Structural Aspects of Ribosomal RNA Recognition by Ribosomal Proteins

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Abstract—This review is focused on the structural aspects of interaction between ribosomal proteins and ribosomal RNA in bacterial ribosomes and complexes of ribosomal proteins with specific fragments of ribosomal RNA. Special attention is given to the recognition of specific spatial architecture of the double-stranded ribosomal RNA by ribosomal proteins and to the role of unstructured protein regions in stabilization of distant ribosomal RNA segments.

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Protein biosynthesis in cells occurs on special cellular organelles – ribosomes. Ribosomes are complex macromolecular machines consisting of several ribosomal RNAs (rRNAs) and over 50 ribosomal proteins (r-proteins) in bacteria (Fig. 1) and up to 80 proteins in eukaryotes [1]. Protein biosynthesis is one of the key cellular processes, and its understanding requires knowing ribosome organization (structure) and interactions between its components. The major break in structural studies of ribosomes happened at the turn of the 21st century, when development of X-ray crystallography and cryo-electron microscopy (cryo-EM) methods allowed to solve the structures of entire bacterial ribosomes and their subunits at a resolution sufficient for detailed description of all ribosomal components.

The structures of isolated r-proteins and their complexes with specific RNA fragments had been determined even before elucidation of the structure of ribosomes and their subunits. The results of the structural studies of the RNA–protein complexes were the first experimental data describing protein interactions with double-stranded RNA and became an important milestone on the pathway to understanding the principles of RNA–protein recognition. Moreover, structures of isolated r-proteins and RNA–protein complexes have served as a basis for interpretation of first electron density maps of ribosomal subunits and their cryo-EM models. Structures of the rRNA–protein complexes have been very useful for deciphering the architecture of functional sites of the 50S (large) ribosomal subunit (LRS), since these sites are not visible in the electron density maps because of their high flexibility. Further investigations of the structure of ribosomal subunits and ribosomes allowed to classify in detail the structures of r-proteins [2].

The majority of r-proteins is conserved and could be found in ribosomes of living organisms from all three domains: bacteria, archaea, and eukaryotes [3, 4]. Comparison of amino acid sequences of r-proteins from different organisms showed that they typically have similar size and do not include nonconserved structural elements [5]. In 2014, a common classification of r-proteins was proposed, that since then has received a general acceptance [6]. Many conserved r-proteins are involved in the formation of functional sites in the ribosome and in initial stages of ribosome assembly. In Escherichia coli, primary RNA-binding r-proteins are S4, S7, S8, S15, S17, and S20 of the 30S (small) ribosomal subunit (SRS) [7, 8] and L1, L2, L3, L4, L9, L20, L23, and L24 of the LRS [9, 10]. During in vitro assembly of ribosomal subunits, binding of these proteins to the corresponding rRNAs leads to a significant compaction of ribonucleoprotein particles [11, 12]. According to the new nomenclature, these proteins are referred to as uS4, uS7, uS8,
uS15, uS17, bS20 and uL1, uL2, uL3, uL4, bL9, bL20, 
uL23, uL24, respectively (here, “u” stands for universally 
conserved r-proteins, whereas “b” denotes proteins spe-
cific to bacterial ribosomes [6]).

In this review, we discuss the structural aspects of the 
interaction between conserved primary binding r-proteins 
and rRNAs. When possible, structures of bacterial r-pro-
teins in the ribosome are compared to those of the isolat-
ed r-proteins or r-proteins complexed with the corre-
sponding rRNA fragments. The “old” nomenclature is 
used for designation of bacterial r-proteins [1].

RIBOSOMAL PROTEINS OF THE L12 STALK
OF THE LRS

Primary binding proteins of the LRS associate with 
different 23S rRNA domains and are situated on both the 
inner (at the intersubunit interface) and outer (solvent 
side) surfaces of the LRS subunit (Fig. 2) [13]. Primary 
binding r-protein L1 forms distinct morphological ele-
ment of the LRS, the so-called L1 protuberance.

On the opposite side of the LRS there is another 
protuberance – the so-called L12 stalk (P stalk in archaea and eukaryotes) formed by the r-proteins L10 
and L11 (uL10 and uL11, according to the new nomen-
clature, respectively) and several dimers of L12 (bL12). 
The L12 stalk of the LRS plays an essential role in the 
ribosome interactions with translation factors and in 
control of translation fidelity by forming ribosome 
GTPase site with the 23S rRNA [14, 15]. The L12 stalk r-
proteins were among the first, whose structures were 
determined by X-ray crystallography and NMR analysis 
in either isolated form or in complexes with rRNA frag-
ments [16, 17].

Ribosomal protein L11. L11 consists of two globular 
domains connected by a short flexible linker [17]. The C-
terminal domain (75 amino acid residues (a.a.) interacts 
with the 23S rRNA domain II (region 1051-1108, E. coli 
numbering) [18]. The N-terminal domain interacts with 
translation factors on the ribosome and does not affect 
the specificity of the 23S rRNA binding by the protein 
[19, 20].

In 1990s, X-ray crystallography was used to solve the 
structures of the C-terminal domain of L11 from 
Geobacillus stearothermophilus complexed with an E. coli 
23S rRNA fragment including helices H43 and H44 [21] 
(Fig. 3a) and of the full-size L11 from Thermotoga mariti-
ma bound to the same 23S rRNA fragment (Fig. 3b) [20]. 
Somewhat later, the structure of L11 complexed with the 
23S rRNA and antibiotic thiostrepton was determined by 
NMR analysis [19].

The N-terminal domain of L11 consists of a three-
stranded β-sheet with two α-helices located on one of its 
sides. According to the Structural Classification of 
Proteins (SCOP) [22], this α+β domain structure is 
unique. The C-terminal domain of L11 consists of three 
α-helices with a short β-hairpin and two long loops, 
which are disordered in the absence of rRNA [20]. 
According to the SCOP, this domain has the DNA/RNA-
binding three-helical bundle fold and belongs to the all-
alpha protein class.
In all the determined structures, L11 protein binds the 23S rRNA via the C-terminal helix $\alpha_5$, that is positioned alongside the minor groove of the rRNA and forms the majority of contacts with it (Fig. 3). The area of contact is shielded by residues of the helix $\alpha_3$ and loops $\alpha_3-\beta_4$ and $\alpha_4-\alpha_5$, which are situated on both sides of the helix [20]. The majority of L11–rRNA contacts are hydrogen bonds between atoms of the protein main chain and the rRNA sugar-phosphate backbone, although there are also a number of contacts between the protein and the rRNA nucleotide bases [21]. Similar mode of recognition via $\alpha$-helix is shared by many DNA-interacting transcription factors (e.g., Cro repressor) [23, 24]. In their cases, protein $\alpha$-helix is located in the major groove of B-DNA, where atoms of amino acid residues interact with atoms of DNA bases, thereby providing specific recognition of the
DNA sequence by the protein [25]. Narrow major groove in the A-form double-stranded RNA does not allow such recognition; therefore, it was assumed that L11 recognizes a spatial fold of the rRNA rather than its nucleotide sequence [20, 21].

**Ribosomal protein L10.** L10 is an important element of the L12 stalk. It acts as a bridge between L12 and the 23S rRNA [15, 26]. L10 consists of two domains. The N-terminal α/β domain has a compact three-layer structure and interacts with a region of the 23S rRNA helices H42-H44 close to L11 [27]. The C-terminal domain is a long α-helix that binds the L12 dimers [15] (Fig. 4). The two domains of L10 are connected by a flexible linker; for this reason, its C-terminal domain with bound L12 dimers is extremely mobile and cannot be seen in the ribosome crystal structures. The structural features of the ribosomal L12 stalk have been described based on the cryo-EM reconstruction of ribosomes. In 2005, the structure of the L10 protein bound to three L12 dimers from the bacterium *T. maritima* was used to create the most interesting structural model of this LRS functional domain in the archaeon *Haloarcula marismortui* [15].

Recently obtained high-resolution structures of bacterial ribosomes from *E. coli* [28, 29] and *Thermus thermophilus* [30, 31] allow to analyze the contacts between the N-terminal domain of L10 and the 23S rRNA in detail. Interaction of L10 with the 23S rRNA is realized through amino acid residues of protein helices α1 and α3, which are located at the rRNA minor groove in the region of nucleotides 1038-1117 in *E. coli* 23S rRNA (Fig. 4). Amino acid residues of L10 in all the available structures do not form hydrogen bonds with atoms of RNA nucleotide bases, but only with the RNA sugar-phosphate backbone. Therefore, L10 recognizes specific spatial rRNA arrangements.

**L1 STALK OF THE LRS**

**Ribosomal protein L1.** The L1 stalk of the LRS is formed by the r-protein L1 and helices H76-H78 of the 23S rRNA. It is located on the opposite side of the LRS from the L12 stalk (Fig. 2). L1 could not be seen on the X-ray structures of ribosomes due to the high mobility of the L1 stalk, which can move a considerable distance and rotate around the axis of the 23S rRNA helix H76 depending on the presence of tRNA in the ribosomal E site [13, 32]. It is believed that the mobility of this region is required for its participation in the tRNA translocation on the ribosome, particularly, in the relocation of deacylated tRNA to the E site with subsequent removal of tRNA from the ribosome [33]. Due to the high mobility of the L1 stalk, the pioneer works on the crystallography of ribosomes had been able to describe only a part of the L1 structure that was a fragment of the 23S rRNA without the protein itself [34, 35]. This gap in the understanding of the ribosome structure was filled by determining the structure of L1 in a complex with the corresponding 23S rRNA fragment.

In 1996, the structure of L1 from the thermophilic bacterium *T. thermophilus* was determined for the first time in our laboratory [36]. It was shown that the protein consists of two domains. The first domain includes both...
the N-terminal and C-terminal parts of the protein, whereas the second domain consists of the middle component of the polypeptide chain (Fig. 5a). The first domain has the two-layer split $\beta$-$\alpha$-$\beta$ structure; the second domain has the three-layer Rossman fold. The domains are connected with a flexible linker and can change mutual orientation in solution [37].

In 2003, we solved the structure of L1 from the archaeon Sulfolobus solfataricus in a complex with a fragment of the 23S rRNA from *T. thermophilus* (Fig. 5b) [38]. Such combination of components from different organisms allowed us to crystallize the complex after almost seven years of searching for proper crystallization conditions after the structure of the isolated L1 had been determined. The obtained structure was used by us and other researchers for describing the L1 stalk in bacterial ribosome models [38].

Fig. 5. a) Structure of L1 from *T. thermophilus* [35]; b) structure of L1 from *S. solfataricus* in a complex with a fragment of the 23S rRNA from *T. thermophilus* (PDB 1MZP).

The structure of the L1–23S rRNA complex has revealed that L1 interacts with RNA through two distant areas located on the surfaces of both domains. The first domain contains four times as many RNA-contacting residues as the second domain, which suggested that it plays the major in the recognition of the 23S rRNA [38]. Similarly to the ribosomal complexes described above, amino acid residues of L1 interact predominantly with atoms of the rRNA sugar-phosphate backbone.

23S rRNA-BINDING RIBOSOMAL PROTEINS OF THE LRS BODY

Ribosomal protein L2. The primary 23S rRNA-binding L2 protein is located close to the base of the L1 stalk in the LRS (Fig. 2) [39]. According to the available data, L2 is required for the association of SRS and LRS and formation of the 70S ribosome, tRNA binding by the ribosome at the A and P sites, and peptide bond formation in the nascent polypeptide [40, 41]. In 70S ribosomes of *E. coli*, L2 forms several contacts with the 16S rRNA during translation initiation stage with the formation of the bridge B7b, which is then disrupted upon binding of the translation factor EF-G to the ribosome [42]. Bacterial L2 protein can be substituted in the ribosomes by its homologs from the archaeon *H. marismortui* (HmaL2) or the L8 protein from the human 60S cytoplasmic ribosomal subunit without loss in the ribosome activity [43].

Most part of the 270-a.a. L2 protein does not possess an apparent secondary structure. However, there are two $\beta$-structured domains in the central part of the protein molecule that might be classified as the nucleotide-binding OB fold and SH3-like domain (Fig. 6).

As a part of the LRS, L2 interacts with distantly located regions of the 23S rRNA, thereby playing a role of a scaffold protein and connecting 23S rRNA helices (Fig. 6). Analysis of the protein–rRNA contacts shows that only 15% of these interactions are hydrogen bonds between the protein atoms and atoms of the rRNA nucleotide bases, whereas the rest are the contacts with the 23S rRNA sugar-phosphate backbone. An interesting feature of the RNA–protein interactions is that the vast majority of them involve amino acid residues situated in disordered protein regions (loops and turns). Hence, it is difficult to believe that the tertiary structure of L2 provides recognition of the 23S rRNA, rather than secures certain spatial arrangement of the 23S rRNA helices during LRS assembly.
Ribosomal protein L3. L3 is one of the two initiator proteins (together with L24) in the LRS assembly [44]. It binds to the 3'-end of the 23S rRNA and nucleates rRNA folding [10]. L3 is involved in the formation of the peptidyl transferase site due to the presence of elongated loops that penetrate the LRS [45]. Mutations in the amino acid sequence of L3 might result in the resistance to the antibiotics tiamulin [46, 47] and linezolid [47]. All the amino acid substitutions in this protein were found in the fragment 144-151. The strongest effects were caused by substitutions of the residues 148 and 149 located in the peptidyl transferase site of the ribosome [46].

No structures of the isolated protein and L3 in a complex with a 23S rRNA fragment have been solved. Examination of the L3 structure in the *E. coli* ribosome shows that L3 is a two-domain protein (Fig. 7). The first domain includes both the N- and C-termini of the protein and represents a β-barrel with a protruding loop of 50 a.a. (residues 115-165). The loop does not form any secondary structure with the exception of a small α-helix and a short β-region. The second domain (residues 30-97) is inserted into the first domain and consists of a β-sheet with few α-helices located on one side of it.

Fig. 6. Left, position of L2 on the 23S rRNA in *E. coli* 70S ribosome (PDB 5IT8). Right, schematic representation of L2 in the ribosome.

Fig. 7. Left, position of L3 on the 23S rRNA in *E. coli* 70S ribosome (PDB 5IT8). Right, schematic representation of L3 protein in the ribosome.
Most of the contacts (approximately 80%) between L3 and the 23S rRNA are made by residues 115-165 of the protein loop; some of them are hydrogen bonds with atoms of rRNA nucleotide bases. The β-strands of the second domain interact with the rRNA sugar-phosphate backbone, whereas the first domain binds rRNA through only five amino acid residues. Therefore, the primary rRNA-recognizing unit in L3 is the second domain that can identify the structure of the 23S rRNA during initiation of the LRS assembly. The loop 115-165 makes the majority of contacts with the rRNA, most probably, during subsequent stages of the LRS assembly.

Ribosomal protein L4. L4 is involved in the early stages of the LRS assembly [10]. It binds to the 5′-terminus of the 23S rRNA and forms multiple contacts with different domains of the 23S rRNA in the assembled LRS [48, 49] (Fig. 8). L4 is located at the solvent side of the LRS. However, it possesses an elongated loop that penetrates the entire subunit toward the peptidyl transferase site of the ribosome. This loop has attracted a lot of interest, since it reaches the ribosomal tunnel through which nascent polypeptide passes after its synthesis on the ribosome [50]. Mutations in this loop might make bacteria resistant to macrolide antibiotics [51, 52]. Though the loop of L4 penetrated the entire LRS, it is dispensable for the LRS assembly and might be deleted [53].

The structure of the isolated L4 from T. maritima was determined in 2000 [54] except the fragment of 41-96 that corresponds to the long loop, which is disordered if the protein is not bound to rRNA (Fig. 8). Because this region possesses high mobility, it is not fixed in the crystal and has poor electron density, which prevents solving its structure.

According to the SCOP, L4 belongs to the three-layer α/β proteins and contains a four-strand parallel β-sheet. The overall fold of its secondary structure elements is unique and was classified as a separate family of L4 r-proteins (SCOP 52167).

Analysis of contacts between amino acid residues of L4 and the 23S rRNA shows that (similarly to L3), the majority of them are made by residues of the long protein loop and adjacent regions of α-helices (a.a. 40-99). The second protein region involved in the RNA–protein interactions is the helix α2. Since it was shown that the loop in L4 protein is not required for the LRS assembly [53], it is reasonable to assume that the rRNA-recognizing unit of L4 is its structured domain, whose most important element is the helix α2.

Ribosomal protein L9. L9 is one of the earliest proteins to bind to the 23S rRNA [10]. It interacts with the rRNA 3′-terminus next to the L1-binding site. L9 was found only in bacterial ribosomes [55]. In the LRS, it is located directly under the L1 stalk (Fig. 9).

The structure of L9 was first published in 1994 [55]. L9 consists of two domains: small N-terminal domain (47 a.a.) with the three-layer α/β/α architecture and C-terminal domain (85 a.a.) with the double-layer α+β architecture connected to a long α-helix (37 a.a., 10 turns) that participates in the formation of the domains. The N-terminal domain is structurally similar to the RNA recognition motif (RRM). Both domains are positively charged, so it was assumed that they both interact with the 23S rRNA [55, 56]. However, according to the X-ray structural data, only the N-terminal domain interacts with the 23S rRNA in bacterial ribosomes, whereas the C-ter-

**Fig. 8.** Left, position of L4 on the 23S rRNA in E. coli 70S ribosome (PDB 5IT8). Central panel, structure of L4 in the ribosome. Right, structure of T. maritima L4 (PDB 1DMG). Residues 41 and 96 are the boundaries of the non-described part of the protein.
minal domain is distant from the ribosome surface and contacts the SRS of neighboring ribosome in a crystal [57]. Cryo-EM of the ribosomes has demonstrated that the α-helix of the L9 protein can bend so that the C-terminal domain approaches the r-protein S6 [58]. One of the hypotheses that explain L9 location in the ribosome is that it controls the distance between ribosomes in polysomes and suppresses translation frameshifting, thereby increasing the fidelity of mRNA translation by the ribosome [59].

Fig. 9. Left, position of L9 on the 23S rRNA in *E. coli* 70S ribosome (PDB 5IT8); view from the intersubunit interface. Right, structure of L9 from *G. stearothermophilus* (PDB 1DIV).

Fig. 10. Left, position of L20 on the 23S rRNA in *E. coli* 70S ribosome (PDB 5IT8); view from the solvent side. Right, zoomed view of the L20-binding site.
resolution of the model and inaccuracy of its refinement at the periphery. However, one may argue that L9 interacts with a region that connects helices H76 and H79 in the 23S rRNA.

**Ribosomal protein L20.** L20 binds to the 5′-terminus of the 23S rRNA early in the process of ribosome assembly and is required for the first stage of the LRS assembly in vitro [60, 61]. L20 is not universally conserved and presents in bacterial ribosomes only. However, deletion of the gene encoding L20 protein in *E. coli* is lethal [61]. It should be noted that L20 can substitute for the L24 protein during initiation of the LRS assembly at low temperatures [62]. It is located on the opposite side of the LRS from the peptidyl transferase site and interacts with several distant regions of the 23S rRNA (Fig. 10).

NMR studies of L20 showed that L20 stays unfolded when not bound to the rRNA [63]. However, it acquires a secondary structure during crystallization [64]. According to SCOP classification, L20 is a single-domain protein containing four orthogonally packed α-helices. Packaging of the helices corresponds to the PAPB domain.

It is believed that L20 becomes structured upon interaction with the 23S rRNA. The C-terminal α-helix of the protein interacts with other r-proteins on the surface of ribosomes, whereas the N-terminal α-helix and long α2-helix interact with rRNA [65]. Indeed, analysis of interactions between L20 and the 23S rRNA in ribosomes shows a large area of contact between these helices and the sugar-phosphate backbone of the 23S rRNA helices H40–H41 on one side of the protein and helices H25–H27 on the other side (Fig. 10). However, amino acid residues of helices α3 and α4 also form a number of contacts with the sugar-phosphate backbone of the 23S rRNA helix H41; therefore, the interactions between the L20 C-terminal domain and the 23S RNA cannot be ignored. Analysis of the L20 interaction with rRNA reveals the role of this protein in the LRS assembly: L20 neutralizes negative charges of phosphates groups in the 23S rRNA and brings its distant domains close together [61, 65].

**Ribosomal protein L23.** L23 binds to the third domain of the 23S rRNA at the early stages of the LRS assembly [10, 49]. It is located on the LRS close to the exit of the nascent polypeptide from the outer side of the ribosome (Fig. 2). Some experimental data indicate that L23 interacts with the trigger factor [66] and signal recognition particle [67, 68] on the ribosomes and might be involved in the folding of the growing polypeptide chain [66].

The spatial structure of isolated L23 was determined by the NMR analysis (Fig. 11) [69]. It was found that L23 belongs to the three-layer α+β protein family (SCOP 54189). Its structure is conserved and does not significantly changes upon protein binding to the 23S rRNA.

In the ribosomal subunit, L23 interacts with the sugar-phosphate backbone of the 23S rRNA helices H6, H9, H23 and H53 (Fig. 11) predominantly through a long loop and adjacent termini of the β-strands and through residues of the α2-helix that form contacts with the sugar-phosphate backbone of H53.

**Ribosomal protein L24.** L24, along with the L3 protein, is the initiator of the LRS assembly in bacteria [44, 70]. It interacts with the 23S rRNA 5′-terminus and nucleates the LRS folding [10]. In the *E. coli* LRS, L24 is located close to L23 in the vicinity of the nascent polypeptide exit from the ribosome and interacts with the 23S rRNA helices H7, H18–H20 and H24 (Fig. 12) [70, 71].

The structure of the isolated L24 has not been determined. In ribosomes, it is a β-protein, whose central part

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**Fig. 11.** Left, position of L23 on the 23S rRNA in *E. coli* 70S ribosome (PDB 5IT8). Right, structure of L23 from *T. thermophilus* as determined by NMR analysis (PDB 1N88).
is folded into the SH3-like barrel (SCOP 50105) (Fig. 12). Interestingly, both the N- and C-termini of the L24 protein are located in the same domain; the rest of the polypeptide chain forms distinct domains and elongated loops. Similar architecture is also observed in the L1, L3, L4, and L23 protein reviewed above. It is possible that such spatial arrangement of the secondary structure elements facilitates the formation of a stable protein core, while “extensions” allow these proteins to reach the regions of the 23S rRNA that are distant from the area of the main interaction, thereby promoting correct folding of rRNA during ribosome assembly.

Analysis of contacts between amino acid residues of L24 and the 23S rRNA shows that the protein mainly interacts with atoms of the sugar-phosphate backbone of the double-stranded rRNA regions.

5S rRNA-BINDING RIBOSOMAL PROTEINS OF THE LRS

Along with the large 23S rRNA (2904 nucleotides in *E. coli*), bacterial LRS also contains small 5S rRNA (120 nucleotides) [72]. There are three r-proteins that bind the 5S rRNA: L5, L18 and L25 [73] (uL5, uL18, and bL25, respectively, according to the new nomenclature [6]). Together they form the central protuberance of the LRS that plays the key role in the ribosome functioning (Fig. 2) [74, 75].

**Ribosomal protein L5.** L5 binds to the loop C that covers the 5S rRNA helix III [76]. Its rRNA-binding properties are significantly promoted in the presence of L18 protein on the 5S rRNA [77, 78]. L5 interacts with the 23S rRNA helices H84-H85, thereby providing attachment of the 5S rRNA—protein complex to the LRS (Fig. 13a) [9, 79].

As a part of the 70S ribosome, L5 protein interacts with the S13 protein of the SRS at the translation initiation stage. According to the 70S ribosome dynamics modeling, this interaction is affected by the binding of the translation elongation factor EF-G [42]. Comparison of the two structures of *E. coli* 70S ribosome determined independently by the X-ray analysis at a 3.5 Å resolution confirmed the dynamic nature of the L5 contacts with S13 and S19 [80].

L5 protein interacts with tRNAs in the ribosomal P site and, together with the 5S rRNA and other 5S rRNA-associated proteins, participates in the tRNA positioning in this ribosomal site [78, 80, 81].

The structure of the isolated L5 from *G. stearothermophilus* was determined in 2001 [82]. In 2003, the structure of the complex of L5 from *T. thermophilus* with a 5S rRNA fragment from *E. coli* was published (Fig. 13) [83]. Comparison of the structures of L5 alone, in a complex with the 5S rRNA fragment, and in ribosome demonstrated that, with an exception of short mobile protein fragments (e.g., helix $\alpha_1$, whose residues forms additional contacts with the 5S rRNA in the ribosome), the protein overall structure remains invariant [83].

L5 is an $\alpha+\beta$ protein with the antiparallel $\beta$-sheet and belongs to the RL5-like protein superfamily (SCOP 55282). Packaging of the secondary structure elements in L5 is similar to that in the RNP (ribonucleoprotein) domain [83] that binds RNA through the $\beta$-sheet surface. In the ribosome, L5 protein makes contacts with the 5S rRNA via amino acid residues of the loop $\alpha_1$-$\alpha_2$, strand $\beta_2$, and terminus of $\alpha_4$ helix (Fig. 13e). Therefore, the 5S rRNA adjoins the edge of the $\beta$-sheet. This interaction involves multiple contacts between the side chains of the L5 amino acid residues and atoms of the 5S rRNA nucleotide bases. This could be explained by the fact that

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**Fig. 12.** Left, position of L24 on the 23S rRNA in *E. coli* 70S ribosome (PDB 5IT8). Right, schematic representation of L24 protein in the ribosome.
the site of L5 binding includes not only the double-stranded 5S rRNA region but also the loop C, which is a single-stranded rRNA region exposed to the intermolecular contacts.

L5 protein interacts with the 23S rRNA via the β-sheet surface that is typical for the RNP domains (Fig. 13f) [83]. It should be noted that the number of L5 contacts with the 23S rRNA is somewhat larger than with the 5S rRNA. These two facts correspond well to the data on the importance of L5 for binding of the 5S rRNA-protein complex to the LRS [78, 82].

Ribosomal protein L18. As it was mentioned above, L18 significantly increases the affinity of the L5 protein to the 5S rRNA [77, 78]. It is believed that L5 and L18 bind to the 5S rRNA in a cooperative manner [84]. Both proteins interact with a region of helix III and loop C of the 5S rRNA (Fig. 14a), so the cooperativity of L5 and L18 binding to the 5S rRNA is not surprising.

The structure of the isolated L18 protein from *T. thermophilus* was determined by NMR in 2002 [85]. Two years later, the structure of L18 from *G. stearothermophilus* was also solved [86]. L18 belongs to the three-layer α/β proteins of the RNase H-like type (SCOP 53138). Twenty N-terminal amino acid residues that do not form any secondary structure in a solution were missing from the both structures (Fig. 14). In ribosomes, the N-terminal part of the protein acquires the α-helical structure and interacts with the 23S rRNA domain V, including the H83-H85 helix junction and helix H87 (Fig. 14c). This rRNA region also contacts with the edge of the β4-strand and

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Fig. 13. a) Interaction of L5 with the 5S rRNA and 23S rRNA in *E. coli* 70S ribosome (PDB 5IT8). b) Crystal structure of L5 from *G. stearothermophilus* (PDB ID 1IQ4). c) The interaction site of *E. coli* L5 with the 5S rRNA (nucleotides 26-60) in the ribosome (PDB 5IT8). d) Crystal structure of L5 from *T. thermophilus* in a complex with the 5S rRNA fragment from *E. coli* (PDB ID 1MJ1). Orientation of L5 in the panels (b)-(d) is the same. e) Zoomed view of the region of interaction of L5 and 5S rRNA. f) Zoomed view of the region of interaction of L5 and 23S rRNA.
the adjacent loop. The area of contact between L18 and the 5S rRNA is much broader and includes the β-sheet surface, which interacts with the 5S rRNA helix I, and the terminal parts of all α-helices that interact with the loops B and C of the 5S rRNA helix III (Fig. 14b). By comparing the interactions of proteins L5 and L18 with the 5S rRNA and 23S rRNA, one may see that they are “mirror-like”: L18 forms more contacts with the 5S rRNA and interacts with it via the β-sheet, whereas L5 makes more contacts with the 23S rRNA and also interacts with it via the β-sheet. At the same time, L18 binds to two spatially distant 5S rRNA helices — I and III, which should stabilize their mutual arrangement and the three-dimensional structure of 5S rRNA in the ribosome [87]. These structural features of the L5 and L18 interactions with rRNA correspond well to the previously defined roles of these proteins in the ribosome assembly [75, 78, 84, 85, 88].

Ribosomal protein L25/TL5. The third 5S rRNA-binding protein is L25 (according to E. coli nomenclature). It binds to the 5S rRNA independently of L5 and L18 and is not required for attachment of the 5S rRNA–protein complex to the 23S rRNA [77, 89]. L25 interacts with the 5S rRNA loop E region situated between helices IV and V, which is the most distant from the site of L5 and L18 binding (Fig. 2) [73, 90]. In contrast to L5 and L18, L25 is not required for the survival of bacterial cells [73, 79]. Its function in the ribosome is not clear, although it was suggested that L25 contributes to the stabilization and proper operation of such functional centers of bacterial LRS as the A and P sites [73, 91].

The structure of the isolated L25 from E. coli was determined by the NMR (Fig. 15) [92]; the structure of L25 bound to the 5S rRNA fragment was solved using X-ray crystallography [93]. According to the SCOP classification, L25 protein belongs to a family of β-proteins with
the β-barrel topology (SCOP 50714), although it also contains three α-helices. In the L25–5S rRNA complex, the protein interacts with the sugar-phosphate backbone of the loop E and recognizes its specific spatial structure [93] via amino acid residues of the β-sheet surface and the α1 helix (Fig. 15).

One year later, the structure of the complex of a 5S rRNA fragment with the TL5 protein from T. thermophilus (Fig. 15d) was determined [94]. The N-terminal domain of TL5 is homologous to the E. coli L25 protein. It was demonstrated that TL5 can functionally substitute L25 in the reconstituted E. coli ribosomes [95]. In the complex, only the N-terminal domain, but not the C-terminal domain, interacts with the 5S rRNA fragment. An interesting feature of the TL5 N-terminal domain is that the helix α1 has less turns as compared to E. coli L25 and almost does not form contacts with the 5S rRNA. Therefore, the recognition of the RNA fragment by the TL5 protein is provided exclusively by the amino acid residues of the β-sheet surface of the N-terminal domain.

Fig. 15. a) Interaction of L25 with the 5S rRNA in E. coli 70S ribosome (PDB 5IT8); b) NMR structure of E. coli L25 (PDB 1B75); c) crystal structure of E. coli L25 in a complex with the 5S rRNA fragment (PDB 1DFU); d) crystal structure of T. thermophilus TL5 in a complex with the 5S rRNA fragment (PDB 1FEU).
16S rRNA-BINDING RIBOSOMAL PROTEINS OF THE SRS

The 16S rRNA is almost twice as short as the 23S rRNA (1542 and 2904 nt in E. coli ribosomes, respectively) [96]. The number of primary 16S rRNA-binding r-proteins is also smaller — S4, S7, S8, S15, S17, and S20 (Fig. 16).

Ribosomal protein S4. S4 is one of the two initiator proteins in the SRS assembly [97]. It binds to the 16S rRNA and nucleates folding of the SRS body [97-99]. The binding site for S4 is situated at a junction of five helices (H3, H4, H16, H17 and H18) at the beginning of the 16S rRNA 5′-domain. The protein interacts with the sugar-phosphate backbones of three of these helices (Fig. 17) and also directly with the 5′-terminus of the 16S rRNA and nucleotides of the helix H21 terminal loop [97, 100]. Such pivotal location of the S4 protein correlates well with the biochemical data on the chemical modifications of nitrogenous bases and protection of the 16S rRNA sugar-phosphate backbone [101].

The structure of the isolated S4 from G. stearothermophilus was determined by X-ray crystallography [102] and NMR [103] (Fig. 17). In both cases, the protein lacked 41 N-terminal amino acid residues, as they are disordered in a solution. In the SRS, the N-terminal part of the protein is partially structured and has two small α-helices.

S4 consists of two domains. Again, the second domain is an insert into the first one, which is typical for some r-proteins. The first domain is entirely α-helical; the second one is an α+β domain that consists of a five-strand antiparallel β-sheet with three α-helices located on one of its sides. The spatial arrangement of the secondary structure in the second domain corresponds to the α-L RNA-binding motif (SCOP 55174).

The 16S rRNA interacts with amino acid residues of four α-helices of both domains and two α-helical regions of the unstructured N-terminal part of the protein. The contacts are formed by atoms of the S4 amino acid residues and atoms of the 16S rRNA sugar-phosphate backbone and also some atoms of its nitrogenous bases. Amino acid residues of the β-sheet do not participate in the interaction with the 16S rRNA.

Ribosomal protein S7. S7 is the second initiator protein of SRSs; its binding to the 16S rRNA leads to the assembly of SRS head [97]. The protein is situated at the interface side of the SRS close to the SRS decoding center, where it interacts with the mRNA [104]. There are data that S7 contacts tRNA in the P and E sites [105].

The S7-binding site in the 16S rRNA includes the junction of 16S rRNA helices H28, H29, and H43 and nearby helix H41 (Fig. 18a). The N-terminal amino acid residues of the protein form single contacts with the helix H37 loop in E. coli ribosomes (PDB 5IT8). S7 interacts with the 16S rRNA sugar-phosphate backbone predominantly via amino acid residues of α-helices. The long N-terminus of the protein is arranged along the groove of the RNA helix H43 and forms the surface of contact with the RNA. The small β-hairpin in S7 forms only a couple of contacts with the 16S rRNA. In general, the interaction of S7 with the 16S rRNA is unique: the protein α-helices are not introduced into the grooves of the RNA helices; the β-hairpin is small and does not form a surface for the RNA helix binding. Apparently, the interaction is provided by a complex spatial structure of the 16S rRNA that is specifically recognized by the S7 protein.

The structures of isolated S7 proteins from G. stearothermophilus [106] and T. thermophilus [107] were determined in 1997 (Fig. 18). S7 protein is classified as an all-α protein consisting of five α-helices with an additional α-helix and β-hairpin outside the protein core (SCOP 47974). Comparison of the structures of both S7 proteins in the isolated state and as parts of the ribosome showed that binding to the 16S rRNA does not noticeably change the structure of the proteins.

Ribosomal protein S8. S8 directly interacts with central domain of the 16S rRNA and coordinates assembly of the SRS platform with participation of the proteins S5 and S12 [108, 109]. In the ribosome, it interacts with the conserved region of helix H21 by binding to the sugar-phosphate backbone of double-stranded RNA. It also interacts with the adjacent helix H20 and forms extensive
contacts with the 16S rRNA helix H25 (Fig. 19a) [101, 110]. Therefore, S8 protein stabilizes the spatial arrangement of this SRS platform region.

It was shown that a small fragment of the 16S rRNA helix H21 is sufficient for the S8 binding [111, 112]. In 2001, the structure of archaeal S8 complexed to the corresponding 16S rRNA fragment from *Methanococcus jannaschii* was published (Fig. 19b) [113] demonstrating that the protein interacts with the RNA minor groove. At the same time, S8 forms a bridge between two turns of the RNA helix by its two domains. It was also established that the majority of contacts are formed between nucleotides and conserved amino acid residues of the C-terminal domain [113].

In 2014, these conclusions were confirmed by the structural studies of the complexes of S8 from *Bacillus anthracis* with RNA aptamers [114]. It was also shown that nucleotides of free RNA aptamers form numerous non-canonical base pairings, whereas interaction with S8 causes these RNAs to acquire the classical S8-recognized fold and also changes interactions between the aptamer nucleotide bases. It was concluded that S8 is able to

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**Fig. 17.** a) Interaction of S4 with the 16S rRNA in *E. coli* SRS (PDB 5IT8); b) S4 structure in *E. coli* ribosome (PDB 5IT8); c) NMR structure of S4 from *G. stearothermophilus* (PDB 1C05).

**Fig. 18.** a) Interaction of S7 with the 16S rRNA in *E. coli* SRS (PDB 5IT8); b) crystal structure of S7 from *G. stearothermophilus* (PDB 1HUS); c) crystal structure of S7 from *T. thermophilus* (PDB 1RSS).
change the secondary and tertiary structures of RNA upon binding [114].

The structure of the isolated S8 was determined in 1996 for *G. stearothermophilus* [115], in 1998 for *T. thermophilus* [116], and in 2012 for *Aquifex aeolicus* [117] (Fig. 19). S8 consists of two domains and belongs to the α+β class of proteins. The arrangement of the secondary structure elements corresponds to the four-layer b/a/b/a structure with two subdomains and, according to SCOP, is characteristic of a separate family (SCOP 56048). It should be noted that S8 from *A. aeolicus* has two additional α-helices in the N-terminal domain (Fig. 19e), which considerably increases its affinity to the region of junction of helices H20, H21, and H22 in the 16S rRNA [117].

**Ribosomal protein S15.** S15 binds to the 16S rRNA in the vicinity of S8 and also participates in the SRS platform assembly [42, 80]. In *E. coli* 70S ribosomes, S15 contacts the 23S rRNA of the LRS, thereby forming the intersubunit bridge B4 [42].

As a part of the SRS, S15 is located along the 16S rRNA helix H22 and forms numerous contacts with its sugar-phosphate backbone (Fig. 20). At the same time, the opposite termini of helices α2 and α3 stabilize two helical junctions formed by helices H20, H21, and H22 and by helices H23a, H22, and H23. As a result, S15 plays a crucial role in the formation of the spatial structure of the 16S rRNA central domain in the region of helix H22 [118, 119].

Analysis of the structure of the S15 protein from *T. thermophilus* in a complex with the 16S rRNA fragment demonstrated that the protein stabilizes mutual orientation of three 16S rRNA helices – helix H22 with adjacent fragments of helices H20 and H21 – via interactions with their sugar-phosphate backbones (Fig. 20b) [118]. Concurrently, the structure of the complex of larger 16S rRNA with proteins S6, S15, and S18 was published that uncovered the precise mechanism of the 16S rRNA central domain formation in the SRS (Fig. 20c) [119]. Until now, this complex remains the largest rRNA fragment complexed with several r-proteins.

The structures of isolated S15 proteins from *T. thermophilus* (Fig. 20d) [120] and *G. stearothermophilus* were also determined (Fig. 20e) [121]. S15 belongs to the class of all-α proteins. In addition, S15 was assigned to a sepa-
rate family of S15/NS1 RNA-binding proteins with irregular packing of three α-helices and additional N-terminal α-helix (SCOP 47064). Indeed, comparison of S15 structures suggests the mobility of the N-terminal α-helix relatively to other helices of the protein, which is especially obvious in structure of S15 from *G. stearothermophilus*.

**Ribosomal protein S17.** S17 specifically binds to the 5′-terminal domain of the 16S rRNA in the region of helix H11 [108] and stabilizes junction between helices H6, H6a, H7, H11, and H12 [100]. In the SRS, S17 is located close to the primary binding proteins S7 and S15. The loops of this protein reach 16S rRNA fragments that are remote from the helix H11 (Fig. 21). S17 forms single contacts with regions of helices H7, H12, H20, H21, and H44. S17 plays an important role in the control of the translation fidelity, as amino acid changes in this protein confer resistance to the antibiotic neomycin, which, in its turn, reduces mRNA misreading *in vitro* [122, 123].

In 1993, the structure of the isolated S17 was determined by the NMR [124], and S17 was classified as an all-β protein. An unusual incident is associated with this PDB entry: the structure of the protein deposited to the data bank turned out to be a mirror image of the real protein structure.

S17 protein is an all-β protein with the widely occurring supersecondary structure of the OB fold type. Its structure represents a β-barrel that belongs to the cold shock DNA-binding domain-like family (SCOP 50282). Despite

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**Fig. 20.** a) Interaction of S15 with the 16S rRNA in *E. coli* SRS (PDB 5IT8); b) crystal structure of S15 from *T. thermophilus* in a complex with the 16S rRNA (PDB 1DK1); c) crystal structure of the complex of *T. thermophilus* r-proteins S6, S15, and S18 with the 16S rRNA fragment (PDB 1G1X); d) NMR structure of S15 from *T. thermophilus* (PDB 1AB3); e) crystal structure of S15 from *G. stearothermophilus* (PDB 1A32).
its resemblance to proteins of this class, virtually all contacts
with the 16S rRNA are formed by amino acid residues of the
loops between the $\beta$-strands, whereas residues of the $\beta$-
sheet form only a small number of contacts with the RNA.

**Ribosomal protein S20.** S20 was found only in bacteri-
al ribosomes [6]. It binds to the 16S rRNA 5'-terminal
region independently of other bacterial r-proteins [7, 8].
Furthermore, there are two alternative sites of initial pro-
tein binding to the rRNA [125]. There are data suggesting
that S20 stabilizes the region of helices H6, H7, H8, H9,
H11, and H13 [100]. In the ribosome, the protein is locat-
ed between the above helices and contacts functionally,

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**Fig. 21.** Left, interaction of S17 with helix H11 of the 16S rRNA in *E. coli* SRS (PDB 5IT8). Right, region of binding of S8, S15 and S17 to the 16S rRNA in *E. coli* SRS (PDB 5IT8).

**Fig. 22.** Interaction of S20 with the 16S rRNA in *E. coli* SRS (PDB 5IT8). Left, side view at the protein $\alpha$-helices; right, top view at the protein $\alpha$-helices.
important 16S rRNA helix H44 (Fig. 22). The deficit of S20 significantly reduces the rate of mRNA binding by ribosomes and decreases the SRS affinity to the LRS, which results in the reduced content of 70S ribosomes in vitro [126, 127]. The absence of S20 protein affects the formation of the SRS–fMet-tRNA–mRNA initiation complex [126, 127]. The absence of S20 protein affects the formation of the SRS–fMet-tRNA–mRNA initiation complex [126, 127]. Bacterial strains with deleted S20 protein grow significantly slower than the control bacteria [126]. Therefore, S20 plays an important role in the SRS functioning.

S20 is a small protein (under 90 a.a.). The structure of the isolated S20 has not been solved yet. S20 is an all-α protein; it contains three α-helices with the spectrin-like fold and, according to the SCOP classification, belongs to a separate protein superfamily (SCOP 46992).

S20 interacts with the 16S rRNA through amino acid residues of its α-helices that are not inserted into the RNA grooves, but situated somewhat aside. For example, the second and the third α-helices of the protein contact the 16S rRNA helix H9 by being positioned together in a close vicinity of “ridges” of the sugar-phosphate backbone (Fig. 22).

Analysis of the structural data on the interactions between primary RNA-binding r-proteins and rRNA indicates that these proteins recognize and bind double-stranded RNA regions. The side chains of conserved amino acid residues typically interact with atoms of the RNA sugar-phosphate backbone, i.e., this interaction is a classic example of the recognition of a specific spatial RNA fold. The contacts between amino acid residues and nucleotide bases are rare and typical for the RNA loops, where the bases may be unpaired and accessible for the intermolecular interactions.

The second feature of the interactions between r-proteins and rRNAs is the multiplicity of contacts for the primary binding r-proteins. These contacts stabilize the spatial structure of the 5S rRNA, 16S rRNA, and 23S rRNA in ribosomes and provide formation of functionally important sites, including the areas of the SRS–LRS interaction.

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