SAM68 interaction with U1A modulates U1 snRNP recruitment and regulates mTor pre-mRNA splicing

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ABSTRACT

Src associated in mitosis (SAM68) plays major roles in regulating RNA processing events, such as alternative splicing and mRNA translation, implicated in several developmental processes. It was previously shown that SAM68 regulates the alternative splicing of the mechanistic target of rapamycin (mTor), but the mechanism regulating this process remains elusive. Here, we report that SAM68 interacts with U1 small nuclear ribonucleoprotein (U1 snRNP) to promote splicing at the 5′ splice site in intron 5 of mTor. We also show that this direct interaction is mediated through U1A, a core-component of U1snRNP. SAM68 was found to bind the RRM1 domain of U1A through its C-terminal tyrosine rich region (YY domain). Deletion of the U1A-SAM68 interaction domain or mutation in SAM68-binding sites in intron 5 of mTor abrogates U1A recruitment and 5′ splice site recognition by the U1 snRNP, leading to premature intron 5 termination and polyadenylation. Taken together, our results provide the first mechanistic study by which SAM68 modulates alternative splicing decision, by affecting U1 snRNP recruitment at 5′ splice sites.

INTRODUCTION

Alternative splicing is a highly regulated event requiring an impressive amount of ribonucleoprotein complexes and associated factors (1,2). In this process, intervening sequences are excised out from nuclear pre-messenger RNA (pre-mRNA) by the macromolecular machinery called the spliceosome (3). The recognition of the 5′ splice sites by U1 small nuclear ribonucleoprotein (U1 snRNP) defines the initial stages of spliceosome assembly. U1 snRNP along with U2, U4, U5 and U6 snRNPs forms the major spliceosome, the core machinery that catalyzes splicing reactions in eukaryotes (4). Although core spliceosomal assembly and its catalytic activity are rather well defined, an increasing number of accessory spliceosomal proteins modulate its activity and specificity, thereby making alternative splicing a highly regulated process (5). The main challenge for efficient intron splicing is the recognition of the 5′ and 3′ splice sites. This is mainly achieved by U1 snRNP (6,7), U2 snRNP and U2AF (8,9). These spliceosome components drive the assembly of the formation of the early spliceosome called complex E (10,11). Now it is well known that regulatory factors can bind sequences neighboring the 5′ splice site to prevent or promote U1 snRNP binding (12). Increasing evidence highlight the importance of RNA-binding proteins in facilitating U1 snRNP recognition of 5′ splice sites and regulating alternative and constitutive splicing. These include FUS (13,14), SF2 (15,16), TIA-1 (17), RBM24 (18), hnRNPs (19,20) and SAM68 (21–24).

Src associated in mitosis of 68 kDa (SAM68), a 443-amino acid polypeptide, belongs to the signal transduction and activation of RNA family of RNA-binding proteins (RBPs) and was identified as a substrate of phosphorylation by c-SRC during mitosis and cellular transformation (25,26). SAM68 was shown to be able to bind mRNA (27), as well as DNA, upon its methylation (28). The multi-functionality of SAM68 can be rightly attributed to its modular organization. The RNA binding activity of SAM68 is confined to its highly conserved GSG domain, comprising of GRP33/GLD-1 domain, comprising of hnRNP KH domain flanked on its N terminus by 80 amino acids (NK) and its C-terminus of 30 amino acids (CK), respectively (29,30). It has been demonstrated by X-ray crystallography that the NK region is required for the RNA-dependent homodimerization of SAM68 (31). In addition, SAM68 has six proline rich sequences on either side of GSG domain along with a tyrosine rich C-terminus that were shown to be targeted by various signaling pathways (32–34). The tyrosine phosphorylation of SAM68 as well as its interaction with SH2 binding proteins has been shown to
impair its affinity for RNA (23,33). Thus, SAM68 is a versatile adaptor and nucleic acid docking protein whose activity is modulated by cell signaling.

SAM68 is known to bind single-stranded U/A-rich mRNA molecules, mainly through U(U/A)AA repeats (35). The RNA-binding activity of SAM68 was shown to be involved in various aspects of mRNA processing including alternative splicing (29). This was initially shown following ERK1/2 signaling pathway activation, which promoted a SAM68-induced inclusion of the variable exon5 in CD44 (24,33). SAM68 has been involved in the alternative splicing of mRNAs implicated in neurogenesis (36,37), adipogenesis (21,38–40), spermatogenesis (41,42) and epithelial-mesenchymal transition (43). SAM68 regulated alternative splicing was further highlighted with SMN2 (44), BCL-x (22), Cyclin D1 (22) and mTor (21) pre-mRNA transcripts. While the mechanisms underlying the splicing of SMN-2, BCL-x and Cyclin D1 are becoming clearer, the mechanisms regulating SAM68-induced alternative splicing of mTor pre-mRNA remains elusive.

mTOR is a central regulator of cell homeostasis, growth, proliferation and survival (45). Its dysregulation occurs in many human diseases such as cancer, obesity, Type 2 diabetes and neurodegeneration (45,46). Hence, it is crucial to understand the mechanism of SAM68 regulated mTor pre-mRNA splicing. Using the Sam68+/− mouse as models, we had previously unveiled a novel role of SAM68 in driving alternative splicing of mTor pre-mRNA (21). We found that impairing SAM68 binding to its target elements found near the 5’ splice site of intron 5 decreases the expression of full-length mTor mRNA by increasing intron 5-induced premature termination leading to the production of a shorter mRNA termed mTor5, with no detectable protein product. The production of mTor5 is increased in Sam68+/− mouse tissues indicating that SAM68 mediates the balance between both isoforms. As a result, Sam68+/− mice have decreased mTOR protein levels and attenuated mTORC1 and mTORC2 activities. RNA-binding assays determined that SAM68 binds multiple U/A-rich sequences distributed throughout intron 5 and enhances splicing at the upstream exon/intron junction (21). These observations suggest that SAM68 has the ability to regulate an important mTor pre-mRNA alternative splicing checkpoint, though the underlying mechanism remains unknown.

Here, we investigated the mechanism by which SAM68 modulates mTor pre-mRNA splicing. First, we found that SAM68 was detected in the immunoprecipitates of the core components of U1 snRNP, namely U1A and U1-70K. Reciprocal immunoprecipitation with Flag-tagged SAM68 showed enrichment of U1 snRNP. Concomitantly, purified recombinant SAM68 can capture U1 snRNP through direct interaction with U1A. Domain mapping experiments revealed that the tyrosine rich C-terminal region of SAM68 (YY domain) was sufficient to interact with U1A. Using endogenous RNA immunoprecipitation assays, we found that SAM68 can recruit U1 snRNP to the 5’ splice site of mTor intron 5. Thus, these results provide the first mechanistic insight on how SAM68 regulates mTor pre-mRNA alternative splicing and could unveil a broader regulatory function of SAM68-mediated 5’ splice site recognition.

MATERIALS AND METHODS

Plasmid constructions

pGEX-6P3-SAM68-Flag and pGEX-6P3-Sam68-Flag were constructed by inserting full-length human and mouse SAM68 coding sequence (cds) into pGEX-6P3 (GE Healthcare) with N-terminal GST tag, a PreScission protease cleavage site (see below ‘Protein purification and GST pulldown’ section) between GST and SAM68. Due to the high sequence homology between human and mouse SAM68, both constructs were produced using EcoRI-SAM68-F and NotI-SAM68-R. Flag tag was then inserted at the C-terminus of SAM68 by annealing the oligos, SacI-Flag-F and SacI-Flag-R, and inserting the adaptor at SacI sites of the plasmid. pGEX-6P2-U1A-His was generated by inserting U1A cds, obtained by polymerase chain reaction (PCR) from HEK-293T total cDNA using EcoRI-U1A-F and Sall-U1A-R at EcoRI-Sall sites. cDNA was amplified from total RNA of HEK-293T using Superscript VILO Master mix (Invitrogen). pGEX-6P2-U1C-His was sub-cloned by PCR from pGEX-2TK-U1C using EcoRI-U1C-F and XhoI-U1C-R and inserted at EcoRI-XhoI sites. pcDNA-Flag-SAM68 and pcDNA-Flag-Sam68 were constructed by inserting corresponding cds, obtained by PCR from pGEX-6P3-SAM68-Flag and pGEX-6P3-Sam68-Flag using EcoRI-SAM68-F and NotI-SAM68-R at EcoRI-NotI sites.

pcDNA-Flag-SAM68y185N and pcDNA-Flag-SAM68y229f were generated by swapping the 679 bp, Agel-XbaI fragment from pcDNA mCherry-SAM68y185N and pcDNA mCherry-SAM68y229f, respectively, to pcDNA-Flag-SAM68 WT. pcDNA-Flag-SAM68-Nter and pEGFP-Sam68-Nter were generated by deletion PCR with SAM68-F and SAM68-Nter-R primers using pcDNA-Flag-SAM68 and pEGFP-SAM68 as templates, respectively. pcDNA-Flag-SAM68-Cter was generated from pcDNA-Flag-SAM68 by PCR using EcoRI-SAM68-Cter-F and NotI-SAM68 Cter-R and cloning the ampiclon at EcoRI-Sall sites of pcEGFP-C1. pcEGFP-C1-SAM68-C1 was obtained by deletion PCR using SAM68-F and SAM68-C1-R, respectively. pcEGFP-SAM68-C2 to C5 were generated by cloning the PCR ampiclons obtained using the reverse primer, Sall-SAM68-R and forward primers namely EcoRI-SAM68-C2, C3, C4, C5 at EcoRI-Sall sites of pcEGFP-C1. pcEGFP-SAM68-NLS was obtained by deletion PCR using EcoRI-SAM68-NLS-F and EcoRI-SAM68-NLS-R. pLKO-shSAM68 was generated by annealing and inserting the oligos, shSAM68-F and shSAM68-R, at Agel-EcoRI restriction sites of pLKO1 (Addgene plasmid #8453). Primer sequences and generated plasmid are listed in supplementary Table S1.

Antibodies, western blotting and immunoprecipitation

The following antibodies were used in this study: anti-Flag (1:2000, 2368S, Cell Signaling Technology), anti-U1–70K (1:1000, 05–1588, EMD-Millipore), anti-U1A (1:1000, ab55751, abcam), anti-U1C (1:1000, ab192028, Abcam), anti-U1A (1:1000, ab55751, abcam), anti-U1C (1:1000, ab192028, Abcam),
anti-SAM68 (1:2000, AD-1, gift from Dr Stéphane Richard), anti-GAPDH (1:2000, MM-0163-P, Médimabs), anti-mTOR (1:1000, 2983S, Cell Signaling Technology), beta-actin (1:1000, 8457L, Cell Signaling Technology), anti-GFP (1:2000, ab290, Abcam) and anti-His (1:1000, 12698S, Cell Signaling Technology). Enzymatic immunoprecipitation was done by immobilizing anti-U1–70K, anti-U1A or control IgG-Mouse (Santa Cruz Biotechnology, sc-2025) on Protein A/G PLUS-Agarose beads (Santa Cruz Biotechnology, #sc-2003). HEK-293T cells were harvested, lysed for 10 min at 4°C in 1× phosphate-buffered saline (PBS), pH 7.4, 1% Triton-X-100, 150 mM NaCl, RNaseA (10 mg/ml; Sigma, #R5503) and 1× protease inhibitor complete ethylenediaminetetraacetic acid (EDTA)-free (Roche). The cell lysates were sonicated five times for 30 s with a Bioruptor ultrasonic cell disruptor and centrifuged at high speed for 30 min at 4°C to remove cell debris. For enzymatic immunoprecipitation, the respective antibodies conjugated to Protein-A/G PLUS-Agarose beads (Santa Cruz Biotechnology, sc-2003) were added to pre-cleared cell lysates. Following 1 h at 4°C, the beads were washed several times with lysis buffer and the immunoprecipitates were eluted with Laemmli buffer. Flag-tagged proteins were immunoprecipitated with Flag-M2 affinity beads (Sigma, #A2220). GFP-tagged proteins were immunoprecipitated with homemade GFP-Trap-A beads.

RNA immunoprecipitation and RT-PCR

RNA from 50% of the Flag-Sam68 and Flag-YFP immunoprecipitates was isolated using TRIzol™ Reagent (Invitrogen, #15596-018) and reverse transcribed using SuperScript™ VILO™ MasterMix (Invitrogen, #11755-050) or M-MuLV reverse transcriptase, according to the manufacturer’s instructions. Extracted RNA was incubated with random hexamers, oligo-dT or gene-specific primer for first strand cDNA synthesis at 25°C for 10 min, 42°C for 1 h and 85°C for 5 min. One-fifth of the reaction product was amplified for U1snRNA transcript using U1snRNA-F and U1snRNA-R primers and GAPDH mRNA using GAPDH-F and GAPDH-R primers, respectively. Primer sequences are listed in Supplementary Table S1.

RNA-binding assay

Purified Sam68 or U1A (stored in 20 mM Tris–HCl, pH 7.4, 200 mM NaCl, 10% Glycerol, 1 mM dithiothreitol) was added at the indicated concentration to a mix containing 10 nM of 32P-labeled U1 snRNA in 1× RNA binding buffer (50 mM Tris–Cl, pH 7.6, 200 mM potassium acetate, 5 mM MgCl2, 2.5 mM dithiothreitol) supplemented with 0.25 μl of RNasin Ribonuclease Inhibitor. The mix was incubated at room temperature for 15 min, then complexes were fixed with 0.5% glutaraldehyde for 10 min at room temperature. The samples were loaded onto a 5.5% Tris/Borate/EDTA (TBE) 1× acrylamide (29:1) gel and run at 150 V for 2 h and 30 min at 4°C, dried onto DE81 filter paper, then visualized by autoradiography. Quantifications were performed on a FLA-5100 phosphorimager system (Fujifilm), and statistics were analyzed with Prism.

Protein purification and GST pulldown

Competent Escherichia coli BL21 DE3 Codon plus RP strain (Stratagene, Agilent technologies # 230255) were transformed with pGEX-6p3-SAM68-flag, pGEX-6p3-Sam68-flag, pGEX-6p3-SAM68-Nter, pGEX-6p3-Cter, pGEX6p2-U1A-His, pGEX6p2-U1–70K-His, pGEX6p2-U1C-His and empty vectors, namely pGEX6p3-flag and pGEX-6p2-His, respectively. Single colony of each construct was then grown in LB media at 37°C until desired density and then induced with 0.3 mM isopropyl B-D-1-thiogalactopyranoside (IPTG) at 30°C overnight. Bacterial pellets were collected and lysed with 50 mM Tris–Cl, pH 7.5, 1% Triton-X-100, 200 mM NaCl, 20% glycerol, 5 mM MgCl2, 1 mM dithiothreitol (DTT) for 30 min, sonicated and centrifuged at 17 000 g for 30 min 4°C. GST fusion proteins were bound to 500 μl of washed and equilibrated Glutathione agarose beads (Sigma, #G4510). For GST-SAM68-flag, the GST-tag was removed by 20 units of PreScission Protease (2U/μl; GE Healthcare, #27-0843-01) at 4°C overnight in 50 mM Tris-HCl, pH 7.0, 150 mM NaCl, 1 mM EDTA, 1 mM DTT. Following this step, SAM68-flag was immunoprecipitated with flag-M2 affinity beads (Sigma, #A2220) for 1 h at 4°C, eluted with 9 μg of 3X FLAG® Peptide (4 mg/ml; Sigma, #F4799), and pooled elutes were dialyzed in 1× PBS, 200 mM NaCl, 0.05% Triton-X-100, 20% glycerol. Aliquots of the preparation were run on sodium dodecylsulphate-polyacrylamide gel electrophoresis (SDS-PAGE) and stained with Coomassie Brilliant Blue to validate preparation purity (Supplementary Figure S1).

For the NMR part of this study, GB1-hSAM68 (C2), GB1-U1A (1-282), U1A linkerRRM2 (156–282) and U1A RRM1 (1-126) were expressed under the control of lactose inducible promoters in E-coli BL21 DE3 at 37°C during 4 h in presence of 1 mM IPTG. To achieve isotope labeling, cells were grown in M9 minimal medium complemented with 15N-NH4Cl and/or 13C-glucose. Bacterial pellets were suspended in buffer A (10 mM HEPES, pH 7.6, 500 mM NaCl, 0.5 M Urea, β-mercapto-ethanol 2.8 mM) in the presence of lysozyme and DNAse I (0.01 mg/ml each). Cells were lysed using a microfluidizer by five cycles at 15 000 psi and the lysates were clarified by centrifugation (30 000 g, 40 min). Proteins were then purified using Ni2+-affinity chromatography and eluted with a gradient of imidazole. The C-terminal histidine tag of U1A RRM1 and U1A RRM2 were cleaved by thrombin (10 U/mg of purified protein, 6 h at room temperature in buffer A), while GB1-hSAM68 (C2) and GB1-U1A were kept as fusion proteins. All the proteins were further purified by size exclusion chromatography in 10 mM sodium phosphate, pH 6.8, NaCl 50 mM. For GST pulldown, 300 ng of purified recombinant hSAM68-Flag was incubated with 150 ng of glutathione bound GST-tagged U1–70K-His, U1A-His, U1-C-His or with negative control GST-His in binding buffer (50 mM Tris, pH 7.5, 200 mM NaCl, 10% glycerol, 0.5% Triton-X-100, 10 mg/ml RNaseA), supplemented with 1× protease inhibitor complete EDTA-free (Roche). The reaction volume was made up to 300 μl in total and incubated for 1 h at 4°C. Beads were washed and bound proteins were resolved by SDS-PAGE and analyzed by western blot.
**Biotinylated probes synthesis and RNP pulldown**

In vitro 5 baits with and without 5′ splice sites and either Wild-Type (WT) or Mutated (Mut) SAM68-binding sequence were cloned in pcDNA-Neo. The baits were in vitro transcribed from EcoRI linearized plasmids using T7-Hi-Scribe kit (Invitrogen) according to manufacturer’s instructions. After DNase I treatment, the reaction mixtures were column purified and 3′ end labeled with UTP-Biotin (Roche) with poly-U-polymerase according to manufacturer’s instructions. Baits were immobilized on streptavidin agarose beads and were washed three times with binding buffer (20 mM HEPES, pH 7.9, 200 mM NaCl, 10% glycerol, 0.5% Triton-X-100, RNase inhibitor, 1× protease inhibitor). The beads were blocked with 100 μg/ml of bovine serum albumin for 30 min at 4°C and washed again in binding buffer. Immobilized baits were incubated with 1 μg of either purified SAM68-Flag or GST-Flag for 30 min at 4°C, then shSAM68 HEK-293T cell lysates were added for 1-h incubation at 4°C. Beads were washed in binding buffer and the retained proteins were eluted in Laemml buffer, run on SDS-PAGE gels and blotted using Flag and U1A antibodies.

**NMR spectroscopy**

NMR data were recorded at 313 K using the 500 MHz Avance III or the 600 MHz Avance III (Bruker), both equipped with cryo-probe. Data were processed with Topspin (Bruker) and analyzed with CARA. Sequential assignment of hSAM68(C2) was deduced from the analysis of classical triple resonance experiments (3D HNCACB, 3D CACB(CO)NH, 3D HNCO and 3D HN(CA)CO). Uniformly 15N labeled hSAM68 (C2) was titrated with unlabeled GB1-U1A, U1A RRM1 (1-126) or U1A linker-RRM2 (156-282) and the formation of the binary complexes was monitored by measuring 2D 15N-1H HSQC spectra after each addition. Reverse NMR titrations of uniformly 15N labeled U1A versions by unlabeled GB1-SAM68 (C2) were performed using a similar strategy. Chemical shift perturbations were plotted onto the surface representation of the structure of the free form of the RRM1 of U1A (47).

**In vitro Xrn-1 protection assay**

XbaI linearized pcDNA 3.1-mTor minigene plasmid was used for in vitro transcription. Plasmid DNA templates were eliminated by DNaseI treatment followed by column purification of the RNA template. RNA templates were either used directly for 5′ Xrn-1 exonuclease assay or 3′ end labeled with poly-U-UTP-biotin using poly-U-polymerase and bound to streptavidin agarose beads. Streptavidin-bound RNA templates were incubated with WT MEFs cell extracts or Sam68−/− MEFS cell extracts supplemented with either in vitro purified GST-Flag or mSAM68-Flag for 30 min at 4°C in binding buffer (10 mM Tris–HCl, pH 7.9, 10 mM MgCl2, 50 mM NaCl, 0.5% Triton-X-100, 1 mM DTT). 5′ monophosphate RNA templates were then treated with 1U of Xrn-1 (1000 U/ml, NEB, M0338L) for 1 h at 37°C in reaction buffer. The reaction was stopped by heating at 70°C for 10 min. cDNA was amplified using primer RRT and ampiclon corresponding to full-length bait was produced using forward (FSS) and reverse (RSB) primers while mSAM68 protected amplicon was produced using forward (FSB) and reverse (RSB) primers.

**RNA immunoprecipitation (RIP)**

RNA immunoprecipitation (RIP) assays were performed using a modified CLIP protocol (48). Nuclear fraction was isolated and lysed in 1× PBS, pH 7.4, 150 mM NaCl, 1% Triton-X-100, 0.5% sodium deoxycholate, 0.1% SDS, protease inhibitor cocktail for 30 min on ice. The extracts were treated with DNase I (10 U/μl, Roche, Cat. No. 04716728001) and RNase A (10 mg/ml, Sigma, R-4875) for 1 h at 4°C under rotation. The samples were diluted in binding buffer (1× PBS, pH 7.4, 150 mM NaCl, 1% Triton-X-100) and supplemented with RNase inhibitor. Samples were precleared and 10% of were saved for inputs, while the rest were used for immunoprecipitation using anti-U1A antibody (Abcam, mab55751) or rabbit IgGs (Jackson ImmunoResearch, 011–000-003). The inputs and immunoprecipitated samples were treated with 50 μg Proteinase K for 30 min at 65°C. RNA was then isolated and used for RT-qPCR analyses that were performed using primers e5-F and e5-R for exon5–intron 5 junction, e4-F and e4-R for exon4–intron4 junction and e37-F and e37-R for exon37–intron37 junction.

**In vivo splicing assays**

Endogenous mTor transcript premature termination and polyadenylation in wild-type (WT) mouse embryonic fibroblasts (MEFs) and Sam68−/− MEFs were rescued using lentivirus-mediated transduction of mouse Sam68 (WT) or mouse Sam68 (∆ARM). Total RNA was extracted using TRIzol® reagent and digested with DNase I to get rid of contaminating genomic DNA. After column purification, cDNA was amplified by reverse transcription using oligo-dT and M-MuLV reverse transcriptase. The following pairs of primers were used F(e4) and R(i5) to amplify mTor8 and F(e4) and R(e6) to amplify mTorExon4−6. A portion of cell lysates were resolved in SDS-PAGE and blotted with mTOR, SAM68 and GAPDH antibodies.

mTor genomic fragment spanning from exon4 to exon6 with corresponding introns 4 and 5 was cloned in pcDNA Neo and referred to as mTorExon4−6 minigene, hereafter. shSAM68 HEK-293T cells were co-transfected with mTorExon4−6 minigene and either Flag-YFP, Flag-hSAM68-WT or Flag-hSAM68ΔARM. Forty-eight hours post-transfection, total RNA was extracted using Trizol® reagent and digested with DNase I to get rid of contaminating plasmid DNA. After column purification, cDNA was amplified by reverse transcription with random hexamers using Superscript VILO according to manufacturer’s instructions. Cell lysates were resolved in SDS-PAGE and
show the presence of two stretches of high-affinity SAM68 used to amplify transcripts. Primer pairs GAPDH-F and GAPDH-R were used to amplify intron 5 Reverse were used to amplify intron 5 retained minigene. Hence, we reasoned that SAM68 could be interacting with the U1 snRNP, the spliceosomal component that recognizes the 5′ splice site, and thus be a determining factor for early splicing assembly within mTor intron 5 (4). While the initial discovery was observed in mice, we reasoned that any SAM68–U1 interaction would also be observed in human cells, as human SAM68 (hSAM68) is almost identical to mouse SAM68 (mSAM68).

To validate if SAM68 could interact with U1 snRNP, flag-tagged hSAM68 (flag-hSAM68) was expressed in HEK-293T cells depleted of endogenous SAM68 using lentiviral-shRNA targeting the 3′ UTR of SAM68 mRNA (shSAM68 HEK-293T) (Supplementary Figure S2A). Flag-tagged yellow fluorescent protein (flag-YFP) was used as negative control. Western blot analyses showed that the three core protein components of U1 snRNPs, namely U1–70K, U1A and U1C, efficiently co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B). RIP followed by RT-PCR using primers specific to U1 snRNA and GAPDH mRNA showed that U1 snRNA only co-immunoprecipitated with Flag-hSAM68, while undetected in Flag-YFP controls (Figure 1B).

Interaction of SAM68 with U1 snRNP is RNA independent

Given that SAM68 is an RNA-binding protein, we first assessed if the SAM68-U1 snRNP interaction was mediated through U1 snRNA. Sequence analyses showed the presence of a potential SAM68-binding site (AUAAUUU) upstream and partially within the Sm domain (49). We tested and compared the affinity of both Sam68 and U1A to the U1 snRNA. While SAM68 can indeed bind U1 snRNA, it showed minimal affinity for this RNA when compared to U1A, a bona fide U1 associated protein (Figure 1F and G). Considering that SAM68 has numerous preferential RNAs targets (21,27,50) and its low affinity for the U1 snRNA (Figure 1F), it would be highly unlikely that its association with the U1 snRNP is mediated by U1 snRNA.

To identify which protein component of the U1 snRNP mediates this interaction, recombinant hSAM68-Flag were incubated with cell extracts from shSAM68 HEK-293T cells in the presence or absence of RNase A. Treatment with RNase A had no effect on SAM68 association with U1 snRNPs (Figure 1F). It would be highly unlikely that its association with the U1 snRNP is mediated by U1 snRNA.
Figure 1. *In vivo* association of SAM68 with U1 snRNP. (A) Schematic representation of a portion of *mTor* pre-mRNA spanning from exon4 to exon6 (upper panel), with a close-up of the 5′ splice site and the subsequent SAM68-binding site (SB-1), as well as the cryptic polyadenylation signal that harbor SAM68-binding site (SB-A). (B) Co-immunoprecipitation of U1 snRNP with Flag-hSAM68. HEK-293T cells depleted of endogenous SAM68 (shSAM68 HEK-293T) were transiently transfected with Flag-hSAM68 or Flag-YFP (yellow-fluorescent protein), the latter serving as negative control. Flag-tagged proteins were immunoprecipitated using anti-Flag M2 agarose beads and immunoprecipitated proteins were detected with antibodies specific to U1–70K, U1A and U1C. β-Actin was used as negative control. Portion of the Flag-immunoprecipitates was used for RNA isolation and RT-PCR using U1 snRNA specific primers. GAPDH (glyceraldehyde 3-phosphate dehydrogenase) RNA was used as negative control of the RT-PCR made from the RNA immunoprecipitation. (C) Co-immunoprecipitation of endogenous hSAM68 with U1–70K. Immunoprecipitated proteins were detected with antibodies directed against SAM68 and U1–70K. β-Actin was used as negative control of immunoprecipitated proteins. (D) Co-immunoprecipitation of endogenous hSAM68 with U1A. Immunoprecipitated proteins were detected with antibodies directed against SAM68 and U1A. β-Actin was used as negative control of immunoprecipitated proteins. (E) Coomassie staining of purified human SAM68 and U1A. (F) RNA binding assay with purified SAM68 and labeled U1snRNA. Reactions contained 10 nM γ-p32 labeled U1snRNA in buffer with no protein (lane 1) or with purified SAM68 (lanes 2–5). Bottom panel: quantification from three independent binding experiments. Error bars represent the corresponding standard error. Unpaired two-tailed t-tests were used to compare the different concentrations of purified protein to the RNA only control. SAM68 P-values are 0.0014, 0.0005, <0.0001, <0.0001 in increasing order of SAM68 concentration. (G) RNA binding assay with purified U1A and labeled U1snRNA. Reactions contained 10 nM γ-p32 labeled U1snRNA in buffer with no protein (lane 1) or with purified U1A (lanes 2–5). Bottom panel: U1snRNA P-values = 0.0008, <0.0001, <0.0001, <0.0001 in increasing order of U1A concentration. **P-value < 0.005, ***P-value < 0.001.
Figure 2. In vitro Purified SAM68 associated with U1 snRNP in an RNA-independent manner. (A) In vitro purified hSAM68-Flag was added to shSAM68 HEK-293T cell lysates for 1 h at 4°C, in the presence or absence of 50 μg/ml RNaseA. hSAM68-Flag and associated proteins were immunoprecipitated using Flag-M2 affinity beads and treated further with RNaseA at 37°C for 30 min. Bound proteins were eluted with Laemmli and immunoblotted with antibodies specific to U1–70K, U1A and U1C. To assess RNaseA treatment efficiency, total RNA from shSAM68 HEK-293T was treated with either Mock or RNaseA for 30 min at 37°C, and the remaining total RNA was assessed on agarose gel. (B) RNA-binding defective mutant hSAM68 I184N interacts with U1 snRNP. shSAM68 HEK-293T were transiently transfected with Flag-hSAM68, Flag-hSAM68 I184N and Flag-YFP (negative control). The Flag-tagged proteins were immunoprecipitated using anti-Flag M2 agarose beads and immunoblotted with antibodies directed against U1–70K, U1A and U1C. (C) Association of hSAM68-Flag with U1 snRNP withstands high salt washes. Purified in vitro produced hSAM68-Flag was added to cell lysates of shSAM68 HEK-293T for 1 h at 4°C. Flag-M2 affinity beads were added to the reaction and left for 1 h at 4°C. The washes were done, by increasing salt concentration, from 150 to 500 mM of NaCl. Bound proteins were eluted with Laemmli and immunoblotted with antibodies directed against U1–70K, U1A and U1C. (D) SAM68 interacts with U1A in vitro. About 300 ng of purified hSAM68-Flag was incubated with 100 μg of glutathione-agarose bound GST-U170k-His, GST-U1A-His, GST-U1C-His and GST-His. Following washes, the beads were washed five times in binding buffer and the bound proteins eluted with Laemmli and immunoblotted using anti-Flag or anti-His antibodies.

SAM68 interaction with U1A is mediated through its tyrosine-rich (YY) domain

Being an adaptor protein, SAM68 comprises many protein–protein interaction domains such as SH3 binding proline-rich motifs and SH2 binding tyrosine-rich domain (28). In order to determine which domain was responsible for the association with U1A, we first truncated hSAM68 in two fragments (Figure 3A). The first fragment (N-term) contains the RNA-binding domain of hSAM68, spans from amino acids 1–280 and comprises the KH domain and proline-rich motifs (P0-P2) and the hSAM68 nuclear-localization signal (NLS). The second fragment (C-term) spanning from amino acids 281–443 comprises proline-rich motifs (P3-P5), the tyrosine-rich domain (YY), the SH2-binding tyrosine residues and the hSAM68 NLS. Both fragments were Flag-tagged at their N-terminus (Supplementary Figure S3A). Both flag-tagged hSAM68 fragments (N- and C-term), along with positive control Flag-hSAM68 and
Figure 3. SAM68 interaction with U1A is mediated through its C-terminal portion. (A) Schematic representation of C-terminus (aa. 1–280) and N-terminus (aa. 281–443) deletion domains of hSAM68 fused to flag. (B) shSAM68 HEK-293T cells were transiently transfected with Flag-SAM68 (N-term), Flag-SAM68 (C-term), Flag-SAM68 (FL) and flag-YFP (negative control). Forty-eight hours post transfection, the flag-tagged proteins were immunoprecipitated using anti-flag M2 agarose beads and immunoblotted with antibodies specific to U1–70K, U1A and U1C. (C) Schematic representation of full-length SAM68, C-terminus deleted SAM68 (NT, aa. 1–280), C-terminus truncated to proline rich C1 (aa. 269–364) and tyrosine rich C2 (aa. 365–443), C3 (aa. 370–443), C4 (aa. 385–443), C5 (aa. 385–443) and NLS (aa. 430–443). Fragments were fused to GFP tag at their N-terminus and all fragments had SAM68 NLS at their C-terminus. (D) GFP-Trap-A pulldown of GFP-tagged proteins. shSAM68 HEK-293T cells were transiently transfected with GFP, GFP-SAM68 (FL), GFP-SAM68 (NT), GFP-SAM68 (C1) and GFP-SAM68 (C2). Forty-eight hours post transfection, cells were lysed and GFP-Trap-A beads were used to pull down GFP-tagged proteins, and their association with U1A was validated by western blot using specific antibodies. (E) Primary amino acid sequence of the various deletion constructs of SAM68 YY domain (GFP-hSAM68 C2 to C5). Underlined indicates YXXY motifs in the YY domain. Also highlighted is the minimal ARM-binding region. (F) GFP-Trap-A pulldown of GFP-tagged proteins. shSAM68 HEK-293T cells were transiently transfected with GFP, GFP-SAM68 (C2), GFP-SAM68 (C3), GFP-SAM68 (C4), GFP-SAM68 (C5) and GFP-SAM68 (NLS). Forty-eight hours post transfection, cells were lysed and GFP-Trap-A beads were used to pull down GFP-tagged proteins, and their association with U1A was validated by western blot using specific antibodies. (G) U1A binds preferentially to the minimal ARM motif (YEGYEY) within the YY domain of SAM68. Flag-hSAM68 (FL) and Flag-hSAM68 (ΔARM) were transiently transfected in shSAM68 HEK-293T cells. Forty-eight hours post transfection, cells were lysed and Flag-tagged proteins were immunoprecipitated using anti-flag M2 agarose beads, and U1A association was assessed using U1A antibody. *: denotes an unspecific band.
negative control flag-YFP, were transfected in shSAM68 HEK-293T cells. Confocal images showed that while the N-terminal fragment remained largely cytoplasmic, it also partially localized to the nucleus, suggesting that the addition of the NLS allows efficient nuclear localization of the N-terminal fragment (Supplementary Figure S3A). Flag-tagged proteins were immunoprecipitated and immunoblotted with U1A antibody, while U1–70K and U1C antibodies were used as positive controls. As expected, full-length hSAM68 showed a strong association with U1 A (Figure 3B) as well as U1–70K and U1C, while they were not detected in Flag-YFP immunoprecipitates (negative control). The U1 snRNP components co-immunoprecipitated with the Flag-hSAM68 C-terminal fragment with as strong of an association as the full-length protein, but not the N-terminal fragment containing the KH RNA-binding domain (Figure 3B). While confocal immunofluorescence detection shows that the N-terminal fragment can also be located in nucleus due to the addition of the NLS (Supplementary Figure S3A), the partial cytoplasmic localization of this fragment could explain the decreased association with U1A. To assess this possibility, equal amounts of in vitro purified N-term and C-term fragments were immobilized and incubated with whole cell lysate taken from shSAM68 HEK-293T, validating our initial observation where U1A specifically binds the C-terminal fragment and not the N-terminal fragment (Supplementary Figure S3B). This result further corroborates our previous observation that the interaction is RNA-independent and implies that the SAM68 binds U1A through its C-terminal region.

To refine our mapping of the SAM68–U1A interaction, we truncated the C-terminal fragment of SAM68 into two smaller parts: GFP-hSAM68 (C1) and GFP-hSAM68 (C2) (Figure 3C). As the fragments were very short, we opted to clone them in frame with the larger, GFP-tag and performed a GFP-binding protein pulldown and as expected showed a strong nuclear localization by confocal imaging (Supplementary Figure S4A). Using these constructs, we found that like the full-length hSAM68, the C2 fragment of hSAM68 was associated with U1A, while the C1 fragment did not (Figure 3D). This initial interaction mapping suggests that the YXXY motif rich domain (Y domain) of SAM68, located in the C2 fragment (six YXXY motifs) but not C1, could be involved in the interaction with U1 snRNP. This domain consists of tyrosine-rich motifs involved in the association between SAM68 and the armadillo repeat domain (ARM) of the adenomatous polyposis coli (APC) protein (54,55). To assess the importance of this motif in the SAM68–U1A interaction, we further divided the GFP-hSAM68 (C2) domain in four different fragments with decreasing number of YXXY motifs (Figure 3E). GFP-hSAM68 fragment (C3) contains five YXXY motifs, (C4) has four, (C5) has two, while (NLS) has none. We found that the co-immunoprecipitation efficiency of U1A was directly dependent on the number of YXXY motifs, since the C4 fragment showed a slight decrease in association strength, while the C5 and the NLS fragments showed little or no association with U1A (Figure 3F). This change in association is unlikely to be related to mislocalization caused by the GFP moiety, since all the fragments showed a predominant nuclear localization (Supplementary Figure S4B).

Interestingly, there are six YXXY motifs within SAM68 YY domain, of which two successive YEGY motifs (SYEGYEGYYS) are defined as the minimal ARM interaction domain (54). Results obtained in Figure 3F suggested that losing this minimal motif could drastically affect the ability of SAM68 to bind U1A, like it was observed with APC. To validate this possibility, we proceeded with the deletion of this ten amino acid stretch (SYEGYEGYYS) within the Flag-hSAM68 (FL) construct. Similarly to what was observed with APC, deletion of the minimal ARM-binding motif (ΔARM) was enough to abrogate most of SAM68 association with U1A (Figure 3G), while it did not affect SAM68 affinity for its RNA target (Supplementary Figure S4C). Taken together, these results strongly indicate that SAM68 interaction with U1A is mediated through a specific sequence found in the YY domain (55).

**SAM68 interact with U1A–RRM1 domain**

To further confirm that SAM68 was directly interacting with U1A and to identify which part of both proteins interact, we used solution-state NMR spectroscopy. The hSAM68 (C2) fragment was isotopically labeled and resonances from backbone atoms were assigned using classical approaches. Analysis of the backbone chemical shifts revealed that in solution, the hSAM68 (C2) fragment adopts a random coil conformation without any secondary structure. Upon addition of unlabeled GB1-U1A, several resonances of the 15N-labeled hSAM68 (C2) experienced chemical shift changes (Figure 4A) that were reproduced with U1A RRM1 (1–126) but not by the C-terminal part of U1A containing RRM2 (156–282). Reverse NMR titration performed with 15N-labeled GB1-U1A revealed that upon addition of unlabeled GB1-hSAM68 (C2), the NMR signals from the RRM1 of U1A experienced strong line broadening and almost disappeared from the spectra (Supplementary Figure S5). However, when the 15N-labeled U1A RRM1 (1–126) was titrated with unlabeled hSAM68 (C2), several signals of the U1A RRM1 shifted from their initial positions (Figure 4B). The chemical shift perturbations observed on the N-terminal part of U1-A reveal an interaction surface with the C-terminal part of hSAM68 located between the edge of β-sheet surface (β2), the C-terminal helix α3 and the interdomain linker (Figure 4C). In addition, the NMR titration of hSAM68 (C2) by unlabeled U1A showed that the NMR signals from the tyrosine-rich sequence (370–400) of SAM68 (C2) are the most affected and thus strongly support that U1A RRM1 interacts with this aromatic rich sequence of hSAM68 in vitro at G-Y-E/D triplets.

**SAM68 recruits U1 snRNP to the 5′ splice site of mTor intron 5**

We next sought to determine if SAM68, through its association to U1A, could serve as an adaptor protein mediating the interaction between mTor pre-mRNA and the U1 snRNP. More specifically, we assessed if this interaction is mediated through the simultaneous association of SAM68 with its binding motifs (SBs) found near the 5′ splice site of
Figure 4. Tyrosine-rich (YY) domain of SAM68 mediates the interaction with U1 snRNP via YXXY repeated motif. (A) Overlay of the 2D $^{15}$N-$^1$H HSQC spectra of GB1-hSAM68 (C2) recorded before and after the addition of unlabeled GB1-U1A. The spectra are colored according to the molar ratio hSAM68 (C2):U1A (1:0; 1:0.6 and 1:1.4 are colored in blue, red and black, respectively). Strongly perturbed signals are marked by red arrows and their assignment is indicated. (B) Overlay of the 2D $^{13}$N-$^1$H HSQC spectra of U1A RRM1 recorded before and after the addition of unlabeled GB1-hSAM68 (C2). The spectra are colored according to the molar ratio U1A RRM1:hSAM68 (C2) (1:0; 1:0.6 and 1:1.4 are colored in blue, red and black, respectively). (C) Plot of the normalized chemical shift perturbations observed in panel (B) in function of the sequence of U1A RRM1. The chemical shift perturbations are then plotted onto the surface representation of the structure of the free form of the RRM1 of U1A (47). Amino acids that experienced chemical shift perturbation between 0.03 and 0.05 are colored in orange while the CSP higher than 0.05 are colored in red.

$mTor$ intron 5 and with U1A. Indeed, while SAM68 binding sequence 1 (SB-1) does not correspond to the bona fide U/AAA consensus sequences identified by SELEX (35), it was previously shown that SAM68 binding to its target sequence closest to the 5′ splice site (SB-1) was essential for $mTor$ normal splicing and expression during adipogenesis (21). Moreover, SB-1 shares a high level of homology to a specific SAM68 binding sequence identified in the β-actin mRNA (27). On the other hand, the SAM68 binding motif embedded within the cryptic poly(A) signal (SB-A) has the U/AAAA consensus sequences. Both SB-1 (UUUUAU) and SB-A (AUAAAAAU) were shown to be bound by mSAM68 in vivo (21). Interestingly, these two sequences are separated by only 11 nucleotides, which correspond to a bipartite pattern (UUUUAU-(n11)-AUAAAAAU), found to favor binding and homodimerization of GSG protein family, including SAM68 (56,57). As such, we reasoned that disrupting SAM68 binding to intron 5 by mutating both these sequences should drastically hinder U1 snRNP recruitment at the 5′ splice site of $mTor$ intron 5 (Figure 5A).

To assess this, we in vitro transcribed different RNA baits using a minimal portion of the $mTor$ minigene that
Figure 5. Both SAM68 and intronic enhancer sequences in mTor intron 5 are required for U1A recruitment to 5′ SS in vitro. (A) Schematic representation of the various in vitro transcribed mTor minigene baits with the 5′ splice site. As shown, the baits span from last 7 nucleotides of exon5 to the poly-adenylation signal in intron 5. WT refers to the wild-type intronic SAM68-binding sequences of SB-1 (UUUUAU) and SB-A (UAAAA), the latter is embedded in the cryptic poly-adenylation signal (AAUAAA). The ‘mut’ denotes the combined mutations of SB-1 (UUUUAU to UUUCAU) and SB-A (AAUAAAA to AAUAACC). (B) SAM68 recruits U1A to 5′ splice site in vitro. Recombinant in vitro purified hSAM68-Flag was tested for its ability to recruit U1A to mTor intron 5 baits with either WT or mutated SAM68-bindingsites. GST-Flag was used as negative control. (C) Schematic representation of the various in vitro transcribed mTor minigene baits that are deleted for the 5′ splice site. As shown, the baits span 18 nucleotides downstream of the 5′ splice site. WT refers to the wild-type intronic SAM68-binding sequences, SB-1 (UUUUAU) and SB-A (AAUAAA). The ‘mut’ denotes the combined mutations of SB-1 (UUUUAU to UUUCAU) and SB-A (AAUAAAA to AAUAACC). (D) SAM68 recruits U1A in the absence of 5′ splice site in vitro. Recombinant in vitro purified hSAM68-Flag was tested for its ability to recruit U1A to mTor intron 5 baits lacking 5′SSs with either WT or mutated SAM68-bindingsites. GST-Flag was used as negative control. (E) Schematic representation of the in vitro transcribed mTor minigene bait and the primers used for the RppH/Xrn1 protection assays. (F) Assessment of the processivity of RppH and Xrn1 enzyme on the naked mRNA bait, showing that RppH treatment is necessary for Xrn1-mediated degradation of the mRNA bait. (G) RppH and Xrn1 protection assays in vitro produced mRNA bait incubated with either WT MEFs cell lysate (lane 1), Sam68−/− MEFs cell lysate (lane 2), in vitro produced mSAM68(WT) + Sam68−/− MEFs cell lysate (lane 3) or in vitro produced mSAM68(WT) + Sam68−/− MEFs cell lysate + U1 snRNAs antisense oligo (lane 4). U1snRNP components (U1A, U1C) and mSAM68 levels were assessed by western blot, while U1 snRNA levels was assessed by RT-PCR. GAPDH served as loading control for the western blot. (H) SAM68 protects the mTor RNA bait from Xrn1 degradation. Biotinylated RNA baits were incubated with buffer (lane 1), 100 ng of GST-Flag (lane 2) or 100 ng of mSAM68-Flag (lane 3) for 30 min on ice. Sam68 levels were assessed by western blotting using anti-Flag, while bait levels were measured by semi-quantitative RT-PCR using FSS-RSB primers for the full-length RNA and FSB-RSB for the SAM68 protected fragment.
span from the last 7 nucleic acid of exon 5 to the cryptic polyadenylation signal at the intron 5 (Figure 5A). To determine if U1 snRNP recognition of the 5′ splice site was driven by SAM68, we mutated SAM68-binding sequence closest the 5′ splice site (SB-1), as well as the binding site embedded in the polyadenylation (SB-A) (Figure 5A). The 3′-end of the RNA baits were labeled with UTP-Biotin tails and immobilized on streptavidin-agarose beads. The baits were first incubated with in vitro purified hSAM68-Flag recombintant protein and then, shSAM68 HEK-293T cell extract was added to the mix. As observed in figure 5B, U1A was mostly detected on the baits harboring WT SAM68-binding sites, while its presence was greatly decreased on baits lacking the SAM68-binding sites. This suggests that not only SAM68 association with mTor intron 5 strengthen the 5′ splice site recognition by U1 snRNP, but also that SAM68 association could recruit U1 snRNP to the 5′ splice site of mTor intron 5 (Figure 5B). To assess this, we used RNA baits lacking the 5′ splice site (Δ5′SS) harboring either WT or mutated SB-1 and SB-A (Figure 5C). Surprisingly, we found that SAM68 was able to recruit the U1A even when the 5′ splice site was absent (Figure 5D).

While these results suggest a SAM68-dependent recruitment of U1A, it remains unclear whether U1 snRNP is only tethered to the RNA bait by a direct protein interaction with SAM68 or if there is an improved recognition by U1 snRNP to the 5′ splice site. To determine that, we incubated a mRNA bait produced in vitro, similar to the one used in the previous pulldown experiments (Figure 5E), which was incubated with different nuclear extracts and subjected to Xrn1, a 5′→3′ exonuclease (58). As shown in figure 5F, Xrn1 can efficiently degrade the RNA bait following treatment with RppH, an RNA 5′ pyrophosphohydrolase that removes pyrophosphate from the 5′ end of triphosphorylated RNA to leave a 5′ monophosphate RNA, the substrate of Xrn1 (59–61). To determine if SAM68 could recruit and enhance U1 snRNP 5′ splice site recognition, we incubated our bait with either WT MEFs cell lysate, Sam68−/−/MEFs cell lysate or Sam68−/−/MEFs cell lysate supplemented with an in vitro produced flag-tagged mSAM68 (Figure 5G). RNP complexes were pulled down using streptavidin agarose beads and treated with RppH prior to their digestion with Xrn1. Using specific primers (Figure 5E), we found that, in the presence of mSAM68 (from WT MEFS cell lysate or in vitro production), 5′ splice site of Sam68−/−/MEFs were protected from RppH and yet exposed to Xrn1, a 5′→3′ exonuclease, which was protected from RppH by SAM68. This was further confirmed with a recruiting U1 snRNA antisense oligo directed against the RNA-binding site of U1 snRNP (Figure 5G, lanes 2 and 5). In both cases, the SAM68-induced U1 snRNP protection of the 5′ splice site was significantly reduced when the cell extracts were supplemented with the U1 snRNA antisense oligo in order to impair U1 snRNA hybridization with the 5′-splice site of the mTor RNA bait.

**SAM68 recruitment of U1 snRNP is specific to endogenous mTor intron 5**

To confirm that SAM68 binds to the endogenous intron 5 sequences of mTOR, we performed cross-linking immunoprecipitation (CLIP) on WT and Sam68−/−/MEFs and assessed mSAM68 binding to different intron of mTOR preRNA. mSAM68 specifically associated in intron 5 near the exon-intron junction in WT MEFS, while no binding was detected in two other introns (4 and 37), which lack mSAM68 binding sites (Supplementary Figure S6A). Furthermore, no signal was observed in the Sam68ΔARM background. To rescue mSAM68 binding, we performed SAM68 CLIP on Sam68−/−/MEFs transfected with mSAM68(WT), mSAM68(ΔARM) or GFP as a negative control. Enrichment of the intron 5 sequence was observed in MEFS expressing either the WT or ΔARM versions of mSAM68 but was undetectable at the other tested intron (Supplementary Figure S6B).

We then sought to determine if mSAM68 could recruit U1 snRNP on endogenous mTOR, as observed with in vitro synthesized baits. To address this, we performed RNA immunoprecipitation (RIP) of U1A and assessed U1 snRNP coverage of the different exon–intron junctions in both WT and Sam68−/−/MEFs. No U1A signal was found at mTOR exon5–intron 5 junction (e15) of Sam68−/−/MEFs (Figure 6A), while U1A signal was easily detectable in WT MEFS, and in the other exon–intron junctions (e4 and e37) of both WT and Sam68−/−/MEFs (Figure 6A). These results confirm that mSAM68 directly facilitates the recruitment of U1 snRNP to e15, while not affecting other exon–intron junctions where there is no SAM68-binding site. Moreover, mSAM68 ability to recruit U1 snRNP was not shared with the mSAM68(ΔARM) mutant in a Sam68−/−/MEFs background, further suggesting that the recruitment of U1 snRNP to e15 is indeed through SAM68 (Figure 6B).

Accordingly, U1A immunoprecipitation in Sam68−/−/MEFs expressing mSAM68(WT) or mSAM68(ΔARM) revealed that only the WT protein allowed the detection of e15, and no change of U1 snRNP association was observed in other exon–intron junctions. Taken together, these results confirm that SAM68 binding to its target intronic sequences is sufficient to recruit the U1 snRNP through U1A to the 5′ splice site of mTor intron 5 and thus facilitates its recognition and the stabilization of the U1 complex at the 5′ splice site.

**SAM68 regulates mTor splicing through the recruitment of U1 snRNP**

To determine the role of SAM68–U1snRNP interaction in the regulation of mTor intron 5 splicing endogenously, we performed in vivo splicing assay on mTor in WT or Sam68−/−/MEFs, using a common forward primer and a splicing specific reverse primer (i5 versus e6) (Figure 7A). As expected, we observed a drastic decrease of mTorΔ4-6 ampiclon upon Sam68 inactivation, which was replaced
by a robust increase of the mTor5 transcript (Figure 7B), confirming our previous observation that Sam68 depletion leads to increased intron 5-induced termination (21). To rescue Sam68 depletion splicing effects, Sam68−/− MEFs were transduced with either Sam68 (WT) or Sam68 (ΔARM). Total RNA was isolated 48 h post transfection and analyzed by semi-quantitative RT-PCR. As expected, upon expression of mSAM68(WT), there was a significant decrease in intron 5-induced premature termination and polyadenylation (Figure 7B). On the other hand, cells that expressed mSAM68(ΔARM) were unable to rescue the splicing defect. This result confirms that reduced association with U1A leads to U1 snRNP recruitment impairments and thus, mTor intron 5 proper splicing. While robust level of intron 5 detection was still observed in cells expressing mSAM68(ΔARM), levels were lower than vector control (Figure 7B). This partial decrease in intron 5-induced termination is assumingly provoked by the remaining hSAM68(ΔARM)–U1A interaction, as observed in Figure 4C. As proposed in our previous work using the 3T3-L1 cell lines, this SAM68 splicing defect was also associated with decreased mTor expression level in MEFs. Indeed, mTOR protein level was highly decreased in Sam68−/− MEFs and Sam68−/− MEFs expressing mSAM68(ΔARM), when compared to WT MEFs or Sam68−/− MEFs expressing mSAM68(WT) (Supplementary Figure S7).

These results were also observed in an in vivo splicing assays using the mTor minigene, a plasmid that drives the expression of the 2.3 kb genomic fragment spanning from exon4 to exon6 of mouse mTor (Supplementary Figure S7A). Indeed, hSAM68-depleted HEK-293T cells co-transfected with the minigene and either Flag-YFP, Flag-hSAM68(WT) or Flag-hSAM68(ΔARM) showed similar results that Flag-hSAM68(WT) was able to revert the increased mTor5/mTor tot ratio, while the Flag-hSAM68(ΔARM) behaved like Flag-YFP (Supplementary Figure S6B).

**DISCUSSION**

The KH domain RNA-binding protein, SAM68, regulates splicing of mTor as well as the ribosomal S6 kinase (Rps6kb1) transcripts in pre-adipocytes (21,39). In turn, pre-adipocytes of Sam68−/− mice do not differentiate to adipocytes due to defective mTOR signaling. Our data show that SAM68 modulates mTor splicing by binding to specific regulatory elements found in intron 5 (SB-1 and SB-A), of which SB-A overlaps with the poly-adenylation signal (AAUAAA). This led us to postulate that these AU-rich cis-acting elements were intronic splicing enhancers to which SAM68 bound with great affinity to modulate the recruitment of U1 snRNP at the upstream 5′ splice site.

In this study, we report that SAM68 functionally interacts with U1 snRNP, the splicing component that recognizes 5′ splice sites (10). While SAM68 can bind with low affinity to the U1 snRNA in vitro, it is highly unlikely that it will happen in vivo. Furthermore, the potential AU-rich SAM68 binding motif found in the U1 snRNA is located in the Sm site, which is rapidly masked by the Sm protein ring during U1 snRNP assembly (49,62). Rather, we found that SAM68 interacts directly with U1A, the stem-
Figure 7. SAM68 deletion of ‘ARM binding region’ shows decrease in U1A binding. (A) Schematic of the pcDNA mTor<sub>4-6</sub> minigene, comprising the mTor genomic fragment from exon4 to exon6. (B) (Left panel) Sam68<sup>−/−</sup> MEFs were infected with Sam68(WT) or Sam68(DARM) and compared to uninfected Sam68<sup>−/−</sup> MEFs or WT MEFs. Total RNA was extracted in each cell lines and semi-quantitative RT-PCRs were performed using endogenous mTor specific primers. Forward(Fe4) and Reverse(Re6) were used to quantify mTor transcripts that were spliced normally (mTor<sub>4-6</sub>), while Forward(Fe4) and Reverse(Ri5) were used to quantify intron 5 including mTor transcripts (mTor<sub>i5</sub>). Gapdh was used to normalize the values obtained. Total protein was also extracted and run on 10% SDS-PAGE and blotted with SAM68, U1A and GAPDH antibodies. (Right panel) Quantification of intron 5-induced termination over normally spliced mRNA based on three independent experiments using endogenous mTor specific primers. **P-value ≤ 0.05 and # = non-significant (two-tailed t-test).

loop II binding protein of U1 snRNA. This SAM68–U1A interaction was shown to be resistant to RNaseA treatment and RNA-binding defective mutant versions of SAM68 bearing point mutations in the KH domain; SAM68<sub>V229F</sub> and SAM68<sub>184N</sub> (36,52) could still bind U1A and the U1snRNP. Domain mapping studies showed that U1A binds to the ‘minimal ARM binding region of SAM68 and that deleting this region (379–389 aa) in SAM68(WT) was sufficient to impair U1A association. This region located within SAM68 YY domain was initially identified to regulate T-cell factor 1 splicing by binding the armadillo repeat (ARM) domain of Adenomatous polyposis coli (APC) (54,55). Hence, our results confirmed that SAM68 can directly interact with U1A, making it the only identified protein interactor of this U1 snRNP core protein to date. NMR spectroscopy also confirmed that this protein–protein interaction is mediated through the tyrosine-rich ‘ARM-binding domain’ of SAM68 and the RRM1 domain of U1A. Conversely, we observed that deleting the ARM domain of SAM68 greatly impaired its association with U1A, resulting in increased intron 5 inclusion. Residual splicing activity could still be observed, which might be due to the remaining YXXY motifs in SAM68(DARM) that may act as weak surrogate binding sites in the absence of the ARM motif. Results obtained with synthetized RNA baits suggest that mSAM68 initiate U1 snRNP recruitment, which then allows the recognition of the sub-optimal splice site. Hence, the presence of mSAM68 increases the recognition rate of the 5’ splice site of mTor intron 5, by increasing U1 snRNP stoichiometry close to the splice site. This was further confirmed by the endogenous mSAM68 CLIP, the U1A RNA immunoprecipitation assays and the mTor splicing disparity observed using either the mTor minigene<sub>exon4-6</sub> or the endogenous mTor. These findings highlight the importance of intronic enhancer mediated binding of SAM68, subsequent recruitment of U1snRNP via U1A and the resultant
splicing of mTor intron 5. This is strikingly similar to the mechanism by which other RNA-binding proteins, such as TIA1 and RBM24, modulate 5′ splice site recognition of Drosophila male specific lethal (msl) gene and inhibitor of κ light polypeptide gene enhancer in B cells, kinase complex-associated protein (IKBΔKAP) gene in familial dysautonomia (FD) by interacting with U1snRNP (17,63). Indeed, both of these RNA-binding proteins were shown to promote splicing of pre-mRNAs containing sub-optimal splicing signals by interacting with components of U1 snRNP. While TIA1 interacts with U1C and SAM68 with U1A, both act as intronic splicing enhancers by binding sequences close to the 5′ splice sites, thus facilitating U1 snRNP recruitment.

As shown here, SAM68 seems to initiate spliceosome assembly at the 5′ splice site of mTor intron 5, through the recruitment the U1 snRNP. This is mainly achieved when SAM68 binds specific sequence within the 5′ portion of mTor intron 5, mainly the bipartite sequences designated as SB-I and SB-A. While these two sequences are clearly involved in U1 snRNP recruitment, the role of the downstream SAM68-binding sites found throughout intron 5 remains to be defined. One main hypothesis could be that SAM68 regulates early steps of spliceosome assembly (e.g. complex E formation) in specific introns. This seems to be in line with the observation that SAM68 binds U2AF65, a subunit of the U2AF complex known to initiate 3′ splice site recognition (64). Moreover SAM68 binding to U2AF65 increased its efficiency. As SAM68 homodimerizes through its association with RNA, it is also possible that multiple binding sites favor its dimerization, thus decreasing the gap between the 5′ and the 3′ splice site, which would again facilitate early spliceosome assembly.

It has been recently reported that introns with weak 5′ splice sites are susceptible to premature cleavage and polyadenylation events in the absence of U1 snRNP binding and RNA-binding proteins are required for U1 snRNP occupancy at such sites (65). Indeed, morpholino oligonucleotides interfering with U1 snRNA binding causes premature cleavage and polyadenylation in numerous pre-mRNAs at cryptic polyA sites in introns and this occurs near the start of the transcripts (66). This mechanism seems to involve core protein components of the U1 snRNP, since both U1–70K and U1A were reported to inhibit polyadenylation by mediating a direct interaction with poly-A-polymerase (67,68). This U1-mediated inhibition of polyadenylation even led to the characterization of a Polyadenylation inhibitory Elements (69). Concomitantly, SAM68 has also recently been reported to regulate 3′ end processing of Aldh1a3 pre-mRNA, a mechanism that is required to maintain glycolytic metabolism and self-renewal of mouse neural progenitor cells (37). While our results show that mSAM68 recruits U1 snRNP to the 5′ splice site of mTor intron 5 and increases the rate of intron 5 excision, it also seems to cooperate in masking crytic polyadenylation sites, a process that is known to be mediated by U1 snRNP (66,70). Indeed, SAM68 alone (without U1 snRNP) is unable to modulate alternative intronic polyadenylation usage, since SAM68 (ΔARM) can bind the poly(A) signal as efficiently as the WT version, but still allows polyadenylation of the mTor15 transcript. Hence, these results suggest that 5′ splice site recognition by the SAM68–U1 snRNP complex precludes intronic cryptic polyadenylation usage, while allowing proper excision of mTor intron 5. This also suggest that SAM68 could also be involved in the modulation of the previously described U1-dependent modulation of alternative poly(A) usage (37,66,70,71).

Our findings illustrate the first mechanistic evidence showing that SAM68 recruits U1 snRNP via direct interaction with U1A to upstream 5′ splice site, while it binds AU-rich regulatory sequences in mTor intron 5, through its RNA-binding KH domain. This is subsequently mediated through its tyrosine-rich ‘ARM-binding domain’, and in turn enhances splicing of intron 5, while SAM68 depletion inhibits U1 snRNP recruitment and promotes premature intron 5-induced termination and polyadenylation. Although the regulatory mechanism of this interaction remains elusive, the presence of numerous tyrosine residues located within the ‘YY region’ of SAM68 suggest that phosphorylation might be involved. Indeed, SAM68 tyrosine phosphorylation was shown to negatively affect its ability to bind RNA (31,33,34), and its splicing functions (i.e. Bel-X, CD-44 and Cyclin D1) (22,23,33).

In the future, the global identification of mRNA targets regulated by this SAM68-dependent alternative splicing should allow us to better understand these possible regulatory mechanisms and determine how a binding site situated 174-nt downstream of the 5′ splice site can modulate its recognition by the U1 snRNP. Indeed, it would be interesting to assess if other alternatively spliced mRNAs targeted by SAM68 could be differentially regulated through distance changes between SAM68-binding sites and the 5′ splice site during cellular processes such as development or cancer. Moreover, determining how the SAM68–U1A interaction is modulated by distance constraints, secondary structures or associated proteins to regulate not only mTor, but also a specific sub-class of mRNA, remains to be investigated.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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