The SBP2 protein central to selenoprotein synthesis contacts the human ribosome at expansion segment 7L of the 28S rRNA

OLGA KOSSINNOVA,1,2 ALEXEY MALYGIN,1 ALAIN KROL,2,3 and GALINA KARPOVA1,3

1Institute of Chemical Biology and Fundamental Medicine, Siberian Branch of the Russian Academy of Sciences, Novosibirsk 630090, Russia
2Architecture et Réactivité de l’ARN, Université de Strasbourg, CNRS, IBMC, 67084 Strasbourg, France

ABSTRACT

SBP2 is a pivotal protein component in selenoprotein synthesis. It binds the SECIS stem–loop in the 3′ UTR of selenoprotein mRNA and interacts with both the specialized translation elongation factor and the ribosome at the 60S subunit. In this work, our goal was to identify the binding partners of SBP2 on the ribosome. Cross-linking experiments with bifunctional reagents demonstrated that the SBP2-binding site on the human ribosome is mainly formed by the 28S rRNA. Direct hydroxyl radical probing of the entire 28S rRNA revealed that SBP2 bound to 80S ribosomes or 60S subunits protects helix ES7L-E in expansion segment 7 of the 28S rRNA. Diepoxybutane cross-linking confirmed the interaction of SBP2 with helix ES7L-E. Additionally, binding of SBP2 to the ribosome led to increased reactivity toward chemical probes of a few bases in ES7L-E and in the universally conserved helix H89, indicative of conformational changes in the 28S rRNA in response to SBP2 binding. This study revealed for the first time that SBP2 makes direct contacts with a discrete region of the human 28S rRNA.

Keywords: mammalian ribosome; cross-linking approach; chemical probing; selenocysteine incorporation; SECIS-binding protein 2

INTRODUCTION

Selenocysteine (Sec) insertion into a growing polypeptide chain occurs in response to a UGA codon redefined as Sec, otherwise acting as a stop codon. To enable UGA recoding to Sec in eukaryotes, several cis- and trans-acting factors are required. All eukaryotic selenoprotein mRNAs contain a stem–loop in the 3′ UTR, called SECIS for SelenoCysteine Insertion Sequence (Berry et al. 1991). Key trans-acting factors are the translation elongation factor eEFSec that binds specifically the Sec-tRNASec complex (Fagegaltier et al. 2000; Tujebajeva et al. 2000; Gonzalez-Flores et al. 2012). The SECIS-binding protein 2 (SBP2) that interacts with the SECIS element (Copeland et al. 2000; Fletcher et al. 2001; Allmang et al. 2002). SBP2 occupies a central role in selenoprotein synthesis because it recruits the eEFSec•Sec-tRNA^Sec•GTP complex (Fagegaltier et al. 2000; Tujebajeva et al. 2000; Gonzalez-Flores et al. 2012). SBP2 was functionally characterized in Drosophila, humans, and rat (Copeland et al. 2000; Lescure et al. 2002; Takeuchi et al. 2009). The mammalian SBP2 can be divided into two domains. The N-terminal domain has an unknown function and is dispensable for selenoprotein synthesis in rabbit reticulocyte lysate (Copeland et al. 2000), while in this system the C-terminal part lacking the 344 N-terminal amino acids (CTSBP2) is able to maintain efficient selenoprotein synthesis (Copeland et al. 2001).

SBP2–SECIS interactions have been studied in great detail. The SBP2 RNA-binding domain as well as the SBP2-binding site on the SECIS element have been investigated. The SBP2 RNA-binding domain is composed of two subdomains, the L7Ae RNA-binding domain (Copeland et al. 2001; Fletcher et al. 2001, Caban et al. 2007; Cléry et al. 2007) which is shared with a family of functionally unrelated proteins (Copeland et al. 2000; Allmang et al. 2002) and a conserved lysine-rich module called either the bipartite RNA-binding domain, the selenocysteine insertion domain (SID), or the K-rich (lysine-rich) region, according to investigators (Bubenik et al. 2007; Donovan et al. 2008; Takeuchi et al. 2009). Copeland et al. (2001) showed that SBP2 co-sediments with ribosomes extracted from cultured cells. We showed that SBP2 binds purified human ribosomes in vitro, and specifically the 60S...
but not the 40S ribosomal subunit (Takeuchi et al. 2009). In that same study, we identified the amino acids that are crucial for the interaction with the 60S subunit, namely the KKPTSLKKIILKER527–540 amino acids (Takeuchi et al. 2009). Also, we and others found that the IILKER residues are important for both SECIS and ribosome binding (Donovan et al. 2008; Takeuchi et al. 2009).

Further studies were carried out to deepen our knowledge on the mechanistic issues leading to selenoprotein synthesis. Upon binding of SBP2 to the 80S ribosome, a few nucleotides in expansion segment ES31L and helix H89 of the 28S rRNA displayed increased accessibilities to a chemical reagent (Caban and Copeland 2012). These changes were proposed to be necessary for Sec-tRNA<sup>Sec</sup> entry to the ribosome aminoacyl (A) site. Recently, with the use of a minimal selenoprotein mRNA, we showed that SBP2 is already associated with the SECIS at the 48S initiation complex formation step; SBP2 mRNA, we showed that SBP2 is already associated with the SECIS at the 48S initiation complex formation step; SBP2 mRNA, we showed that SBP2 is already associated with the SECIS at the 48S initiation complex formation step; SBP2 mRNA. Also, we and others found that the IILKER residues are important for both SECIS and ribosome binding (Donovan et al. 2008; Takeuchi et al. 2009).

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In this work, we sought to identify the nature of the contacts that SBP2 establishes with the ribosome. To address this question in humans, we have applied several approaches in which the CTSBP2 recombinant protein was used. Using a cross-linking technique with bifunctional reagents, we showed that the ribosomal SBP2 binding site is mainly formed by the human 28S ribosomal RNA. Chemical probing of the 28S rRNA structure in complexes of CTSBP2 with the human 80S ribosomes or 60S subunits led us to determine the 28S rRNA regions changing accessibility to the probes upon CTSBP2 binding. Hydroxyl radical footprinting revealed that in both 80S and 60S ribosomes CTSBP2 protects a discrete region in helix ES7L-E in expansion segment ES7L of the 28S rRNA. Diepoxybutane cross-linking proved that the ribosome-bound CTSBP2 indeed interacts with ES7L-E. We showed that the ES7L-E 2D structure is exquisitely conserved in vertebrates and a few invertebrates. Additionally, a few bases in helices H89 and ES7L-E displayed increased reactivity toward dimethylsulfate upon CTSBP2 binding. Altogether, when mapped on the cryo-EM structure model of the human 80S ribosome (Anger et al. 2013) our data suggest that the SBP2-binding site is located on the solvent side of the 60S subunit, close to the ribosomal A site. Our results reported herein provide for the first time the evidence that SBP2 contacts a discrete region of the 28S rRNA.

**RESULTS**

**SBP2 contacts the 28S ribosomal rRNA in the 60S ribosomal subunit**

In a previous work (Takeuchi et al. 2009) we showed that CTSBP2 binds exclusively the 60S but not to 40S ribosomal subunit. However, as the actual localization of the CTSBP2-binding site on the ribosome was not identified at the time, we set out to determine it. At the onset, we established conditions for optimal formation of the SBP2-ribosome complex. The human recombinant CTSBP2 protein used in this and further experiments contained amino acids 345–856 and, as shown earlier, it is active in selenoprotein synthesis in rabbit reticulocyte lysate (Copeland et al. 2000; Lescure et al. 2002; Takeuchi et al. 2009). We determined that the binding of CTSBP2 to the purified human 60S ribosomal subunits is almost stoichiometric at a threefold excess of CTSBP2 over purified human ribosomes used at 0.5 μM (Fig. 1, lane 2). These conditions were used in all of the experiments described below.

To identify the ribosomal components interacting with SBP2, 80S-CTSBP2, 60S-CTSBP2 (and 40S-CTSBP2 as the control) complexes were treated with two bifunctional reagents, diepoxybutane or 2-iminothiolane. Diepoxybutane is highly reactive toward –NH<sub>2</sub> and –SH groups of proteins and G, C, and A residues in RNAs, forming near 4 Å-length RNA–protein and protein–protein cross-links (Baumert et al. 1978; Koskinen and Plna 2000); 2-iminothiolane can act either as an RNA–protein (cross-linking radius 5 Å) or a protein–protein cross-linker (cross-linking radius 14 Å), depending on the reaction conditions used to generate the cross-link; it reacts with the –NH<sub>2</sub> groups of lysines and U residues in RNAs (Traut et al. 1973). After treatment of the complexes with the bifunctional reagents, the distribution of cross-linked CTSBP2 between subunits as well as between ribosomal proteins and rRNAs was analyzed by dot-blotting with anti-SBP2 antibodies (Fig. 2). With diepoxybutane, we found that CTSBP2 cross-links only to the 60S and not the 40S subunit in the 80S ribosome (Fig. 2A). Furthermore, analysis of the cross-link distribution in the 60S-CTSBP2 complex revealed that the SBP2 signal was ~10-fold higher in the rRNA than in the ribosomal protein fraction, both with diepoxybutane and 2-iminothiolane (Fig. 2B). This strongly suggests that the rRNA contributes mainly to the formation of the SBP2-binding site. We cannot rule out, though, that the lower...
Hydroxyl radical probing of the 28S rRNA shows protection of discrete regions in expansion segment ES7L-E in SBP2-bound 80S and 60S human ribosomes

We next sought to determine the regions of the 28S RNA that are protected by SBP2 in the ribosome. To do so, hydroxyl radical probing of the 28S rRNA was carried out with the 80S- and 60S-SBP2 complexes. Hydroxyl radicals attack the C4′ atom of exposed ribose residues and thus induce strand scissions in the RNA phosphodiester bonds (Wu et al. 1983). Positions of the scissions were detected by primer extension of 5′-32P-labeled oligodeoxynucleotides. Almost all of the about 5000 nucleotides of the human 28S rRNA sequence were analyzed, with the exception of three regions: 50 nucleotides and 80 nucleotides at the very 5′ and 3′ ends, respectively, and 230 nucleotides in expansion segment ES27L; this was due to the presence of highly GC-rich regions impeding primer hybridization. The location of the 28S RNA regions protected by SBP2 against hydroxyl attack was mapped on the 28S rRNA 2D structure model derived from the cryo-EM structure of the human ribosome (Anger et al. 2013). The footprints reside in two regions of expansion segment ES7L whose accessibilities to hydroxyl radical attack were altered in the 60S-SBP2 complex vs. the free 60S subunits: Both regions were highly accessible in the free 60S subunits but displayed decreased accessibilities in the presence of SBP2 (Fig. 3A, cf. lane 2 with 3 and lane 9 with 10). These regions are 1133–1141, 1173–1186, and 1189–1190 at the apex of helices ES7L-F and ES7L-E, respectively (Fig. 3C). Using the SHAPE strategy, which leads to acylation of accessible 2′OH ribose residues, Caban and Copeland (2012) reported that SBP2 binding to the ribosome altered the reactivity of certain residues encompassing positions C4421 and U4419 in helix H89 and 4080–4166 in ES31L (numbering according to Anger et al. 2013). However, our footprinting experiments showed no differences in the accessibilities to hydroxyl radicals of these regions in the 60S-SBP2 complex (Supplemental Fig. S1), implying that at least the C4′ atoms of riboses (which are sensed by OH radicals) in those regions of H89 and ES31L are not involved in the SBP2-induced conformational changes. The same hydroxyl radical treatment was performed with the 80S-SBP2 complex to ask whether association of the 40S subunit could change the pattern of SBP2 protection on the 28S rRNA. To establish specificity of the SBP2 interactions with the ribosome, the same treatment was carried out with the 80S in the presence of the SBP2 mutant (MutSBP2) containing AAA 529–531 instead of TSL 529–531 and which is unable to bind the ribosome (Takeuchi et al. 2009). Supplemental Figure S2A (lane 8) showed that MutSBP2 was unable to generate a footprint, indicating that the footprints observed with ES7L-E were indeed due to the specific SBP2-ribosome interaction. Positions 1133–1141 in ES7L-F were not protected in the 80S ribosomal complex (Supplemental Fig. S2B); lack of protections in this
region is likely due to structural rearrangements following subunit association. Again, as with the 60S-CTSBP2 complex, no change in the protection pattern was detected either in H89 or ES31L (Supplemental Fig. S2C,D).

Diepoxybutane cross-linking validates the CTSBP2–ES7L-E interaction

To validate the hydroxyl radical probing data, we asked whether ES7L-E is in close proximity to CTSBP2 in the 80S-CTSBP2 complex. Besides, as H89 was found to adopt a conformation change upon CTSBP2 binding (Caban and Copeland 2012), we sought to determine whether CTSBP2 contacts H89. To answer these questions, we analyzed by reverse transcription the 28S rRNA isolated from 80S ribosomes treated by diepoxybutane in the presence of either wild-type CTSBP2 or the SBP2 mutant MutSBP2. Nucleotides C1183, G1189, G1194, and G1195 were cross-linked to CTSBP2 (Fig. 4, cf. lanes 6 and 7). These positions overlap the area found to be protected against hydroxyl radical attack. The lack of reverse transcription stops with MutSBP2 at these positions argues for the specificity of the cross-links (Fig. 4, cf. lanes 7 and 8). Taking into account that diepoxybutane forms near 4 Å–length RNA-protein cross-links, one can conclude that the four identified nucleotides tightly surround CTSBP2 on the ribosome or even contact it. Remarkably, no nucleotide was found to cross-link in the H89 region (which displayed increased accessibility toward benzoyl cyanide in the 80S-CTSBP2 complex) (Caban and Copeland 2012), and in the ES7L-F region protected by CTSBP2 from hydroxyl radical attack in the 60S-CTSBP2 complex (Supplemental Fig. S3A,B).

Chemical probing of helices ES7L-E and H89 reveals conformation changes upon SBP2 binding to the human 60S subunit

The protections against hydroxyl radical attack observed at the sugar-phosphate backbone of helix ES7L-E led us to ask whether changes in the accessibility of the chemical groups of the bases could also be observed in this helix upon SBP2 binding to the 60S subunit. To this end, we performed chemical probing of the 28S rRNA in the 60S-CTSBP2 complex with kethoxal or dimethylsulfate (DMS). Kethoxal treatment creates a cyclic adduct between the N1 and N2 positions of unpaired guanines, DMS methylates N1 of adenines, and N3 of cytosines when they are unpaired as well. Modified positions were identified by the primer extension method. Figure 5A shows the result of the DMS and kethoxal treatments at ES7L-E. Three positions were modified by DMS, C1180, C1181, and C1182. C1182 is highly modified in the free and complexed subunit; however, only C1180 and C1181 displayed an increased reactivity in the 60S-CTSBP2 complex vs. the free subunit (Fig. 5A, cf. lanes 6 and 7). G1185 reactivity toward

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**FIGURE 3.** Direct hydroxyl radical probing of the human 28S rRNA in 60S-CTSBP2 (A) and 80S-CTSBP2 (B) complexes. The sites of the hydroxyl radical-induced cleavages were mapped on the 28S rRNA by reverse transcription; reverse transcriptase stopped 3′ to the nucleotides whose ribose moieties were subjected to hydroxyl radical attack. Lanes 2,3,9,10 contained hydroxyl radicals (·OH). Lanes 1,8 lack hydroxyl radicals; lanes 2 and 9 lack CTSBP2. The extent of protection by CTSBP2 is shown on the left by a vertical bar indicating also the name of the protected helix. (C) The sites of protection identified in A and B are mapped onto the secondary structure of the 28S rRNA expansion segment ES7L (Anger et al. 2013); a blow-up of helices E and F of ES7L is displayed in the inset. Protections observed in both the 60S-CTSBP2 and 80S-CTSBP2 complexes are shown as closed circles, those found only in the 60S-CTSBP2 complex as open circles. Larger circles indicate higher protection. U,G,C,A lanes are sequencing markers.
kethoxal modification was observed upon SBP2 binding (Fig. 5B, lanes 9,10).

Structure-based sequence alignment reveals the extreme evolutionary conservation of a part of helix ES7L-E 2D structure

The footprints that were observed in ES7L-E in the presence of CTSBP2 raised the question of whether the 2D structure of this helix is conserved in selenoprotein-making organisms to enable SBP2 to bind. To answer this question, we analyzed the sequences of several 28S rRNAs between positions 956 and 1284 encompassing helices ES7L-E and ES7L-F (numbering according to the human 28S rRNA sequence from Anger et al. [2013]). Sequences that could not be manually folded as ES7L-E because of high-sequence dissimilarities and/or the presence of blocks of extended deletions, were discarded. For example, the sequences of Anopheles gambiae (Cannone et al. 2002), Plasmodium falciparum (Cannone et al. 2002), Drosophilia melanogaster (Anger et al. 2013), and Tetrahymena thermophila (Klinge et al. 2011) could not be considered because they possess much shorter ES7L sequences. Twelve organisms (10 vertebrates, one echinoderm, and one mollusk) were submitted to LocARNA (Will et al. 2007), a web-based tool, which simultaneously folds the input sequences and aligns them. Figure 6, A and B, reveals the perfect evolutionary conservation of the 2D structure of helix ES7L-E between positions 1170 and 1191 owing to a number of base covariations. The sequence of the G1171–C1190 and C1176–G1185 base pairs (human numbering) is conserved in the 12 organisms. Such a 2D structure conservation is indicative of a functional role of this helix.

Correlation between the sequence of the SBP2 K-rich region and the structure of the human 28S rRNA ES7L-E expansion segment

In an earlier work, we showed that the KKPTSLKKILK ER527–540 motif of the K (lysine)-rich domain in the human SBP2 is crucial for binding to the 60S ribosomal subunit (Takeuchi et al. 2009); these amino acids are conserved in the majority of the SBP2 sequenced so far, from sea urchin to mammals; however, insect SBP2 possesses the KK ITRLKKSVRVY instead of the KKPTSLKKIILKER sequence, and the sequence difference in the C-terminal part of this motif (SVRVY instead of ILKER) is by itself sufficient to change the pattern of SECIS recognition (Takeuchi et al. 2009). Interestingly, the length of the D. melanogaster 28S RNA ES7L is shorter and its sequence and 2D structure differ from the human counterpart due to deletions/point mutations (Anger et al. 2013). This led us to ask whether a correlation exists between the type of ES7L sequence/2D structure and the amino acid sequence in the SBP2 K-rich domain. To this end, we performed a multiple alignment of SBP2 amino acid sequences that are annotated in databases and
Modified nucleotides (one nucleotide prior to the reverse transcriptase stops) are marked on the right of the KKPTSLKKIILKER sequence in the K-rich region of SBP2 and the existence (or the lack) of the ES7L-E helix in 28S (26S) rRNAs.

DISCUSSION

The SBP2 protein plays a pivotal role in selenoprotein synthesis. It binds the SECIS element in the 3′ UTR of selenoprotein mRNAs and the translation elongation factor eEFSec to carry the eEFSec·Sec-rRNA Sec·GTP ternary complex to the ribosomal A site (Copeland et al. 2000; Tujebajeva et al. 2000; Gonzalez-Flores et al. 2012). Our previous work established that SBP2 binds to the 60S ribosomal subunit (Takeuchi et al. 2009) and we and others identified the SBP2 amino acids responsible for the interaction (Caban et al. 2007; Donovan et al. 2008; Takeuchi et al. 2009). However, the important question of the localization of the SBP2-binding site on the ribosome had not been solved. We addressed it in this work by using various approaches utilizing cross-linking and diverse chemical probes. We have found by hydroxyl radical footprinting experiments that the 80S ribosome-bound SBP2 protects a discrete region of the sugar-phosphate backbone of the 28S rRNA, namely, nucleotides 1173–1186 and 1189–1190 in helix ES7L-E. Diepoxybutane cross-linking validated the hydroxyl radical probing data and highlighted nucleotides C1183, G1189, G1194, and G1195 of ES7L-E, which tightly surround SBP2 on the 80S ribosome. Similar protection from hydroxyl radicals was observed in the 60S·CTSBP2 complex, supporting this finding. Base-specific probing in the 60S·CTSBP2 complex revealed enhanced accessibilities of the ES7L-E nucleotides C1180 and C1181 to DMS, and G1185 to kethoxal. Such a reactivity would imply that SBP2 induces structural rearrangements at the apex of ES7L-E upon binding. Interestingly, G1185 is conserved in the 12 organisms analyzed as the partner of the conserved C1176–G1185 base pair, and could therefore play an important as yet unknown role. However, the kethoxal reactivity of G1185 must be taken with care because G1185 is not single stranded, and kethoxal can occasionally generate base-pair disruption. In H89, A4414 and A4422 showed enhanced accessibility to DMS. Altogether the findings in this report, extending our previous ones (Kossinova et al. 2013), constitute the first evidence that the 28S rRNA plays a key role in the binding of SBP2 to the ribosome by implicating nucleotides in helix ES7L-E. As we can judge from the cryo-EM structure of the human ribosome (Anger et al. 2013), ES7L-E originating from two protozoa, three insects, and five vertebrate selenoprotein-synthesizing organisms (other vertebrate sequences are available but did not bring new information because of the high-sequence conservation between vertebrate species). Figure 7 shows that the finding observed in Drosophila (see above) extends to other species, while the KKPTSLKKIILKER sequence occurs only in vertebrates. In addition, an insertion of variable length is found in protozoan SBP2.

ES7L segments differ dramatically in length and nucleotide sequence between protozoa and insects as well as with vertebrates (Cannone et al. 2002). As mentioned above, these organisms have no helix homologous to ES7L-E; the lack of obvious sequence conservation, and the presence of deletions in these organisms did not enable us to propose a 2D structure model similar or homologous to helix ES7L-E.

In conclusion, our sequence analysis revealed the existence of a correlation between the concomitant presence or absence of the KKPTSLKKIILKER sequence in the K-rich region of the RNAs, Vol. 20, No. 7

FIGURE 5. Chemical probing of the 28S rRNA in 60S·CTSBP2 complexes. Reverse transcription analysis of the 28S rRNA in the ES7L-E and H89 regions is displayed in A and B, respectively. Modified nucleotides (one nucleotide prior to the reverse transcriptase stops) are marked on the right of the gels. Reactions contained DMS (A,B, lanes 6,7) or kethoxal (Ket: A, lanes 13,14; B, lanes 9,10). Control lanes 5,12 (A) and 5,8 (B) did not contain DMS or kethoxal; CTSBP2 was not added in lanes 6,13 (A) and 6,9 (B). Modified positions are mapped onto the secondary structures of ES7L-E and H89 (Anger et al. 2013). Bases modified by DMS are in open circles, bases displaying increased reactivity toward DMS are displayed in a dark background; G1185 (boxed in ES7L-E) has an increased accessibility toward kethoxal modification. U,G,C,A lanes are sequencing markers. A gel compression led to stacking of the CG sequence at positions 1188–1189; resolution is better with the kethoxal gel.
is moderately involved in the contact with other ribosomal components. In particular, only three nucleotides (of the 13 ones we have found protected by SBP2 against hydroxyl radicals) are located in close proximity to the amino acid side chains of ribosomal protein L5e.

Helix ES7L-E is part of the large expansion segment ES7L (866 nt) of the human 28S rRNA. We showed that a portion of the 2D structure of helix ES7L-E is conserved among vertebrate and several invertebrate species (at least in echinoderms and mollusks). SBP2 amino acids 527–540 in the K-rich domain important for SBP2-ribosome interactions are conserved in vertebrates as well (Takeuchi et al. 2009). Remarkably, the 28S rRNAs of land plants that do not synthesize selenoproteins have no ES7L-E at all (Cannone et al. 2002). The above and our findings suggest that during selenocysteine insertion, the conserved K-rich motif of SBP2 interacts with the conserved ES7L-E helix. This suggestion is also supported by the observations of Gupta et al. (2013).

**FIGURE 6.** Structure-based alignment of expansion segment ES7L-E sequences in 28S rRNAs arising from 12 vertebrates and invertebrates. (A) The region boxed shows 2D structure conservation through base covariations of a section of helix ES7L-E. Color code: The hues show sequence conservation (red corresponds to absolutely conserved and green indicates the least conserved base pairs) and saturation decreases with the number of compatible base pairs, thus showing structural conservation (generated automatically by LocARNA). Parentheses indicate base pairs. Because most of the sequences are partial, numbering according to the human sequence was not possible. The first nucleotide in the sequence of *Homo sapiens* corresponds to C1148. (B) Secondary structure model of a section of the human ES7L-E helix supported by the base covariations shown in A. The number of covariations for each base pair is indicated on the right. Color code as in A and numbering as in Figure 5A. Open circles represent nonconserved nucleotides in the apical loop and base pairs that do not covary.

**FIGURE 7.** Multiple sequence alignment of the SBP2 K-rich domain. The K-rich domain, corresponding to residues 517–544 in the human SBP2, is boxed. SBP2 sequences are from *H. sapiens* (UniprotKB accession number Q96T21), *Pan troglodytes* (H2QXFS), *Mus musculus* (Q3U1C4), *Gallus gallus* (XP_424425.4), *Xenopus tropicalis* (NP_001090731.1), *A. gambiae* (Donovan and Copeland 2009), *D. melanogaster* (NP_648204.1), *Drosophila virilis* (XP_002046871.1), *P. falciparum* (XP_001351170.1), and *T. thermophila* (XP_001011959) and were aligned by MUSCLE (Edgar 2004) and displayed with JalView.
showing that ribosomes from non-Sec-utilizing organisms are unable to support Sec incorporation. As a matter of fact, to carry out selenoprotein synthesis in wheat-germ extract (that is unable to support selenoprotein synthesis), these investigators had to complement it not only with specialized factors and tRNA<sub>Sec</sub> but also with mammalian ribosomes. One can see from the available 28S rRNA 2D structures (Cannone et al. 2002) that ES7L-E differs or is absent in selenoprotein-making organisms of algae, protists, and most invertebrates. Interestingly, the findings described in this work establish that sequence/structure changes in ES7L-E are accompanied by amino acid changes in the K-rich region of SBP2, very likely for maintaining the ability of the protein to interact with the 60S subunit. In other words, a correlation exists between the structure of ES7L-E and the SBP2 amino acid composition in the K-rich region. The lack of correlation between Drosophila SBP2 and mammalian ribosomes could explain why this protein, which carries different amino acids from mammals in the K-rich region, displayed a drastically lower selenoprotein synthesis activity in rabbit reticulocyte lysate compared with human SBP2 (Takeuchi et al. 2009).

This minor activity of dSBP2 could have related to its eukaryote-specific L7Ae domain, whose mammalian homolog was shown to be able to weakly bind to the ribosome (Donovan et al. 2008). Our data, combined with those of the literature cited above, strongly suggest that in vertebrates and a few invertebrates the K-rich domain of SBP2 interacts with helix ES7L-E of the 28S rRNA. Concerning lower eukaryotes, we can hypothesize that the lack of helix ES7L-E in the 28S rRNA is compensated by changes in the amino acids of the SBP2 K-rich domain in order to provide SBP2 binding to the shortened ES7L.

According to the cryo-EM model of the human 80S ribosome (Anger et al. 2013), ES7L-E resides in the 80S ribosome on the solvent side of the 60S subunit, near the ribosomal A site (Fig. 8A). Such a location for the SBP2-binding site would be in good accordance with the function of SBP2 proposed by Gonzalez-Flores et al. (2012), which is to facilitate delivery of the eEFSec-Sec-tRNA<sub>Sec</sub>-GTP ternary complex to the ribosomal A site. Helix H89 of the 28S rRNA resides in the 80S ribosome at the 60S subunit surface, facing the 40S subunit and extending from the peptidyl transferase center (PTC) loop (Anger et al. 2013). The secondary structure of helix H89 is extremely well conserved in the three domains of life (Cannone et al. 2002). Its apical loop is involved in the binding of IF2 in bacteria (La Teana et al. 2001) and eIF5B in eukaryotes (Unbehaun et al. 2007); the helix itself is essential for maintaining translation fidelity in bacteria (O’Connor and Dahlberg 1995) and its structure is important for ribosome peptidyl transferase function in E. coli (Burakovskaya et al. 2011). Moreover, H89, together with helices H90–H92, forms the aminoacyl-tRNA accommodation corridor (Sanbonmatsu et al. 2005). Thus, H89 is involved in the different steps of the translation process. During Sec insertion, the conformational changes implicating H89, namely the increased reactivity to DMS of A4414 and A4422 (this study) and of the 2′OH ribose of U4419 and C4421 to benzoyl cynaide (Caban and Copeland 2012), could be a consequence of the structural transitions in the 60S subunits caused by the contacts of SBP2 with ES7L-E inducing conformational changes in this helix at C1180, C1181, and G1185. In this respect, we found that at least 85Å separate ES7L-E from H89 (measured with PyMol) (DeLano 2002) in the cryo-EM model of the human 80S ribosome (Anger et al. 2013). To explain how SBP2 could induce conformational changes in H89 at a distance, we propose the following hypothesis. As ES7L-E is in close proximity to ribosomal protein L5e, which in turn contacts L10e, and as the N-terminal “hook” of L10e inserts into the H89 bulge (Anger et al. 2013), the SBP2-induced conformational changes at the apex of helix ES7L-E could propagate to H89 (Fig. 8B). Remarkably, the yeast L10e has been already considered as the protein playing a key role in the allosteric transmission of information by...
conformational rearrangements through the ribosome (Petrov et al. 2008; Sulima et al. 2013). Gonzalez-Flores et al. (2012) propose that the conformational changes in H89 provide the ability of H89 to interact with Domain IV of the elongation factor eEFSec in order to dissociate the ternary complex eEFSec-Sec-ribosome-SecGTP and to deliver the Sec-ribosome to the A site.

The results of our study shed light on structural aspects of the SBP2-ribosome interactions in the course of selenoprotein synthesis. We have shown for the first time that the 28S rRNA is involved in SBP2 binding and identified ES7L-E as the helix contacting SBP2. This helix is moderately involved in intraribosomal interactions and its 2D structure is highly conserved in vertebrates and a few invertebrates by virtue of a number of base covariations. We have also determined the bases of the 28S rRNA involved in the conformational changes induced by SBP2 binding to the ribosome. Additionally, the findings presented herein led us to propose the localization of the SBP2 binding site on the human 80S ribosome.

MATERIALS AND METHODS

Preparation of recombinant CTSBP2

CTSBP2 (amino acids 343–854) fused to a 6xHis N-terminal tag was expressed in E. coli BL21(DE3)RIL strain (Novagene) as described in Takeuchi et al. (2009).

Purification of human ribosomal subunits

Human ribosomal 40S and 60S subunits were isolated from full-term placenta as described in Matasova et al. (1991). The concentration of subunits was determined assuming that one A260-unit equals 50 pmol and 25 pmol in the case of 40S subunits and 60S subunits, respectively (Matasova et al. 1991). The activities of purified ribosomes were validated by poly-U-dependent synthesis of 14C-polyphenylalanine in vitro.

Formation of CTSBP2-ribosome complex

The 0.5 μM 60S subunits were incubated with 1.5 μM CTSBP2 in 50 μl of buffer A (20 mM HEPES-KOH at pH 7.5, 100 mM KCl, 2 mM MgCl₂, 1 mM DTT) at 25°C for 30 min. The complex was isolated by centrifugation in 10%–30% sucrose gradient in buffer A containing 450 mM KCl (rotor SW41, 23,000 rpm, 17 h, 4°C). Fractions corresponding to 40S and 60S subunits were ethanol precipitated, dissolved in buffer A, blotted onto nitrocellulose membrane, and analyzed for CTSBP2 content with rabbit polyclonal anti-CTSBP2 antibodies, which were shown beforehand not to cross-react with ribosomal proteins (1/2500 dilution). Membranes were then treated with anti-rabbit HRP-conjugated secondary antibody (1/10000 dilution), revealed with the ECL-plus kit (GE Healthcare) and exposed to either X-ray film (GE Healthcare) or ChemiDoc XRS (BioRad). To analyze the cross-link distribution between rRNAs and proteins in 60S subunits, the corresponding subunit fraction prepared as described above was divided into two halves. One half was used for protein extraction with acetic acid according to Hardy et al. (1969), and total rRNA was isolated from another half by phenol extraction as in Graifer et al. (1994). Both ribosomal protein and total rRNA fractions were analyzed with anti-SBP2 antibody as described above. To analyze cross-link distribution between rRNAs in the 60S subunit, the corresponding subunit fraction prepared as described above was incubated for 30 min at 37°C in 0.5% SDS, 5 mM NaOH-EDTA (pH 7.5). The mixture was layered onto a 5%–20% sucrose gradient in 20 mM Tris-HCl (pH 7.5), 2 mM EDTA, 0.1% SDS, and centrifuged for 17 h at 27,000 rpm (rotor SW41). After centrifugation, fractions were ethanol precipitated, the pellets dissolved in a minimal amount of water, and blotted onto a nitrocellulose membrane. The SBP2 signal was detected as above.

Alternatively after quenching, the mixture was supplemented with 0.1% SDS, 4 mM EDTA, 0.05 mg/ml proteinase K, and incubated for 20 min at 37°C. rRNA was isolated by phenol extraction. Reverse transcription and product analysis were carried out as described in Malygin et al. (2013).

2-iminothiolane cross-linking

CTSBP2-ribosome complexes were formed as described above, CTSBP2-rRNA cross-links and SBP2-ribosomal proteins cross-links with 2-iminothiolane were performed as described (Kenny et al. 1979; Brimacombe et al. 1988). CTSBP2-rRNA cross-links and CTSBP2-ribosomal proteins cross-links were analyzed as described above.

Hydroxyl radical, kethoxal, and DMS reactions

Hydroxyl radical cleavage of 28S rRNA in 60S or 80S ribosomes (0.5 μM) and their complexes with CTSBP2 (1.5 μM CTSBP2) was performed as described in Malygin et al. (2013). Kethoxal and dimethylsulfate (DMS) modifications were performed according to Xu and Culver (2009) with the only modification that the final DMS concentration was 0.15% (v/v). The RNA was isolated by phenol extraction. For reverse transcription, 5′-32P-labeled primers complementary to the human 28S rRNA sequence regions 196–212, 394–413, 508–525, 693–709, 735–753, 948–968, 1178–1195, 1221–1240, 1298–1315, 1435–1454, 1589–1605, 1792–1809, 1991–2008, 2160–2178, 2313–2331, 2406–2422, 2618–2639, 2795–2811, 2889–2907, 3047–3064, 3260–3277, 3626–3644, 3809–3829, 4036–4055, 4184–4200, 4361–4378, 4495–4515, 4540–4558, 4721–4739, 4880–4898, 5014–5032, 5254–5271, and 5408–5425 was conducted.
and 5010–5028 were used. Reverse transcription and product analysis were carried out as described in Malygin et al. (2013).

**Structure-based sequence alignment of ES7L**


**SUPPLEMENTAL MATERIAL**

Supplemental material is available for this article and can also be accessed in PDF format at http://www.niboch.nsc.ru/lib/exe/fetch.php/r2/structure/labs/kar/2014_sd_kossinova_et_al.pdf.

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SBP2 binding site on the human ribosome


The SBP2 protein central to selenoprotein synthesis contacts the human ribosome at expansion segment 7L of the 28S rRNA

Olga Kossinova, Alexey Malygin, Alain Krol, et al.

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