SRSF1 interactome determined by proximity labeling reveals direct interaction with spliceosomal RNA helicase DDX23

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SRSF1 is the founding member of the SR protein family. It is required—interchangeably with other SR proteins—for pre-mRNA splicing in vitro, and it regulates various alternative splicing events. Dysregulation of SRSF1 expression contributes to cancer and other pathologies. Here, we characterized SRSF1’s interactome using proximity labeling and mass spectrometry. This approach yielded 190 proteins enriched in the SRSF1 samples, independently of the N- or C-terminal location of the biotin-labeling domain. The detected proteins reflect established functions of SRSF1 in pre-mRNA splicing and reveal additional connections to spliceosome proteins, in addition to other recently identified functions. We validated a robust interaction with the spliceosomal RNA helicase DDX23/PRP28 using bimolecular fluorescence complementation and in vitro binding assays. The interaction is mediated by the N-terminal RS-like domain of DDX23 and both RRM1 and the RS domain of SRSF1. During pre-mRNA splicing, DDX23’s ATPase activity is essential for the pre-B to B spliceosome complex transition and for release of U1 snRNP from the 5′ splice site. We show that the RS-like region of DDX23’s N-terminal domain is important for spliceosome incorporation, while larger deletions in this domain alter subnuclear localization. We discuss how the identified interaction of DDX23 with SRSF1 and other SR proteins may be involved in the regulation of these processes.

Human SRSF1 was the first member of the SR protein family to be discovered in the early 1990s, when its activities in the complementation of pre-messenger RNA (pre-mRNA) splicing in vitro and in the regulation of 5′-splice-site selection were described (1–3). SRSF1 is a 248-residue protein with two RNA-Recognition Motifs (RRMs) at its amino-terminal end, followed by an RS domain with multiple consecutive RS or SR dipeptides—a defining feature of the SR protein family (4). The RRM2s are responsible for specific RNA binding, but they can also engage in key interactions with SRSF1 functional partners and with the RS domain (5–7). The SRSF1 RS domain consists of 51 amino-terminal residues, with 37% arginine (R) and 39% serine (S) content, organized as blocks of consecutive dipeptide repeats. Many of the serine residues in the RS domain are phosphorylated by kinases that regulate SR protein localization and recruitment to nascent transcripts (8). At steady state, the SRSF1 RS domain is highly phosphorylated (9). In general, RS domains are responsible for protein–protein interactions, though unphosphorylated RS domains have also been proposed to interact with RNA (10, 11).

During early spliceosome assembly, SRSF1 engages in protein–protein interactions with a U1 small nuclear Ribonucleoprotein (snRNP) polypeptide (U1-70K) at the 5′ splice site (5′ss) and with the U2AF1/U2AF2 heterodimer at the 3′ splice site (12), promoting the commitment of the pre-mRNA to splicing (13). SRSF1 also specifically binds U1 snRNA stem-loop 3 and can be recruited to the pre-mRNA via U1 snRNA, to participate in exon definition (14). Recent studies found additional connections of SRSF1 with later-stage spliceosome and associated components, including the SF3 complex (15) and the exon-junction complex (16–18). SRSF1’s RRM2 was recently crosslinked and modeled into a cryo-EM structure of the pre-B→B spliceosome complex, proximal to the 5′ end of the upstream exon, near the cap-binding complex and additional B-box-specific proteins, including TCFRG1, MPA1, and UBL5 (19).

In this work, we used a proximity-labeling method (20) to characterize SRSF1’s interactome, and we then validated a strong direct interaction between SRSF1 and DDX23 (also known as PRP28 or U5-100kD). DDX23 is an 820-residue protein that belongs to the DEAD-box family of ATP-dependent RNA helicases and is involved in the displacement of U1 snRNP from the 5′ss during the pre-B to B spliceosome-complex transition (21, 22). Mapping of the regions required for the interaction revealed that both RRM1 and DDX23 are implicated in the regulation of SRSF1’s splicing activity.

**Significance**

SR proteins are essential for pre-mRNA splicing, but the molecular details underlying their mechanism of action within the spliceosome remain unclear. We used proximity labeling to identify many proteins that are proximal to SRSF1 in HeLa cells. Among them, we uncovered and characterized a direct interaction between SRSF1 and the spliceosomal RNA helicase DDX23/PRP28. This interaction has potential functional implications for a discrete stage of spliceosome assembly. Detailed knowledge about molecular interactions between key components of the splicing machinery may yield unique approaches to target the spliceosome in pathologies that involve dysregulation of pre-mRNA splicing.

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and the RS domain of SRSF1 contact the N-terminal (Nt) RS-like domain of DDX23. Deletions within this DDX23 domain disrupted the interaction with SRSF1. Phosphorylation of the RS repeats in a minimal DDX23 construct also inhibited the interaction. Furthermore, deletion of the first 327 amino acids of DDX23 resulted in altered subnuclear localization, and a 37-residue deletion within the RS-like domain showed decreased spliceosome recruitment during in vitro splicing. Therefore, the Nt domain of DDX23 is involved in its subnuclear targeting, including incorporation into the spliceosome, and its interaction with SRSF1 likely facilitates or regulates these processes.

Results

SRSF1 Interactome Analysis by Proximity Labeling. We designed two SRSF1 cDNA constructs fused to BioID2—a small, promiscuous biotin ligase developed for proximity labeling (23); in these constructs, the biotin-ligase domain is located at the N-terminus (BS) or at the C-terminus (SB), respectively (Fig. 1A).

We designed an additional construct to express BioID2-EGFP-3xNLS (BG) as a nuclear nonspecifically biotinylating control (Fig. 1A). Retrotransduction of HeLa cells followed by puromycin selection yielded three cell lines that express the proteins of interest. The expression level attained for the SRSF1 fusion constructs was markedly lower than the endogenous SRSF1 level, so as to limit unspecific biotinylation due to overexpression (Fig. 1B). Immunofluorescence staining demonstrated that the addition of the BioID2 domain at the N- or C-terminus of SRSF1 did not alter the predominantly nuclear localization pattern of SRSF1 (SI Appendix, Fig. S1A). Moreover, fusion of the BioID2 domain to SRSF1 did not affect its dose-dependent ability to regulate the expression of endogenous SRSF1 (24) or alternative splicing of the endogenous target PEA15 (25), suggesting that the fusions do not hinder SRSF1’s splicing-regulatory activity (SI Appendix, Fig S1 B and C).

We used these HeLa cell line derivatives (BG, BS, SB) to study SRSF1’s interactome by proximity labeling. After a 24 h biotin pulse in live cells and streptavidin purification of the biotinylated proteins, we analyzed triplicate samples for each condition by Liquid Chromatography with tandem Mass Spectrometry (LC/MS-MS) (Dataset S1). We compared the samples expressing the SRSF1 fusions to the nuclear EGFP control. We observed a general enrichment of proteins in the samples expressing SRSF1-BioID2 fusions (Fig. 1C and D). SRSF1 was the most enriched protein in both samples, with an enrichment value of more than 4,000-fold (Fig. 1 E and F). A total of 353 proteins were enriched ≥5 times in the BS dataset with respect to the BG control, whereas 246 proteins were similarly enriched in the SB dataset. Of these proteins, 190 were present in both datasets, highlighting the strong overlap between the two lists, and suggesting that the retrieved candidates represent functionally relevant proteins proximal to SRSF1 (Fig. 1G, Dataset S1, and SI Appendix, Table S1).

Gene Ontology (GO)-term analysis of the overlap list revealed an enrichment for proteins related to RNA splicing (GO:0006380, FDR: 1.97E-36), which confirms that SRSF1 is still able to engage in pre-mRNA splicing in the presence of the BioID2 fusions. In addition, enrichment of proteins related to nuclear speckles (GO:00016607, FDR: 2.32E-20) suggests that the retrieved proteins colocalize with SRSF1 in its predominant subcellular compartment. The significant enrichment of proteins susceptible to ubiquitin-like conjugation (KW:0832, FDR: 7.97E-24) may reflect the reported role of SRSF1 in the SUMO conjugation pathway (26, 27), whereas the enrichment of proteins related to mRNA transport (GO:00051028, FDR:2.01E-6) is consistent with the ability of SRSF1 to promote mRNA export, and its known association with postsplicing events like translation and nonsense-mediated mRNA decay (NMD) (17, 18, 28–32) (Fig. 1H).

Among the proteins with the highest enrichment score, we found well-characterized SRSF1 interactors like U2AF1 (BS: 500× enrichment; SB: 68×) and core spliceosome proteins like SF3A2 (BS: 125×; SB: 156×), SF3B2 (BS: 13×; SB: 59×), TCERG1 (BS: 28×; SB: 39×), and CACTIN (BS: 127×; SB: 175×), proteins associated with the EJC like RNPS1 (BS:19×; SB: 66×) and PNN (BS: 32×; SB: 28×), and other splicing-associated proteins like NONO (BS: 32×; SB: 24×). In addition to splicing-related proteins, other functional aspects of SRSF1 were reflected. For instance, several histone proteins including H2A.1 (BS: 490×; SB: 1276×), Histone H2B type 3-B (H2BUI) (BS: 444×; SB: 741×), H3.2 (BS: 18×; SB: 54×), and H3.3 (BS: 24×; SB: 5×) were specifically enriched. These interactions are likely relevant, as SRSF1 was reported to be associated with chromatin, specifically with the histone H3 tail during interphase (33). Strong enrichment of proteins related to other RNA-processing events, like the alternative cap-binding protein NCBP3 (BS: 29×; SB: 287×) and the mRNA 3′-end processing protein FIP1L1 (BS: 226×; SB: 297×), suggests that coordination between splicing and these processing events could be mediated by SRSF1.

The proximity-labeling approach led to strong enrichment of several proteins that were not previously known to physically or functionally interact with SRSF1, including AHNAK2 (BS: 796×; SB: 217×), HAUS2 (BS: 950×; SB: 849×), RABL6 (BS: 374×; SB: 40×), HMGXB4 (BS: 397×; SB: 41×), and PPHLN1 (BS: 26×; SB: 471×). AHNAK2 is a 616-kDa PDZ-domain scaffold protein that interacts with many proteins (34) and was computationally inferred, using the Phylogenetic Annotation and Inference Tool (PAINT), to have a role in splicing regulation (35). HAUS2 is involved in mitotic-spindle assembly, maintenance of centrosome integrity, and cytokinesis (36). SRSF1 was reported to promote translation of mitosis-associated genes (28, 37), so a potential functional connection with HAUS2 during the events leading to mitosis warrants further exploration. Finally, connections between SRSF1 and the GTPase RABL6, the epigenetic repressor PPHLN1, and the transcription regulator HMGXB4 were not previously described, but their high enrichment scores suggest possible functional connections that merit further study.

The overlap list reflected all stages of spliceosome assembly, including E-complex proteins (U2AF1, U2AF2, and SF1), A-complex proteins (SF3A2, SF3B2, U2SURLP), B/C-complex proteins (DDX23, TCERG1, SART1, CACTIN, SRRM1, SNW1, RBM42), and components of the EJC (RNPS1, PNN). This observation is consistent with spliceosome-composition studies that found that SRSF1 is associated with the spliceosome from the earliest stages of assembly and remains bound to the mature mRNA after splicing is completed (38). In addition, we detected several proteins that participate in alternative splicing, including other SR proteins (SRSF3 and SRSF11) and hnRNPsuperfamily members (HNRNPC, HNRNPA3, HNRNPC, HNRNP1, HNRNP2, HNRNPAB). Proximity to these proteins likely reflects a shared functional environment, which is expected for RNA-splicing regulators (Fig. 1H). It is possible that some of these interactions take place when these proteins are bound to the same RNAs in close proximity.

Finally, two additional cytoplasmic functions of SRSF1 were reflected in the overlapping protein list, namely its proposed connection with mTOR kinase during translation (29) and the effect of SRSF1 on NMD (32). In particular, we detected enrichment of the mTOR-pathway components RICTOR (BS: 10×; SB: 11×)
Fig. 1. Proximity labeling highlights connections of SRSF1 to the spliceosome in addition to recently identified functions. (A) Constructs used for proximity labeling. M: Myc tag, H: HA tag. (B) Western blot analysis of lysates from HeLa cells transduced with the indicated retroviruses, after puromycin selection. Arrows point to the specific immunoreactive bands. (C) Volcano plot showing protein enrichment in the BS sample, compared to the BioID2-GFP control. Selected proteins are highlighted. (D) Volcano plot showing protein enrichment in the SB sample, compared to the BioID2-GFP control. Selected proteins are highlighted. (E) Representation of 353 proteins enriched in the BioID2-SRSF1 dataset. (F) Representation of 246 proteins enriched in the SRSF1-BioID2 dataset. (G) Venn diagram of proteins detected in the BS and SB datasets. (H) Protein–network visualization generated with Cytoscape and STRING database, derived from the 190 proteins in common between BS and SB datasets. Colors represent the indicated GO terms (RNA splicing, Ubi conjugation, mRNA transport, nuclear speckles). Proteins were clustered according to their functions. The thickness of the connecting lines represents the degree of confidence for protein–protein functional interaction, obtained from the STRING database.
proteins identified by proximity labeling can physically interact with proteins that were recently modeled near SRSF1 in a cryo-EM structure of the pre-B complex (deletion mutants Δ127 and Δ77 completely abolished the interaction (Fig. 3C). In contrast, removal of the first 40 residues of DDx23 did not affect the interaction, suggesting that the region of DDx23 that contacts SRSF1 involves at least 37 residues, from S41 to R77 (Fig. 3D). To further test whether these 37 residues are sufficient for the interaction with SRSF1, we expressed TrxA (E. coli Thioredoxin 1) fused at its C terminus to these residues and tested the interaction with untagged SRSF1. The PD analysis confirms that these DDx23 residues interact with SRSF1 (Fig. 3F). This region is conserved in higher eukaryotes (but is missing in Caenorhabditis elegans and Schizosaccharomyces pombe) and includes motifs with SK/KD/RK/ER/RR and RS consecutive repeats (Fig. 3F). The presence of these negatively and positively charged sequence features suggests potential electrostatic interactions with SRSF1.

To further define the interaction region between SRSF1 and DDx23, we generated a construct [Ddx23-Δ(S41-R77)] that lacks the S41-R77 region that on its own binds SRSF1. Notably, this construct retained the ability to bind SRSF1, suggesting that redundant regions within the Nt-domain of DDx23 can still interact with SRSF1 in the absence of the S41-R77 region (SI Appendix, Fig. 6D). Indeed, the first 40-residue peptide (M1-P40) also comprises many charged residues (35% K+R, 27.5% D+E), including 10 consecutive alternating-charge residues, and is conserved in higher eukaryotes (SI Appendix, Fig. S7). Our results are consistent with a flexible interaction between the unstructured Nt-region of DDx23 (M1-R77) and SRSF1, which may involve internally redundant segments of the former.

Deletion of the RS domain of SRSF1 did not abolish the interaction with FL-DDx23 (SI Appendix, Fig. S8A). We also tested isolated domains of SRSF1—fused to the Ct of His-TrxA—for interaction with DDx23, and both TrxA-RRM1 and the
Fig. 2. BiFC validates interactions of SRSF1 with selected spliceosomal proteins. (A) Schematic representation of the BiFC approach. SRSF1 is shown fused to mVenus-173-Nt. mVenus-82-Ct is shown fused to a circle that represents the various interactor candidates tested. The interaction between the candidate and SRSF1 brings the two mVenus halves closer together, resulting in stronger green fluorescence. (B) BiFC results for DDX23 visualized by fluorescence microscopy. BF, bright field. Scale bar represents 130 μm in 10× images and 15 μm in 40× images. (C–F) BiFC results for the indicated proteins, plotted as Forward scatter (linear scale) vs. Green fluorescence (logarithmic scale). The vertical line indicates the threshold for cells that were considered mVenus positive. The bar graphs on the right of each panel represent % mV+ cells and mean green fluorescence, respectively, for Interactor-mV Ct coexpressed with Nt-mVenus vs. Interactor-mV Ct coexpressed with SRSF1-Nt-mVenus. An unpaired t test was performed to compare the %mV+ cells and MGF values between the experimental conditions and the controls. ****P ≤ 0.0001. Error bars represent SD.
phosphorylated TrxA-RS domain—generated by coexpression with CLK1—successfully pulled down DDX23. In contrast, the isolated TrxA-RRM2 did not show detectable binding to DDX23 (Fig. 3G).

The presence of RS-like regions in spliceosomal proteins from higher eukaryotes is usually correlated with the evolutionary emergence of SR proteins (43). Therefore, it has been proposed that RS-like regions could bind to the RS domains of SR proteins and that these interactions could help regulate splicing in these organisms. The DDX23 RS-like domain has a conserved nine-residue stretch of RS repeats (S57 to S65), whereas the remaining RS repeats are shorter and dispersed (Fig. 3B). To test the relevance of the long RS-repeat stretch for the interaction, we generated a

Fig. 3. A direct interaction between SRSF1 and DDX23 can be recapitulated using recombinant proteins in vitro. (A) Schematic representation of domains of DDX23. (B) AlphaFold prediction of the structure of DDX23 (AF-Q9BUQ8-F1). RS/SP residues present in the RS-like domain are highlighted as sticks. RecA1 domain is shown in orange and RecA2 domain in yellow. (C) Representative His Pulldown assay using His-SUMO-SRSF1 and untagged DDX23, DDX23-Δ77, DDX23-Δ127, with or without benzonase treatment. (D) Representative His Pulldown assay using His-SUMO-SRSF1 and the indicated DDX23 constructs. (E) Sequence alignment of the SRSF1-interacting fragment of the RS-like domain of human DDX23 and its orthologs. This region is absent in the proteins from C. elegans and S. pombe. The location of the 9 SR residues is marked by a horizontal line. Charged residues conserved in the top five species are in red font. Numbers represent residue position in the human protein. (F) Representative His Pulldown assay using His-TrxA or His-TrxA-DDX23 (S41-R77) and untagged SRSF1. (G) Representative His Pulldown assay using the indicated truncated versions of SRSF1 or His-TrxA and FL-DDX23. (H) Representative His Pulldown assay using His-TrxA-RRM1, or His-TrxA and WT or 9GA-mutant DDX23.
Phosphorylation of the RS Repeats in DDX23 (S41-ΔR77) Inhibits SRSF1 Binding. DDX23 was previously shown to undergo phosphorylation of its RS repeats, with one of the kinases responsible for this phosphorylation being SRPK2 (44). Although the precise residues that undergo phosphorylation were not mapped, DDX23 phosphorylation was shown to be required for B-complex formation and for stable association of DDX23 with the tri-snRNP (44). Considering that the SRSF1-interaction region we mapped harbors several RS repeats, including the above-mentioned nine-residue stretch, we tested whether phosphorylation of these residues affects the interaction with SRSF1. We coexpressed His-TrxA-DDX23 (S41-R77) with CLK1 kinase in bacteria, to obtain a phosphorylated version of the protein. CLK1 was previously shown to phosphorylate DDX23 in vitro (21), and we determined the phosphorylation pattern by MS analysis (SI Appendix, Fig. S9). In addition, we generated a mutant version of the minimal protein fused to His-TrxA, with the 9RS motif (S57 to S65) substituted by GA repeats (9GA). Phosphorylation of the RS repeats in the minimal DDX23 construct markedly reduced the efficiency of the PD with SRSF1. In the context of this minimal version of DDX23, the 9GA mutant had no detectable interaction with SRSF1 (Fig. 4A). We conclude that the phosphorylation state of the conserved RS repeats affects the interaction with SRSF1 mediated by the positively charged arginines. Moreover, the fact that the 9GA mutant in this context impairs binding to SRSF1, whereas it has no effect in the context of FL-DDX23, is consistent with other charged regions within the DDX23 Nt domain helping to stabilize the interaction with SRSF1.

DDX23’s (S41-R77) Region Is Important for Recruitment to the Spliceosome. To better understand the possible functional consequences of the DDX23-SRSF1 interaction, we characterized the role of the Nt-domain of DDX23 in spliceosome assembly. A previous study demonstrated that an ATPase catalytically inactive mutant of DDX23 (DEAD to AAD mutation) results in stalling of the spliceosome at the pre-B stage; therefore, the addition of recombinant DDX23-AAD to in vitro splicing reactions results in splicing inhibition (41). We expressed and purified recombinant DDX23-AAD and DDX23-AAD-Δ(S41-R77)—a version of the protein that is both catalytically dead and lacks the (S41-R77) region. We performed in vitro splicing reactions using HeLa cell nuclear extract preincubated with recombinant DDX23-wt, DDX23-AAD, or DDX23-AAD-Δ(S41-R77). Splicing reactions with added DDX23-wt proceeded normally, whereas incubation with the DDX23-AAD mutant resulted in splicing inhibition, as expected, reflecting competition of the AAD mutant with endogenous DDX23-wt for spliceosome incorporation. In contrast, incubation with the DDX23-AAD-Δ(S41-R77) mutant did not inhibit splicing. This result is consistent with the DDX23-AAD-Δ(S41-R77) mutant failing to incorporate into the spliceosome (Fig. 4B). Given that SRSF1 interacts with this region of DDX23, it is possible that SR protein binding has a role in regulating DDX23’s recruitment to the spliceosome.

The Conserved DDX23 Nt Domain Is Required for Proper Localization. To further characterize the function of DDX23’s Nt RS-like domain, we generated a series of mutants that lack the first 77, 127, or 327 residues of DDX23 and are fused at their N-terminus to mVenus. We used these constructs to transfect 293T cells and track the localization pattern of the DDX23 deletion mutants. Full-length DDX23 had an exclusively nuclear localization, with a nuclear speckled pattern, which we also observed with the smaller deletions (Δ77, Δ127). In contrast, deletion of the first 327 residues resulted in a diffused nuclear localization, lacking the characteristic speckled pattern, and with occasional cytoplasmic localization (Fig. 4C). These localization patterns indicate a role for the Nt domain of DDX23 in proper subnuclear targeting, which might involve interaction with SRSF1 and/or other SR proteins.

DDX23 Interacts with Additional SR Proteins. SR proteins have related and, in some cases, overlapping functions, so we tested whether DDX23 can also interact with other family members. We tested seven additional SR proteins using BiFC and found that DDX23 interacted with SRSF3, SRSF4, SRSF5, SRSF6, SRSF7, SRSF9, and SRSF10 (Fig. 4D and SI Appendix, Fig. S10). Therefore, there may be a conserved function(s) among members of the SR protein family that involves direct interaction with DDX23. These results are consistent with our observation that RRM1 and the phosphorylated RS domain of SRSF1 can each efficiently pull down DDX23 and point to other SR proteins harboring these domains as candidates for interaction with DDX23.

Discussion

SRSF1 is an essential splicing factor that has been implicated in cancer, autoimmune diseases, and tautopathies. Therefore, further investigation of SRSF1 functions and its potential interactors can lead to unique insights into SRSF1 dysregulation in disease (45). Our previous study used IP-MS and computational approaches to study the SRSF1 interactome (15). However, many of the interactors we detected using BioID2 were not previously observed using IP-MS. Several advantages of proximity labeling may account for this observation. First, the labeling takes place in live cells, capturing interactions in the intact cellular compartments. Second, proximity labeling allows the detection of dynamic and transient interactions, such as those that occur during spliceosome assembly and splicing catalysis, which may be hard to capture in a copurification assay. Third, the protein isolation for proximity labeling is done under highly denaturing conditions, improving the solubility and recovery of labeled proteins and eliminating interference from bound nucleic acids. However, the proximity-labeling interactome reflects adjacency (up to 10 Å), so not all recovered proteins are necessarily part of a complex with SRSF1 (46). This is why complementary methods, such as BiFC and recombinant-protein interaction assays, can help to validate and map relevant protein–protein interactions.

Our datasets reflect SRSF1’s varied and dynamic subcellular localization, as well as SRSF1’s roles in splicing and in additional pathways like NMD and enhancement of translation (45). The detection of spliceosomal proteins involved in all stages of splicing reinforces the idea that SRSF1 remains associated with the
Fig. 4. Possible functional roles of DDX23’s Nt-domain and conservation of the interaction with other SR protein family members. (A) Representative His PD assay using the indicated versions of His-TrxA-DDX23 (41 to 77). (B) In vitro splicing reactions using nuclear extract supplemented with the indicated versions of recombinant DDX23 protein. Calculated PSI values are reported for each condition, with their indicated SD. Triplicate samples were used for unpaired t tests. **P ≤ 0.01, ns = not significant. (C) Live fluorescence microscopy imaging of HEK293T cells transfected with the indicated mVenus-fused DDX23 constructs. Magnification was 40× (Top images). Scale bar represents ~15 μm. Cropped images (Bottom) are focused on one or two nuclei, so that the nuclear distribution of DDX23 can be observed more clearly. (D) BiFC results for the indicated proteins, plotted as Forward scatter (linear scale) vs. Green fluorescence (logarithmic scale). The vertical line indicates the threshold for cells that were considered mVenus-positive. The bar graphs below each panel represent %mVenus-positive cells and mean green fluorescence, respectively, for DDX23-mVCt coexpressed with Nt-mVenus vs. DDX23-mVCt coexpressed with SRSF3/5/6-Nt-mVenus. An unpaired t test was performed to compare the %mV+ cells and MGF values between the experimental conditions and the controls. *P ≤ 0.05; ****P ≤ 0.0001. Error bars represent SD.
spliceosome and later accompanies the spliced mRNA to the cytoplasm. Further characterization of these dynamic interactions should help clarify the molecular mechanisms underlying SR protein functions in pre-mRNA splicing and additional cellular functions of SRSF1.

Previous studies suggested that SR proteins play a role during the prespliceosome to spliceosome transition, helping to recruit the tri-snRNP to the spliceosome (47). We provide experimental evidence of a direct physical interaction between SRSF1 and DDX23, an essential protein that drives the pre-B to B complex transition, where stabilization of the tri-snRNP occurs. We detected the DDX23-SRSF1 interaction in live cells by two different approaches, as well as in vitro between purified recombinant proteins. The interaction involves several charged, conserved segments of the Nt-domain of DDX23 that function as alternative contact surfaces with SRSF1.

In SRSF1, the interaction involves two domains: RRM1 and the phosphorylated RS domain. The RS domain was previously characterized as a protein-interaction domain (10). Regarding RRM1, several studies likewise identified this domain as a mediator of protein–protein interactions, in addition to binding RNA (48–51).

In the Nt domain of DDX23, most of the RS repeats are relatively dispersed, except for a nine-residue stretch of RS repeats (S57–S65). A recent study of SRSF1 identified a negatively charged surface of RRM1 as a potential mediator of interactions with RS-repeat-containing peptides that is conserved in all SR protein family members (7). In the present study, mutation of the DDX23 RS repeats to GA (9GA mutant) decreased the interaction with RRM1, confirming that the long RS-repeat region contributes to SRSF1 binding through interactions with RRM1. However, binding was not completely abolished by the mutations, suggesting that other regions of the FL-DDX23 protein also contribute.

In contrast, in the context of the minimal DDX23 version (S41–R77), the 9GA mutation completely abolished binding to FL-SRSF1, whereas in the context of FL-DDX23, it had no effect. These results suggest that the flexibility of the disordered Nt domain of DDX23 permits multiple interactions with SRSF1 in the context of the whole protein. This notion is also confirmed by our results with the DDX23-Δ(S41–R77) mutant, which still binds SRSF1. We suggest that in the absence of the conserved (S57–S65) RS repeats, other charged repeats within the Nt domain of FL-DDX23 can interact instead, reflecting the intrinsic flexibility of the Nt domain.

Phosphorylation of the RS repeats of DDX23 was previously reported and proposed to be important for DDX23’s association with the tri-snRNP and for spliceosome B complex formation. It is not clear whether DDX23 is already phosphorylated when it associates with U5 snRNP or whether it becomes phosphorylated after tri-snRNP assembly (44). In addition, it seems plausible that DDX23 needs to be dephosphorylated to dissociate from the tri-snRNP after U1’s departure from the spliceosome. Our results suggest that the interaction with SRSF1 mediated by the RS repeats in DDX23 is inhibited by phosphorylation. Therefore, it is possible that regulation of the interaction with SRSF1 is mediated by the phosphorylation/dephosphorylation dynamics of DDX23, which would control the recruitment of DDX23 and its subsequent exit during spliceosome remodeling.

Several regions within the DDX23 Nt domain interact with spliceosome components, including a PRP8–SNU114 binding region (Y286 to H356), and a helical region (E158–L255) that connects BRR2 with PRP8. The Nt region also comprises the so-called 5′ exon mimic (D269–S283), a disordered loop that occupies the position of the 5′ exon in the pre-B spliceosome and is displaced by the 5′ exon upon DDX23’s departure (52) (Fig. 5A). Our results indicate that the S41–R77 region of DDX23 is important for spliceosome incorporation and that the M1–R77 region can interact with SRSF1 in multiple ways. This region also undergoes serine phosphorylation, which can affect the interactions with components of the tri-snRNP and with SRSF1.

Larger deletions of the DDX23 Nt domain resulted in striking changes in the nuclear localization of DDX23 (Δ327 mutant). These changes may reflect the loss of protein–protein interactions that allow DDX23 to properly engage with spliceosome components, or to participate in liquid–liquid phase separation. A recent report on DDX23 from C. elegans showed that—similar to our observations with the human protein—the Nt domain is involved in the formation of nuclear foci, which involves liquid–liquid phase separation (53). Therefore, in higher eukaryotes, the presence of the extended Nt domain could play a role in several aspects of DDX23 regulation, from localization and phase separation, to phosphorylation and spliceosome incorporation. Dissecting the protein interactions involving this domain can therefore reveal key regulatory mechanisms that may be exclusive to higher eukaryotes.

Both human DDX23 and yeast Ptp28p have very low RNA-dependent ATPase activity in vitro, and their presumed RNA-helicase activity has never been detected. Therefore, it is believed that key interactions between this enzyme and other spliceosome components are essential to achieve proper activation of DDX23 (22, 54). A recent study identified an interaction between Ptp28p and phosphorylated Npl3p (P-S411) that is important for Ptp28p’s ATPase activation and recruitment to the budding yeast spliceosome (55). Given that SR proteins are evolutionarily related to Npl3p, our results suggest that the reported contact observed in yeast is conserved in higher eukaryotes as a contact between DDX23 and SR proteins. We performed preliminary assays to investigate whether SRSF1 could stimulate DDX23’s very low basal RNA-dependent ATPase activity, but we did not observe activation under the conditions tested. However, one key difference between yeast Ptp28p and human DDX23 is the lack of the RS-like Nt domain in the yeast protein. In addition, yeast Npl3p has RRM’s with clear homology with those in SR proteins, but it lacks an RS domain (56), which in SRSF1 enhances the interaction with DDX23, as shown in the present study. Therefore, notwithstanding the conserved interaction between Ptp28p–Npl3p and DDX23–SRSF1, the underlying interaction mechanisms are different, and so the functional consequences may also differ, such that additional protein interactions with human DDX23 may be required to stimulate its ATPase activity.

In addition to its crucial role in the pre-B to B spliceosomal–complex transition, yeast Ptp28p was implicated in the earliest stage of spliceosome assembly (57). As SRSF1 also participates in early spliceosome assembly (58), it seems possible that the initial interaction with DDX23 could take place at an early stage. However, information derived from the cryo-EM structures of the human pre-B and pre-Bcret spliceosomes places DDX23 in the pre-B complex and SRSF1’s RRM2 in the pre-Bcret complex in the same vicinity, proximal to the 5′s (19, 52) (Fig. 5 B and C). In addition, the fact that SRSF1 binds U1 snRNA and is involved in U1 snRNP recruitment, whereas DDX23 promotes U1 snRNP release, functionally links these two proteins and suggests that their physical interaction could help coordinate these sequential events, either by recruiting DDX23 or affecting the conformation of its Nt domain. Further studies, particularly structural approaches and analysis of DDX23 regulation by phosphorylation, will be necessary to fully understand the functional implications of this interaction.
Materials and Methods

Proximity Labeling and Streptavidin Pulldown. Details about the generation of cell lines used for proximity labeling are in SI Appendix, Supplementary Methods section. For proximity labeling, we performed a 50-μM biotin pulse for 24 h, followed by streptavidin PD under denaturing conditions (Lysis buffer: 50 mM Tris-HCl pH 7.5, 8 M urea, 1 mM DTT, 1× protease inhibitor cocktail (Roche)). We subjected the washed beads corresponding to triplicate samples for each condition to on-bead trypsin digestion and analyzed the resulting peptides by LC/MS, followed by label-free quantification. We detected a total of 2,706 proteins, and for each one, we extracted the peptide ion chromatogram (XIC) and calculated the area under the curve ratio between SRSF1 samples and the GFP control. We filtered the proteins by their combined abundance across samples and set a lower threshold of 6 × 10^7, yielding 1,699 candidates for further analysis.

BioC. We seeded HEK-293 cells in 6-well plates (5 × 10^4 cells/well) and transfected them with the BioC vectors at 70% confluence. For transfection, we mixed 100 μL of opti-MEM with X-treme gene 9 and the corresponding BioC plasmid. We normalized the amount of plasmid to use by the size of the plasmid, as follows: ng plasmid = 0.059 x length of plasmid (bp), and added 3 μL of X-treme gene 9 per μg of plasmid. After 18 h, we imaged live cells by fluorescence microscopy, collected them using 0.25% trypsin (500 μL), and resuspended them in 13 mL of DMEM. We transferred 200 μL of cells into 96-well round-bottom plates and analyzed them by flow cytometry (Guava). We analyzed eight technical replicates per transfection in parallel and compared them to an EGFP transfection control. We repeated each transfection at least 3 times. We analyzed the flow cytometry data using Guava Soft 4.5, with InCyte or Express pro. We gated the cells from a Forward versus Side scatter plot and obtained the percentage of mVenus-positive cells and mean green fluorescence values, which we plotted on Graphpad Prism.

His Pulldown Assay. Details of expression and purification of recombinant proteins are in SI Appendix, Supplementary Methods section. We incubated with gentle rotation 10 μg of His-tagged protein for 1 h at 4 °C with IMAC beads (Chelating Sepharose Fast Flow beads charged with Ni2+) equilibrated in Pulldown buffer (50 mM Tris-HCl pH 7.5, 200 mM NaCl, 2 mM MgCl2, 0.5% CHAPs, 20 mM imidazole), in the presence of 10 μg BSA and, where indicated, 10 units of benzonase nuclease (Millipore Sigma). We then added 20 μg of untagged protein and incubated the suspension for an extra 2 h at 4 °C. Next, we spun the beads for 5 min at 500 × g and washed them three times with 500 μL of Pulldown buffer for 3 min, with rotation at 4 °C. After the last wash, we removed all the remaining buffer and resuspended the washed beads in 20 μL of Laemmli buffer with 5 mM EDTA. We heated the samples at 95 °C for 10 min and analyzed the proteins by SDS-PAGE and Coomassie Blue staining.

Sequence Alignment. We retrieved the protein sequences of human DDX23 and its orthologs from the Uniprot database, visualized them in Seaview, and aligned them using Muscle (60). We generated a structural sequence alignment using ESPript 3.0 (61).

In Vitro Pre-mRNA Splicing Assay. We transcribed and gel-purified the AdML pre-mRNA from BamHI-digested AdML-M3 plasmid (Addgene #11244) as described in SI Appendix, Supplementary Methods section. Pre-mRNA splicing was carried out with
Helix cell nuclear extract. (Ipracell). For each version of DDX23 [DDX23-27. F. Pelisch et al. The cellular EJC interactome reveals higher order mRNP structure and an EJC loop-formation mechanism in splicing factor SRSF1 and its role in mitosis. Elife 10, e65104 (2021).]

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Graphics. We generated protein-protein structure images with Pymol 1.7.4.0, and protein-domain graphics with IBS illustrator (62).

Data, Materials, and Software Availability. Interactome data have been deposited in ProteomeXchange, PRIDE database (Project accession: PXD0048180, and is included as Dataset S1 (63). All other data are included in the manuscript and/or supporting information.


