shown) for both crystal structures and selected the fraction of IRs that were the closest spatial neighbors of SDPs. For this fraction the IR–SDP distances varied from 3 to 7 Å. Solvent accessibility of SDPs and IRs was calculated using SYBYL (http://www.tripos.com) and FANTOM (http://www.scsb.utmb.edu/fantom/fm_home.html).

**RESULTS**

**Comparative analysis of the RF1 and RF2 multiple alignment**

The MSA of the RF1/RF2 training set was subjected to phylogenetic analysis. The phylogenetic tree, as expected, split into two branches formed separately by RF1s and RF2s. Thus we excluded the possibility of RF misannotation in the training set. The RF1 (UAG specific) and RF2 (UGA specific) parts of the seeds for the BLAST search. RF-like proteins (1334) were found in the current protein sequence databases (GenBank translations, PIR and others). We used the ClustalX software (ftp://ftp-igm.c.u-strasbg.fr/pub/ClustalX/) to align all these sequences. A maximum-likelihood species tree was reconstructed with the PROTML with the JTT amino acid replacement model from the PHYLIP package (http://evolution.genetics.washington.edu/phylip.html) (20). RF sequences from mycoplasmal or mitochondrial genomes were not included in the training set, as they are deficient in RF2 and could therefore distort the pattern of RF1 specificity. Based on the multiple alignment and the phylogenetic tree we discarded all weak homologs, and all RFs containing large N- or C-terminal deletions. Then, among the remaining organisms only those with both RF1 and RF2 sequences were used for further analysis. Thus we obtained the multiple alignment of 234 RF1/RF2 orthologs from the genomes of various prokaryotes, consisting of 117 RF1s and 117 RF2s. The final alignment was manually controlled and edited using the BIOEDIT multiple alignment analysis package (http://www.mbio.ncsu.edu/BioEdit/page2.html). Selected sequences and alignments are available as Supplementary Data.

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The MSA of the RF1/RF2 training set was subjected to phylogenetic analysis. The phylogenetic tree, as expected, split into two branches formed separately by RF1s and RF2s. Thus we excluded the possibility of RF misannotation in the training set. The RF1 (UAG specific) and RF2 (UGA specific) parts of
function. For this reason, we further considered separately the subgroup of SDPs with the highest Z-scores (Table 1). These positions formed a distinct group preceding the first local minimum on the Bernoulli estimator plot (data not shown). The second SDP group consisted of middle-scoring SDPs (positions 143, 144, 201, 202, 207, 208, 213, 217, 320, 325, 327, 334). These SDPs were clustered in space with the highest-scoring SDPs in superdomain II/IV (Figure 2) and probably were of secondary role in specific functions of RF1/RF2 family.

The generated SDP profile was tested at RF1 and RF2 sequences not included in the training set (as mentioned in Materials and Methods). All of them were attributed to RF1 or RF2 specificity group as anticipated from their annotations. We also compared the selected SDPs of RF1 with the vertebrate mitochondrial genetic code, where it is believed that not only RF1-specific UAG and UAA stop codons serve as termination signals but in addition also the non-canonical AGA and AGG triplets (summarized in the NCBI Taxonomy Browser http://www.ncbi.nlm.nih.gov/Taxonomy/Utils/wprintgc.cgi?mode=c) (5).

Initially, experiments on the prokaryotic RF1/RF2 stop codon decoding sites were based solely on the amino acid sequences (12) without knowing the 3D structures of RFs. Now the crystal structures became known for RF2 (22) and very recently for RF1 (21). The previously described ‘protein anticodon’ (12) or ‘tripeptide discriminator’ (13) motifs are located in domain II in the RF1/RF2 crystal structures. SDPs 201, 202, 205, 207, 208, 213 and 217 are located in the same region (Figure 2). The Ser205 residue (the first position of the PxT/SPF motif) belongs to the highest-scoring SDP group whereas the Phe207 residue (the third position of PxT/SPF motif) is one of the middle-scoring SDPs (Table 1). The structure of superdomain II/IV in both RFs appeared to be very similar if not identical (21,22). The location of SDPs on these structures is shown in Figure 3.

We compared the locations of the SDPs on the E.coli RF2 (22) and Thermotoga maritima RF1 (21) crystal structures, and on the cryo-EM-based model of the ribosome-bound E.coli RF2 (23,24). The main difference between the crystal and the ribosome-bound structures is associated with positioning of domain III that contains the universal GGQ tripeptide (25) against the other domains of RFs. Superdomain II/IV is similar in all three structures. There are no SDPs outside this superdomain (Table 1). In space, SDPs 143 and 144 are located proximal to the SPF-containing region. Therefore, these residues were attributed to the same spatial zone as SDPs in positions 201–217 (Figure 2).

SDPs of domain IV are located in two distinct regions. SDPs 322, 325, 327 and 334 clustered with domain II SDPs 201–217 (Figure 2). None of these SDPs are high-scoring (Table 1). SDPs 319 and 320 possess the highest Z-scores (Table 1) indicating their importance for RF1/RF2 specificity. They are located in the distal part of the domain IV of the RF 3D structure (Figure 3). Comparing the spatial location of the highest-scoring SDPs we observed two distinct zones at the 20–30 Å distance from each other. Zone 1 included position 205 while zone 2 contained amino acid residues 202, 205, 207, 208, 213 and 217 (Figure 2). The middle- and lower-scoring SDPs were distributed between these two zones indicating the probable involvement of the entire superdomain II/IV in the stop codon decoding.

An interesting feature of zone 2 is the profound amino acid differences between the RF1 and RF2 SDPs. RF1s contain the ‘RS’ motif in contrast to the ‘WG’ motif of RF2s (Table 1). Obviously, arginine and serine residues are chemically very different from tryptophan and glycine residues, respectively. Surprisingly, in the RF2 subfamily, the conserved hydrophobic tryptophan residue was located on the protein surface (Figure 3C).

### Table 1. Predicted SDPs in the RF subfamilies

| Position of SDPs (E.coli RF numbering) | Amino acid residues | Z-score | Human mRF1 |
|--------------------------------------|---------------------|---------|-------------|
| 143                   | Leu or Ile         | 27.83   | Gin         |
| 144                   | Phe                | 27.98   | Trp         |
| 201                   | Val                | 30.77   | Val         |
| 202                   | Gin                | 28.45   | Gin         |
| 205                   | Pro                | 38.32   | Pro         |
| 207                   | Thr                | 27.43   | Thr         |
| 208                   | Glu                | 22.90   | Glu         |
| 213                   | Ile or Val         | 22.74   | Ile         |
| 217                   | Ala or Thr         | 27.51   | Thr         |
| 319                   | Arg                | 38.56   | Gln         |
| 320                   | Ser                | 40.32   | Ser         |
| 322                   | Arg                | 27.00   | Arg         |
| 325                   | Thr                | 25.23   | Thr         |
| 327                   | Asn                | 25.85   | Asn         |
| 334                   | Thr                | 23.54   | Thr         |

*aCalculated as described by Kalinina et al. (15). The highest-scoring SDPs are highlighted.

*bThe amino acids of human mRF1 different from RF1 SDPs are shaded.

*cIdentified earlier (12) as parts of ‘protein anticodon’ that determines RF1/RF2 stop codon specificity towards purines.
on RFs crystal structures. One IR cluster contained the functionally important GGQ loop (25–28) in domain III (Figures 2 and 3). For detecting IRs putatively involved in the stop codon recognition, we searched for IRs clustered with SDPs. Five IRs, namely Arg200, Arg203, Thr215, Arg324 and Tyr326, were located 3–7 Å from SDPs (according to centroids of these amino acid residues) (Table 2). These residues could be involved in RNA–protein interactions via H-bond formation, electrostatic interactions and/or stacking in both RF1 (1r0x) and RF2 (1gqe) crystal structures (Figure 3). The arginine was the most frequent residue found in RNA–protein interacting sites (29,30). The presence of invariant arginines

Figure 1. Multiple alignment of the RF1/RF2 family. Subset taken from MSA of RF1s and RF2s from eubacterial genomes is presented: gi|15604741 (RF1, Chlamydia trachomatis), gi|33235957 (RF1, Chlamydophila pneumoniae), gi|3322309 (RF1, Treponema pallidum), gi|45656013 (RF1, Leptospira interrogans), gi|16273461 (RF1, Haemophilus influenzae), gi|45546433 (RF2, Rubrobacter xylanophilus), gi|49236241 (RF2, Moorella thermoacetica), gi|3322871 (RF2, Treponema pallidum), gi|33236657 (RF2, Prochlorococcus marinus) and gi|33866841 (RF2, Synechococcus sp.). The RF1s are shown as ‘1gi...’ and RF2s as ‘2gi...’. Accession numbers and gi-identificators are shown at the left. The alignment is highlighted according to the conservancy level of this MSA subset: RF1-conserved positions are red, RF2-conserved are green and RF1 + RF2 conserved are blue. SDPs are marked above the alignment by black boxes.
Figure 2. Location of putatively important amino acid residues in RF2 crystal structure. Ribbon models of RF2 3D structure. (A) SDPs (blue), IRs (violet) and RSPs (green). (B) The Cα-atoms of the highest-scoring (blue) and middle-scoring (gray) SDPs. (C) The Cα-atoms of IRs (violet).

Figure 3. SDP/IR areas in the 3D structures of RF1 and RF2. Ribbon (A and C) and surface (B and D) models of RF1 (A and B) and RF2 (C and D) based on 1r0x and 1gqe crystal structures. (A and C) The Cα-atoms of the highest-scoring SDPs (blue) and IRs clustered with SDPs (violet). The same color code is used for surface presentation (B and D).
Amino acid positions in *E. coli* (Figure 3). Zone 1 contains position 205 (numbering as in (12–14) is not the only region involved in the stop codon recognition. Two tripeptides (PxT and SPF motif) are involved in discrimination of the stop codon but we believe that all of them may contribute for high fidelity of stop codon discrimination (31).

The previously described PxT/SPF ‘tripetide anticodon’ represents an example of differences in mutation rates for position 206 in RF1s (variable X) and RF2s (invariant P). This may imply that the selection pressure at this position is different in these subfamilies. This RS may indicate sites, in which functional constraints have changed in the course of evolution and differ in branches of the phylogenetic tree. Since the SDPpred method is not suitable to uncover motifs of this kind, we have applied the DIVERGE approach (17). In contrast to MSA-dependent cutoff of Z-scores, estimated by the SDPpred program, the DIVERGE program utilizes a user-defined threshold of ‘probability score’ for putatively important residues. We set this threshold to 0.99 and obtained a list of 22 potentially significant residues. These residues correspond to sites of the highest differences in mutation rates between RF1 and RF2 subfamilies. The best ‘probability score’ (1.00) was reached by only one amino acid residue and as anticipated, it was position 206. Positions with lower scores were spread over entire superdomain II/IV (Figure 2). Generally, SDPpred searched for positions that are conserved in each subfamily but differ between them. DIVERGE, in contrast, searches for positions that are, for example, conserved in one subfamily and variable in the other. It seems that the latter approach is somewhat less specific. Still, about a half of RSPs occur within or close to spatial zones defined by SDPs and IRs. Therefore, both methods point to the superdomain II/IV as the discriminator area in stop codon decoding.

The SDPs of zone 1 are within 3–7 Å distance from IRsArg200, Arg203, Thr215, Arg325 and Tyr326 (Table 2 and Figure 3). Zone 1 contains position 205 (numbering as in *E. coli* RF2) identified earlier as the first residue of the ‘protein anticodon’. Position 207 from the same ‘protein anticodon’ is among middle-scoring SDPs (Table 1). As visualized by cryo-EM (23,24) this region is located in the decoding centre of the ribosome. These data are consistent with hydroxyl radical mapping of the *E. coli* RF2 SPF motif onto small ribosomal subunit (34). Although the SDPs are spread along two domains of RF1/RF2 (Table 1 and Figure 1), nevertheless, they are clustered in space (Figure 2). Properties of zone 1 are consistent with the suggestion that it forms an ‘extended’ stop codon recognition site which includes the ‘protein anticodon’ as an essential part. Probably, not all SDPs located in the vicinity of zone 1 are in direct contact with the second and/or third positions of the stop codon but we believe that all of them may contribute for high fidelity of stop codon discrimination (31).

The current method of filtering the evolutionary signal among families of orthologous proteins by comparing paralogous protein subfamilies differing in ligand-binding specificity, SDPpred, has been developed and successfully applied to several protein subfamilies (15). This approach permits identification of RF1- and RF2-specific positions in the RF family because of their structural and functional resemblance. Application of the SDPpred to RF protein subfamilies reveals that all SDPs (Table 1) are located in the superdomain II/IV forming an SDP-enriched area (Figure 2). We divided all SDPs into groups according to their Z-scores. Three SDPs with the highest Z-scores map to two distinct spatial zones in domains II and IV (in both RF1 and RF2 3D structures) (Figure 3). Zone 1 contains position 205 (numbering as in *E. coli* RF2) identified earlier as the first residue of the ‘protein anticodon’. Invariant amino acids (IRs) in the RF family are shown in Table 2. Invariant amino acids (IRs) in the RF family are shown in Table 2. IRs clustered with SDPs are highlighted.
machinery. Some data argue in favor of this assumption. Both zones are located in the same superdomain and oriented toward the decoding centre of the ribosome at the small ribosomal subunit (23,24). Most of the SDPs and IRs clustered with SDPs are located at the surface of the RF2 and RF1 crystal structures (Figure 3). The solvent accessibility of SDP/IR clusters reveals that, most probably, these amino acid residues are unshielded and accessible for interaction with stop codon and/or rRNA.

It has been proposed from genetic (35) and biochemical (36) data that RF1/2 binds to the bacterial ribosome in a two-step fashion; at the first step the binding is weak, whereas at the second, codon-specific step, it is strong. For eRF1, a two-step binding model is supported experimentally (37). If so, one may speculate that at the initial step of the RF1/2 binding to the ribosome, distal SDPs in domain IV play an essential role, probably interacting with nucleotides of rRNA or mRNA in the decoding site (Figure 4). The first step induces a conformational change in the RF1/2–ribosome complex. The only proved structural change in RF1/2 is the relocation of the domain III, shown by cryo-EM (23,24). It is not unlikely that after the zone 2 binding at the first step, some unfolding of the RF occurs. This could lead to direct specific contact of zone 1 with the stop signal in mRNA.

Experimental data available so far are consistent with the proposed functional role for two zones of the SDP/IR area. After random mutagenesis nine mutations of Salmonella thyphimurium RF2 have been selected (35). These mutations alter the RF2 specificity towards stop codons in vivo and enable it to recognize the RF1-specific UAG codon. Four of nine point mutations were located in zones 1 and 2 of the SDP/IR area.

The essential role of selected SDPs in the stop codon recognition follows also from the data on mRF1s. Vertebrate mRF1s differ from both bacterial and eukaryotic RFs in their stop codon responses as mentioned in the Results (3,5,38). In parallel, SDPs in human mRF1 and bacterial RF1/RF2s differ in their patterns (Table 1). Unfortunately, at present, the set of mRF1 sequences from evolutionarily distant organisms is not sufficient for systematic analysis of this group of class 1 RFs by SDPred with good statistical validity.

Domains I and III are implicated in functions common for both factors, e.g. interactions with RF3 and the ribosomal subparticles including the peptidyl-transferase centre. This explains why these domains contain no SDPs but exhibit several IRs.

Only five IRs are clustered with SDPs (Table 2) while others are located in different regions of RF with only one IR cluster in domain III including GGQ motif, shown to be functionally essential both for eukaryotes (25–27) and prokaryotes (28,39).

In conclusion, based on bioinformatics approach we have shown in accordance with earlier biochemical and genetic data that superdomain II/IV in RF family is implicated in specific stop codon discrimination. We excluded participation of domains I and III in this process. We predict that 'stop codon discriminator' site is 3D and composed of two distinct regions. We have predicted the involvement of invariant amino acids clustered with SDPs in recognition of the first base of all stop codons. We assume that stop codon discrimination is a two-step process which is accompanied by a conformational change in superdomain II/IV or alteration of mutual orientation of RF and the stop codon. Finally, we predict that positions 319 and 320 (numbering as in E.coli Figure 4. Hypothetical model of prokaryotic class 1 release factor stop codon decoding. Step 1: initial binding of the RF to the pretermination complex. The zone 2 amino acid residues are proposed to be essential for this step. Step 2: The conformational change of RF makes it possible for zone 1 residues to decode specifically the stop codon (UAA is given as an example). The specific binding provokes the signal transduction to the ribosomal peptidyl-transferase center and the release of the nascent polypeptide. Zone 1 (brown), zone 2 (black) and GGQ-containing IR cluster (blue) are schematically represented as circles on the RF structure.
RF2) in RF family are functionally essential. Obviously, our predictions have to be tested in future experiments.

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