Shwachman–Bodian–Diamond syndrome (SBDS) protein deficiency impairs translation re-initiation from C/EBPα and C/EBPβ mRNAs

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ABSTRACT

Mutations in the Shwachman–Bodian–Diamond Syndrome (SBDS) gene cause Shwachman–Diamond Syndrome (SDS), a rare congenital disease characterized by bone marrow failure with neutropenia, exocrine pancreatic dysfunction and skeletal abnormalities. The SBDS protein is important for ribosome maturation and therefore SDS belongs to the ribosomopathies. It is unknown, however, if loss of SBDS functionality affects the translation of specific mRNAs and whether this could play a role in the development of the clinical features of SDS. Here, we report that translation of the C/EBPα and -β mRNAs, that are indispensable regulators of granulocytic differentiation, is altered by SBDS mutations or knockdown. We show that SBDS function is specifically required for efficient translation re-initiation into the protein isoforms C/EBPα-p30 and C/EBPβ-LIP, which is controlled by a single cis-regulatory upstream open reading frame (uORF) in the 5′ untranslated regions (5′ UTRs) of both mRNAs. Furthermore, we show that as a consequence of the C/EBPα and -β deregulation the expression of MYC is decreased with associated reduction in proliferation, suggesting that failure of progenitor proliferation contributes to the haematological phenotype of SDS. Therefore, our study provides the first indication that disturbance of specific translation by loss of SBDS function may contribute to the development of the SDS phenotype.

INTRODUCTION

The autosomal recessive disorder Shwachman–Diamond syndrome (SDS) is caused by the expression of hypomorphic alleles carrying mutations in the Shwachman–Bodian–Diamond syndrome (SBDS) gene (1). SDS is characterized by bone marrow failure with neutropenia, exocrine pancreatic insufficiency and skeletal abnormalities (2). In mice, complete loss of SBDS function is embryonic lethal (3), indicating that SBDS is an essential gene. Over the past decade, diverse functions for SBDS have been described, including mitotic spindle stabilization (4), chemotaxis (5), Fas ligand-induced apoptosis (6), cellular stress response (7) and Rac2-mediated monocyte migration (8). Nonetheless, there is now compelling evidence that SBDS functions in cytoplasmic ribosome maturation (9–13). Thus, SDS should be considered a ribosomopathy caused by defective maturation of the large ribosomal subunit. Studies with eukaryotic SBDS and its yeast ortholog Sdo1 showed that SBDS cooperates with the GTPase elongation factor-like 1 (EFL1) to catalyse removal of the eukaryotic initiation factor 6 (eIF6) from the 60S ribosome subunit. eIF6 is critical for biogenesis and nuclear export of pre-60S subunits and prevents ribosomal subunit association. Therefore, its release is required for ribosomal subunit association during translation initiation (9,10,13–15). Currently, it is not known whether SBDS deficiency mainly causes a general effect on mRNA translation, or whether it results in aberrant translation of specific mRNAs that contributes to the SDS phenotype.

Neutropenia is the most prominent haematopoietic abnormality seen in almost all SDS patients (16). Myeloid progenitors derived from the bone marrow of SDS patients have a reduced proliferation capacity with low frequency of CD34+ cells and reduced colony forming ability (17).

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The CCAAT enhancer binding proteins C/EBPα and C/EBPβ are critical transcription factors for myelomonocytic lineage commitment, granulocyte differentiation and macrophage function (18–20). Expression of C/EBPα and -β proteins are strictly controlled at the mRNA-translation initiation level (21–23). From consecutive initiation codons in the C/EBPα mRNA three different protein isoforms are synthesised. Extended-C/EBPα or full-length C/EBPα-p42 is expressed from a cap-proximal GUG- (CUG for rodents) or AUG-codon, respectively. A shorter N-terminally truncated C/EBPα-p30 isoform is translated from a distal AUG-codon. Translation from the distal AUG into C/EBPα-p30 requires re-association of ribosomes following translation of a cis-regulatory upstream open reading frame (uORF) in the 5′ untranslated region (5′ UTRs) of the C/EBPα mRNA (Figure 1A) (22). Extended-C/EBPα is not further considered here since its expression from the non-canonical GUG codon is usually very low.

C/EBPα-p42 expression and induction of target genes such as the CSF3R (colony stimulating factor 3 receptor (granulocyte)) is essential for granulocytic differentiation (24). In addition, C/EBPα-p42 inhibits MYC expression, which causes proliferating myeloid precursor cells to undergo cell cycle arrest and entry into terminal differentiation (25). C/EBPα-p30 lacks the major part of the N-terminal transactivation sequences but retains the C-terminal DNA-binding domain and therefore competes with C/EBPα-p42 or other C/EBPs for DNA binding (20). Moreover, solitary expression of C/EBPα-p30 blocks granulocytic differentiation and results in an Acute Myeloid Leukemia (AML) like disease in mice (26,27). Translation of the C/EBPβ mRNA into different protein isoforms is regulated in a similar way (22,28). C/EBPβ-LAP* and -LAP proteins (LAP: liver activating protein) are translated from cap-proximal AUG-codons, while the truncated C/EBPβ-LIP (LIP: liver inhibitory protein) protein is translated from a distal AUG-codon through an uORF-dependent mechanism. LAP* expression is often very low since the LAP*-AUG codon lacks a Kozak-consensus sequence that is required for efficient translation initiation (29).

Translation of the C/EBPα and -β mRNAs is highly sensitive to alterations in the translation machinery. Since both factors play such a crucial role in the development of the myelomonocytic lineage, we examined whether C/EBPα and -β translation is regulated by SBDS and whether C/EBPα and -β isoform expression is altered by SBDS mutations. Here, we show that SBDS is required for efficient translation of the truncated p30/LIP isoforms and that lymphoblastoid cell lines derived from SBDS patients have reduced C/EBPβ-LIP levels. In addition, our study suggests that SBDS deficiency indirectly suppresses the expression of MYC by increasing the C/EBPα-p42/p30 isoform ratio. Furthermore, our study suggests that this regulatory connection between SBDS, C/EBPα/β and MYC is decisive for myeloid cell proliferation and differentiation. Our finding is the first evidence of specific defective mRNA translation in SDS, highlights the importance of the uORF-mediated translation re-initiation mechanism that directs C/EBPα and -β isoform expression for hematopoietic proliferation and differentiation control.

**MATERIALS AND METHODS**

**Cell culture**

C33A cells were maintained in DMEM (Gibco) supplemented with 10% fetal calf serum (FCS) and 1% Penicillin/Streptomycin. HL-60 and HL-60 pCMV C/EBPα-p30flag overexpressing cells were commonly maintained in RPMI (Gibco) supplemented with 10% FCS and 1% Penicillin/Streptomycin. SDS patient-derived lymphoblast cell lines including healthy wild type cell were maintained in RPMI (Gibco) with 15% FCS and 1% Penicillin/Streptomycin at 37°C with 5% CO2. For neutrophil differentiation, 5.0 × 105 cells/ml were treated with 5 μM ATRA (Sigma) for 7–10 days. After treatment of all-trans retinoic acid (ATRA), cells were harvested and stained with Giemsasolution for morphologic differentiation determination. The HL-60 cells expressing C/EBPα-p30flag were generated by retroviral transduction with a pMSCV-neo based construct and subsequent selection of the transduced cells with G418 (1.8 mg/ml). For knockdown of SBDS in HL-60 and C33A cells, the cells were transduced with the lentiviral plKO.1 shRNA vector against human SBDS (Sigma Aldrich) or plKO.1-scrambled shRNA as control followed by selection of the transduced cells with puromycin (1.5 μg/ml). For ectopic SBDS expression HL-60 or C33A cells were transduced with a lentiviral pLVX-ires-Neo vector and selected with G418 (1.8 mg/ml for HL60 and 0.8 mg/ml for C33A). Cell number and viability was determined using the CASY electric field multi-channel cell counting system following the manufacturer’s instruction.

**Plasmid cloning and transfection**

pLKO.1 shRNA vector against human SBDS was purchased from Sigma Aldrich. shSBDS sequence: 5′-CCG AGA AAT TGA TGA GCT AAT ctc gag ATT AGC TCA TCA ATC ATT CTT CGG-3′. For construction of the SBDS overexpression vector, total mRNA was isolated from human lymphoblast cells using the RNeasy kit (Qiagen) and SBDS cDNA was synthesized using the Transcriptor First-strand cDNA Synthesis kit (Roche), which was used as a PCR template. Forward primer: 5′-GTG AGA AAT TCA TGT CGA TCT TCA CCC CCA C-3′, Reverse primer: 5′-TCT GAA TCA TAC AAA TTT CTC ATC TTC TT-3′. The PCR product was cloned into the pLVX-ires-Neo plasmid (Clontech) for lentiviral transduction and into the pCDNA3 vector (Invitrogen) for transient transfection using the EcoRI and XbaI restriction sites. Wild-type rat C/ebpα and -β cDNAs and uORF start codon mutant constructs (rat C/ebpα : ATG> ATC, rat C/ebpβ : ATG> ATT) were previously cloned (22). C33A stable SBDS knockdown or SBDS overexpression cells were transiently transfected with wild-type or mutant rat C/EBPα and -β expression plasmids using FugeneHD (Roche) following the manufacturer’s instructions.

**Lentiviral and retroviral transduction**

A total of 3.5–3.8 × 10⁶ Hek293T cells were used for co-transfection with pLKO.1 shSBDS vector (from Sigma
Figure 1. Deregulated C/EBPβ protein isoform expression in SDS. (A) The human C/EBPα and -β mRNAs are presented with consecutive translation initiation sites (arrowheads) and each of the protein isoforms and its size (*size of murine orthologs). Extended, p42, LAP* or LIP proteins are expressed through regular translation initiation, omitting the uORF. Truncated p30 or LIP proteins are expressed through translation re-initiation by post-translation ribosomes that have first translated the uORF. For detailed description of the uORFs and surrounding sequences, see (21–23). Expression of the Extended-C/EBPα isoform is generally weak because it uses the alternative GUG (CUG for murine) codon. Similarly, expression of the C/EBPβ-LAP* from a non-Kozak AUG codon is mostly weak. (B) SBDS protein levels were detected in SDS patient-derived (SW18, SW74) and healthy control-derived (wt) lymphoblastoid cells by immunoblotting. Long exposure shows the very low expression of wt SBDS in SW74 cells harbouring the homozygous 258 +2T>C mutation. (C) The upper panels show immunoblots of C/EBPβ isoforms, SBDS and α-tubulin as loading control in both SDS patient-derived cells (SW18, SW74) and healthy control-derived cells (wt). The lower panels show immunoblots of 4E-BP1, phosphorylated-4E-BP1 (P-4E-BP1), S6K1, phosphorylated-S6K1 (P-S6K1) and β-actin as loading control to monitor alterations in mTORC1 signalling (D) qRT-PCR analysis for endogenous C/EBPβ-mRNA levels in patient-derived cells (SW18, SW74) and healthy control cells (wt). (E) Immunoblots for MYC, SBDS and α-tubulin (loading control) in SDS patient-derived and healthy control-derived cells. (F) qRT-PCR analysis of MYC transcript levels in SDS patient-derived and healthy control-derived cells. (G) Cell multiplication assay with patient-derived cells (SW18, SW74) and healthy control-derived cells (wt). Seven days after seeding 5 x 10^5 cells, cells were harvested and counted. Statistical differences were determined by Student’s t-tests. Error bars represent ±SD (n = 3), *P < 0.05, **P < 0.01.
Aldrich) or pLVX SBDS expression vector (10 μg) together with plasmid for viral packaging factors including pMDL-RRE (6.5 μg), pCMV-VSVg (3.5 μg) and pRSV-Rev (2.5 μg) were transfected using the calcium phosphate method. For retroviral transduction Phoenix A packaging cells were transfected with the C/EBPα-p30flag pMSCV-neo vector with the calcium phosphate method. After 48 h, lentiviral-or retrovirus-containing medium was harvested from the transfected cells. Supernatant was collected by centrifugation and filtered using a 0.45 μm size PVDF filter. A total of 5.0 × 10⁶ cells were incubated with virus-containing supernatant with 4 μg/ml (HL-60) or 8 μg/ml (C33A) polybrene for 24 h. To establish stable cell lines, puromycin or G418 was added to cells in fresh medium.

**Immunoblotting**

Cells were washed with cold phosphate buffered saline (PBS) and harvested by scraping or centrifugation. Cells were incubated with lysis buffer (50 mM Tris pH 7.5, 150 mM NaCl, 1mM EDTA pH 8.0, 1% TritonX-100 with Protease- and Phosphatase Inhibitors (Roche)) for 30 min at 4°C or sonicated with a Bioruptor for 5 min at 4°C. After centrifugation with 13 000 rpm for 30 min at 4°C, the supernatant was obtained and used for immunoblot analysis. The protein concentration measurement was done with Bradford assay (BioRad). Equal amount of protein extracts were loaded to 10 or 12% SDS-PAGE and transferred to PVDF membrane (Biorad) by semi-dry blotting for 1 h. Membranes were washed with washing buffer (Tris buffered saline with 0.1% tween 20) and blocked for 1 h at room temperature with blocking buffer (5% skim milk solution). The primary antibodies, SBDS (S-15), C/EBPα C18 (sc-9314) for human C/EBPα and C/EBPα 14AA (SC-61) for rat C/EBPα, C/EBPβ C19 (SC-150) for both human and rat C/EBPβ, MYC (sc-40), α-Tubulin (sc-8035), 4E-BP1 (C-19) (all Santa Cruz Biotechnology) and Phosphorylated 4E-BP1 (Thr37/46) (9459), Phosphorylated p70-S6K (Thr389) (9234), p70-S6K (9202) (Cell Signaling) and β-actin (691001) (MP Biomedicals) were diluted in blocking buffer (1:400–1:1000). Incubation with primary antibody was carried out at 4°C overnight using an orbital shaker. After washing with washing buffer for 10 min x 3 times, secondary antibody (1:5000) was added for 1 h at RT. Protein signals were visualised using X-ray films (Amersham) with chemiluminescence detection (Perkin Elmer). Signals were visualised using X-ray films (Amersham) with chemiluminescence detection (Perkin Elmer).

C/EBPα and-β isoform ratio measurement

Immunoblotting was performed to visualise C/EBPα and -β isoform expression as described above. To analyse isoform ratio, band intensity in immunoblot images was measured and quantified by densitometric analysis using AlphaImager software (Cell Bioscience) following the manufacturer’s instruction.

**Luciferase assay**

pcDNA3-based p42-Renilla and p30-Firefly (C/EBPα) reporters were generated based on the previously described pGL3-FL-Cα and pGL3-Tr-Cα (30): the Firefly luciferase gene was exchanged by the Renilla luciferase gene in the pGL3-FL-Cα and both reporters were re-cloned from pGL3 into pcDNA3. For the pcDNA3-based LAP-Renilla and LIP-Firefly (C/EBPβ) reporters, C/EBPβ-5’ UTR until LAP initiation codon was cloned together with Renilla (from pGL3) in pcDNA3 and C/EBPβ-sequences spanning the 5’ UTR and sequences until the LIP initiation codon with a +1 frame shift (7 nt upstream of the AUG) was cloned together with Firefly in pcDNA3. The control reporters were generated by cloning Renilla (from pGL3) or Firefly (from pGL4) genes in pcDNA3. Details of the cloning strategy will be provided upon request. For the Luciferase assays 2.5 × 10⁵/well of C33A cells were seeded into 96 well plates. After 24 h, cells were transfected with p42-, p30-, LAP or LIP reporters using FugeneHD (Roche). For restoration of SBDS function a pcDNA3-based SBDS construct or empty vector as control was co-transfected. Renilla or Firefly luciferase vectors were co-transfected for normalization of the reporter expression. After additional 48 h, cells were harvested and the Stop and Glow kit (Promega) was used to measure luciferase activity.

**Quantitative real time PCR (qRT-PCR) of mRNA**

To detect mRNA transcription, total RNA was isolated using the RNAeasy kit (Qiagen). cDNA synthesis was performed with oligo dT primers using the Transcriptor First Strand cDNA synthesis kit (Roche). GAPDH house keeping gene was used for normalization of gene expression. C/EBPα: Forward primer: 5′-GGT TTT GCC CTT CTT GGG AAT GGT GTT CAC-3′, Reverse primer 5′-CAG CTC ATT GGT CCC CCA G-3′; C/EBPβ: Forward primer: 5′-TTT CGA AGT TGG ATG CAA TCG-3′, Reverse primer: 5′-CAG CAC AAT GGG AGG AAC AT-3′; MYC: Forward primer: 5′-TCA AGA GGT GCC ACC ACG TCT CC-3′; Reverse primer: 5′-TCT TGG CAG GAT AGT CCT-3′; GAPDH: Forward primer: 5′-GTC AGT GGA CCT GAC CT-3′, Reverse primer: 5′-ACCTGAG TGC TCA GTC TAG CC-3′; SBDS: Forward primer: 5′-ACG TGC TCA CAT GAG TCT CG-3′, Reverse primer: 5′-CAG CCC GGG TCA ATC AGA CAT-3′.

**RESULTS**

C/EBPβ-LIP expression is diminished in SDS-derived lymphoblast cells

To investigate whether SBDS mutations may affect C/EBP-mRNA translation, we examined C/EBPβ isoform expression in two SDS patient-derived lymphoblast cell lines compared to lymphoblasts derived from a healthy individual by immunoblotting. The SW74 cell line carries the homozygous SBDS mutation 258 + 2T > C, a common mutation in SDS resulting in very low expression of wild type SBDS compared to the healthy control (wt) (Figure 1B); the SW18 cell line carries the heterozygous SBDS mutations 258 + 2T > C and 505C > T resulting in reduced levels of mutant SBDS (Figure 1B) (13). The C/EBPβ isoforms LAP and LIP were both expressed in the control cells (wt). By contrast, LIP expression was hardly detectable in the patient-derived cells (SW18 and SW74) while LAP levels were similar to the control (Figure 1C). We have shown...
before that expression of LIP is activated by mTORC1 signalling and that inhibition of mTORC1 results in suppression of LIP both in cell culture and in vivo (22,28,31). To examine whether SBDS deficiency reduces mTORC1 activity and thereby regulates LIP expression, we analysed the expression and the phosphorylation status of two mTORC1 targets, the 4E-binding protein 1 (4E-BP1) and the p70 ribosomal protein S6 kinase 1 (S6K1). SBDS deficiency does not significantly diminish the phosphorylation status of 4E-BP1 or S6K1 (Figure 1C, lower panels). Thus, mTORC1 signalling and its major downstream translation control pathways through 4E-BP1 or S6K1 p70 seem not to be involved in the SBDS-dependent regulation of LIP expression. Quantitative Real-Time PCR (qRT-PCR) analysis showed that SBDS-deficient cells secrete more LIP protein than their wild-type counterparts (Figure 1C, lower panels). No significant change was observed in the phosphorylation status of 4E-BP1 or S6K1, suggesting that mTORC1 signalling is not involved in the regulation of LIP expression. Hence, two different pathological SBDS mutations correlate with reduced C/EBPβ-LIP and MYC expression and reduced proliferation.

uORF-dependent regulation of C/EBPα-p30 and C/EBPβ-LIP expression by SBDS levels

Lymphoid cells do not express C/EBPα. To further examine the relation between SBDS expression and C/EBPα and -β expression, we performed SBDS knockdown in the human promyelocytic leukaemia cell line HL-60 that express endogenous C/EBPα and C/EBPβ. SBDS knockdown by lentiviral SBDS-shRNA resulted in reduced expression of C/EBPα-p30 and C/EBPβ-LIP (Figure 2A and B) and concomitant lower p30/p42 and LIP/LAP ratios. Similar as described for the patient-derived lymphoblasts SBDS knockdown did not reduce phosphorylation of 4E-BP1 or S6K1, suggesting that mTORC1 signalling is not involved in the downregulation of p30/LIP upon SBDS knockdown (Figure 2A, lower panels). No significant change was observed in C/EBPα and C/EBPβ mRNA levels. Overexpression of SBDS in HL-60 cells led to an increased expression of p30 and LIP resulting in higher LIP/LAP ratios without changing C/EBPα or C/EBPβ mRNA levels (Supplementary Figure S1A and B). These data indicate that SBDS and expression of p30 or LIP might be functionally correlated.

To examine the relevance of uORF-controlled translation of the C/EBPα and -β mRNAs by SBDS levels we transiently expressed wild-type cDNA constructs of rat C/EBPα and -β as well as constructs thereof carrying mutations that interfere with uORF-function in C33A human cervix carcinoma cells that expressed either SBDS-shRNA for knockdown or SBDS-cDNA for overexpression. We used the C33A cell line for these studies because these cells tolerate high expression of p42 without inducing the usual p42-associated cell cycle arrest that would complicate analysis (33). Similar to what we observed for endogenous C/EBPs, knockdown of SBDS resulted in a reduction of p30/p42 and LIP/LAP ratios through downregulation of p30 or LIP, respectively (Figure 2C and D), while SBDS overexpression upregulated p30 and LIP expression (Supplementary Figure S1C and D). Also in this case, SBDS knockdown did not alter mTORC1 signalling (Figure 2C, lower panels).

We have shown before that the conserved uORF in the vertebrate C/EBPα and -β mRNAs is required for translation re-initiation from the p30 or LIP translation-initiation codon, respectively (21,22). Mutation of the uORF-AUG into a non-AUG codon (ATC for C/ebpα; ATT for C/ebpβ) that eliminates uORF-function resulted in complete loss of p30 or LIP expression independent of SBDS knockdown or overexpression but did not affect p42 or LAP expression (Figure 2E and F). These data indicate that reduction of p30/LIP by SBDS levels depends on the cis-regulatory function of the uORFs in the C/EBP mRNAs as was shown before for other translational control pathways (22,28).

**Loss of SBDS function specifically affects translation re-initiation**

To examine the effect of SBDS function on translation initiation versus uORF-mediated re-initiation we used a reporter system based on the rat C/EBPα-mRNA or C/EBPβ-mRNA structures (30,34,35). Briefly, from the p42- or LAP-reporter, expression of Renilla luciferase can only be achieved by translation initiation and omission of uORF-translation, reminiscent to C/EBPα-p30 or C/EBPβ-LIP translation (Figure 3A). From the p30- or LIP-reporter, expression of Firefly luciferase can only be achieved through a translation re-initiation mechanism by ribosomes that have first translated the uORF, resumed scanning and initiated at the downstream AUG codons, which is reminiscent of C/EBPα-p30 or C/EBPβ-LIP translation. In the latter case, eventual translation from the C/EBPα-p42 or C/EBPβ-LIP AUG-codons do not run into the Firefly luciferase coding frames and therefore are not detected since a reading frame shift is introduced just upstream of the p30/LIP initiation sites (Figure 3A) (22). To normalize against effects not related to initiation/re-initiation efficiency, Renilla and Firefly luciferase expression control vectors were co-transfected along with the Renilla and Firefly reporters, respectively.

C33A cell lines expressing either SBDS-shRNA or SBDS-cDNA were transiently transfected with the reporter and control constructs and used for the luciferase assay. In the SBDS knockdown cells, translation initiation into p42-Renilla was not altered while translation re-initiation into p30-Firefly is strongly reduced, resulting in significant lower p30-Firefly/p42-Renilla ratio (Figure 3B, left side). Similarly, in the SBDS knockdown cells initiation into LAP-Renilla was slightly enhanced while re-initiation was strongly reduced, resulting in significant lower LIP-Firefly/LAP-Renilla ratio (Figure 3B, right side). Moreover, the negative effects of SBDS knockdown on p30-Firefly and LIP-Firefly expression could be rescued by additional overexpression of SBDS resulting in normalised p30-Firefly/p42-Renilla and LIP-Firefly/LAP-Renilla ratios comparable to the control (Figure 3C). Contrary to the SBDS knockdown, overexpression of SBDS did not greatly
Figure 2. SBDS is required for efficient C/EBPα-p30 and C/EBPβ-LIP expression. Expression of the endogenous (A) human C/EBPα (p42 and p30) or (B) human C/EBPβ proteins (43 kDa LAP and 20 kDa LIP) in HL-60 cells with stable SBDS knockdown (shSBDS) or control (scrambled shRNA) by immunoblotting, and quantification of p30/p42 or LIP/LAP ratio and qRT-PCR analysis of transcript levels at the right. The lower panels in (A) show immunoblots of 4E-BP1, phosphorylated-4E-BP1 (P-4E-BP1), S6K1, phosphorylated-S6K1 (P-S6K1) and β-actin as loading control to monitor alterations in mTORC1 signalling. Immunoblots of (C) r a t C/EBPα (p42 and p30) or (D) r a t C/EBPβ (38 kDa LAP*, 34 kDa LAP and 20 kDa LAP) expression in C33A cells with stable SBDS knockdown (shSBDS) and control (scrambled shRNA) transiently transfected with wild-type rat C/ebpα or wild-type rat C/ebpβ expression plasmids, respectively. Quantification of isoform ratio is shown at the top. The lower panels show immunoblots of 4E-BP1, phosphorylated-4E-BP1 (P-4E-BP1), S6K1, phosphorylated-S6K1 (P-S6K1) and β-actin as loading control to monitor alterations in mTORC1 signalling. (E) Immunoblot of rat C/EBPα or (F) rat C/EBPβ proteins in C33A cells transiently transfected with mutated uORF-AUG (dysfunctional uORF) rat C/ebpα or -β cDNA expression vectors. Statistical differences in the mRNA levels and isoform ratios were determined by Student’s t-tests. Error bars represent ±SD (n = 3), *P < 0.05, **P < 0.01, ***P < 0.005.
Figure 3. Loss of SBDS function specifically affects translation re-initiation. (A) Representation of the p42- or LAP-Renilla luciferase and p30- or LIP-Firefly luciferase reporters. Renilla luciferase expression from the p42/LAP-reporters results from translation-initiation omitting the uORF, while Firefly luciferase expression from the p30/LIP-reporters is achieved through uORF-mediated translation re-initiation (30). (B) C33A cells with stable SBDS knockdown (shSBDS) or control cells (scramble shSBDS) were transiently transfected with the p42-Renilla or p30-Firefly (graphs at the left) or LAP-Renilla or LIP-Firefly (graphs at the right) reporters, and co-transfected with respective complementary Renilla or Firefly luciferase control expression vectors. The bar graphs show luciferase values of the p42- and p30-reporters and the p30-Firefly/p42-Renilla ratio, and of the LAP- and LIP-reporters and the LIP-Firefly/LAP-Renilla ratio. Immunoblots of SBDS and β-actin loading control are shown at the far right. (C) C33A cells with stable SBDS knockdown (shSBDS) were either cotransfected with empty vector control (EV) or SBDS expression vector and subjected to transient transfection with reporter constructs as described in (B). (D) C33A cells were stably transfected with SBDS expression vector or EV and subjected to transient transfection with reporter constructs as described in (B). Statistical differences were analysed by Student’s t-tests. Error bars represent ±SD (n = 4), * P < 0.05, *** P < 0.005.
affect the levels of the individual reporters, resulting in similar p30-Firefly/p42-Renilla or LIP-Firefly/LAP-Renilla ratios (Figure 3D).

Therefore, collectively our experiments with C/EBPα and -β cDNAs and the related reporter systems suggest that SBDS levels affect uORF-mediated translation re-initiation but have no significant effect on regular translation initiation.

SBDS knockdown reduces MYC expression through C/EBPα p30/p42 ratio

MYC is required for the proper balance between hematopoietic stem cell self-renewal and differentiation (36). Conditional elimination of MYC in the bone marrow (BM) of mice results in severe cytopenia and accumulation of HSCs (37). Deregulated activation of MYC has been found in many types of human lymphoma and leukemia (38). MYC mRNA expression is regulated by p42 through the inhibitory interaction with E2F at the MYC promoter while p30 fails to do so (25,39,40). For example, treatment of HL-60 cells with the growth arrest and apoptosis inducing drug CDDO (2-Cyano-3,12-dioxooleana-1,9-dien-28-oic acid) causes a translational upregulation of C/EBPα accompanied by a translational downregulation of p30 and an increase in the p42/p30 ratio, which results in strong reduction of MYC mRNA levels associated with inhibition of proliferation and induction of neutrophil differentiation (41). In HL-60 cells with stable SBDS knockdown and concomitant reduction of the C/EBPα p30/p42 and C/EBPβ LIP/LAP ratios (Figure 2A and B) the expression of MYC protein and mRNA was decreased (Figure 4A and B). To examine whether down-regulation of p30 is required for MYC suppression, we overexpressed a flag-tagged C/EBPα-p30 (p30flag) protein in HL-60 cells. Overexpression of p30flag rescued MYC protein and mRNA expression in HL-60 cells with stable SBDS knockdown (Figure 4C and B), showing that MYC expression depends on p30. Similarly, cell multiplication was reduced upon SBDS knockdown in HL-60 cells, which could be restored by p30flag overexpression (Figure 4D).

To corroborate our data and literature (39) suggesting that p42 inhibits MYC expression, we examined MYC expression in a K562 chronic myeloid leukaemia (CML) cell line stably expressing a C/EBPα-p42-estrogen receptor (p42ER) fusion protein (39). Proteins fused to an oestrogen receptor domain (ER) are sequestered in the cytoplasm through binding to heat shock proteins (HSPs). Addition of the oestrogen receptor ligand β-estradiol to the cell culture medium results in relocation of p42ER to the nucleus and activation of C/EBPα-p42-transcriptional functions. Activation of p42ER by β-estradiol addition resulted in strong repression of MYC mRNA and protein, which was associated with reduced proliferation capacity (Supplementary Figure S2A–C). In the control experiment, activation of the unfused ER domain had no effect on MYC expression or cell proliferation (Supplementary Figure S2D–F).

To substantiate the association between expression of p30 and MYC, we reduced p30 expression by inhibition of mTOR signalling (22). Treatment of HL-60 cells with the mTOR inhibitor PP242 resulted in a reduced p30/p42 ratio, reduction of MYC mRNA and protein expression and reduced proliferation capacity (Figure 4E-G). Overexpression of p30flag rescued both MYC mRNA and protein levels despite the inhibition of mTOR by PP242 (Figure 4H and I). The proliferation capacity was partially restored by p30 overexpression (Figure 4J) suggesting that the C/EBPα isoform ratio is decisive for MYC expression in HL-60 cells which however only partially contributes to mTORC1 mediated proliferation control. Collectively the data suggest that MYC expression is regulated by SBDS through translational regulation of the p30/p42 C/EBPα isoform ratio.

Deregulation of C/EBPα p30 expression impedes neutrophil differentiation

HL-60 cells differentiate into the neutrophil lineage in response to ATRA treatment. During differentiation, the expression of p30, but not of p42 was reduced, resulting in a strong reduction in the p30/p42 ratio (Figure 5A). Transcript levels of C/EBPα were reduced at the final day 6 (Figure 5B). MYC mRNA and protein levels rapidly and strongly declined during differentiation (Figure 5A and B). Intriguingly, SBDS mRNA and protein levels were also reduced during differentiation (Figure 5A and B), as reported for other cell lines (42–44). Overexpression of p30flag resulted in delayed suppression of MYC mRNA (Figure 5A) and protein (Figure 5C). In addition, endogenous p30 and SBDS expression are delayed (Figure 5B and C). Furthermore, overexpression of p30flag is sufficient to sustain proliferation of HL-60 cells during ATRA treatment (Figure 5D). Compared with control HL-60 cells, at day seven of ATRA treatment p30flag overexpression induced a significant reduction in the number of terminally differentiated, proliferation-arrested cells, with expansion of proliferative myelocytes to more than 20% (Figure 5E). Finally, SBDS knockdown did not alter ATRA-induced differentiation of HL-60 cells (Figure 5F).

DISCUSSION

Recent studies have defined the role of SBDS in ribosomal maturation (9,10,13). However, so far a clear connection between reduced ribosomal SBDS function in SDS patients, deregulated translation and the disease phenotype is lacking. Our data for the first time demonstrate that translation of specific mRNAs, namely the C/EBPα and -β mRNAs, is affected by loss of SBDS function. Different isoforms, full-length (p42 and LAP) and truncated (p30 and LIP), are translated from both C/EBPα and -β mRNAs through the use of different translation initiation sites. Interestingly, reduced SBDS function by either shRNA mediated knockdown or by mutations in SDS patient-derived B-lymphoid cells affected in particular translation re-initiation into the truncated p30 and LIP isoforms leaving translation into the full-length isoforms by regular initiation largely unaffected. The same reduction in re-initiation but not initiation by SBDS knockdown was observed with luciferase reporters mimicking C/EBPα-mRNA (p42/p30 reporters) or C/EBPβ-mRNA (LAP/LIP reporters) regulation, rendering it independent of effects on protein turnover. Moreover, re-initiation from the reporters was restored in SBDS...
Figure 4. SBDS-dependent C/EBPα p30/p42 isoform ratio regulates MYC expression and cell proliferation in HL-60 cells. (A) Immunoblots showing endogenous MYC, C/EBPα, SBDS and α-Tubulin (loading control) protein expression in HL-60 cells with stable knockdown of SBDS (shSBDS) or control (scrambled shRNA). Quantification of the p30/p42 isoform ratio and MYC protein levels are shown on the right panel. (B) qRT-PCR analysis of endogenous MYC mRNA expression level in HL-60 cells and HL-60-p30flag overexpressing cells with stable knockdown of SBDS (shSBDS) or control (scrambled shRNA). Immunoblot shows endogenous MYC, C/EBPα, SBDS and α-tubulin (loading control) protein expression. Quantification of the p30/p42 isoform ratio and MYC protein levels are shown on the right panel. (D) Cell multiplication assay with HL-60 cells expressing scrambled shRNA (control), shSBDS or shSBDS-shCTRL. Seven days after seeding 5 × 10^5 cells, cells were harvested and counted. Different letters (a, b, c, d) indicate significant differences in cell numbers determined by one-way-ANOVA. Error bars represent ±SD (n = 3), P < 0.05. ** P < 0.01, *** P < 0.005.

Table: Relative transcript level

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SBDS-dependent C/EBPα p30/p42 isoform ratio regulates MYC expression and cell proliferation in HL-60 cells. (A) Immunoblots showing endogenous MYC, C/EBPα, SBDS and α-Tubulin (loading control) protein expression in HL-60 cells with stable knockdown of SBDS (shSBDS) or control (scrambled shRNA). Quantification of the p30/p42 isoform ratio and MYC protein levels are shown on the right panel. (B) qRT-PCR analysis of endogenous MYC mRNA expression level in HL-60 cells and HL-60-p30flag overexpressing cells with stable knockdown of SBDS (shSBDS) or control (scrambled shRNA). Immunoblot shows endogenous MYC, C/EBPα, SBDS and α-tubulin (loading control) protein expression. Quantification of the p30/p42 isoform ratio and MYC protein levels are shown on the right panel. (D) Cell multiplication assay with HL-60 cells expressing scrambled shRNA (control), shSBDS or shSBDS-shCTRL. Seven days after seeding 5 × 10^5 cells, cells were harvested and counted. Different letters (a, b, c, d) indicate significant differences in cell numbers determined by one-way-ANOVA. Error bars represent ±SD (n = 3), P < 0.05. ** P < 0.01, *** P < 0.005.

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Figure 5. SBDS dependent C/EBPa-p30 expression regulates MYC expression and cell proliferation during neutrophil differentiation. (A) HL-60 cells undergo neutrophil differentiation with 5 µM ATRA treatment. Immunoblots show expression of MYC, p42, p30, SBDS and α-tubulin (loading control) at days 0, 2, 4 and 6 of treatment (D = days). (B) qRT-PCR analysis of mRNA levels of MYC, C/EBPa, and SBDS during neutrophil differentiation. (C) Immunoblots show expression of MYC, p42, p30, SBDS or α-tubulin (loading control) in HL-60 control cells (left) or HL-60 cells stably overexpressing C/EBPa-p30flag (right). (D) Cell multiplication assay with HL-60 or HL-60 cells expressing C/EBPa-p30flag treated with 5 µM ATRA at day 0. At day 0, 5 × 10⁵ cells were seeded and cell numbers were determined at the indicated time points (D = day). (E) HL-60 or HL-60 expressing C/EBPa-p30flag treated with 5 µM ATRA for 7 days (D7). (F) HL-60 cells expressing shSBDS for stable knockdown of SBDS or expressing control scrambled shRNA treated with 5 µM ATRA for 7 days (D7). For (E) and (F) Giemsa staining was used for morphological analysis (magnification x200). At the right, the distribution of neutrophil phenotypes is shown. All asterisks indicate significant differences between control cells and p30flag overexpressing cells or control and shSBDS expressing cells determined by Student’s t-tests. Error bars represent ±SD (n = 3), *P < 0.05, **P < 0.01, ***P < 0.005.
knockdown cells by ectopic overexpression of SBDS. This suggests that reduced SBDS function does not in general affect all translation events but may specifically affect translation re-initiation. Prerequisite for translation re-initiation at the p30 and LIP initiation codons is an initial translation of a cis-regulatory small uORF in the C/EBPα and -β mRNAs. This generates post-termination ribosomal subunits that are able to scan the mRNA further to the downstream initiation codon and re-initiate translation.

So far we do not know which step of the C/EBPα/β mRNA-specific translational event is inhibited by the reduced SBDS function. It may involve reduced recognition of the uORF initiation codon, efficiency of re-initiation at the downstream initiation codon or scanning along the mRNA following uORF translation. It seems that re-initiation after uORF translation is regulated by different or additional mechanisms compared to general cap-dependent translation initiation and we are only at the beginning of understanding the underlying mechanisms. For example, for Drosophila it was shown that the DENR-MCT-1 complex specifically affects re-initiation events after uORF translation of a specific set of mRNAs in proliferating cells (45), and in Arabidopsis eIF3H is required for mTOR-mediated translation re-initiation (46). Future experiments have to address the molecular basis of the effect of SBDS depletion on C/EBPα and -β mRNA translation. In addition it will be interesting to see whether the translation of other mRNAs harbouring uORFs that promote re-initiation are sensitive to reduced SBDS function. Genome-wide ribosome profiling could reveal transcripts and specific translation modes that are affected by loss of SBDS function. SBDS promotes a conformational switch in the GTPase EFL1 that displaces eIF6 from the 60S subunit by competing for an overlapping binding site (12). Displacement of eIF6 allows joining of the 60S and 40S subunits to generate actively translating 80S ribosomes. In addition, it was postulated based on in vitro reconstitution experiments that eIF6 may act in the dissociation of post-termination ribosomes and thereby stimulates ribosomal subunit recycling (47). It has been shown recently that experimental reduction of eIF6 levels reduce expression levels of LIP (48). The authors discuss the possibility that the anti-association activity of eIF6 is required to circumvent formation of inactive 80S after uORF translation termination and allowing post-termination 40S to re-initiate at a downstream initiation codon. If eIF6 is indeed required for keeping post-termination 40S ribosomal subunits free for re-initiation at downstream sites then SBDS would facilitate this by releasing eIF6. However, other yet to be identified mechanisms could be responsible for the regulation of p30 and LIP expression by SBDS.

Another important question addresses the biological consequence of reduced C/EBPα and -β truncated isoform expression in the context of SBDS deficiency. Unfortunately, the SDS patient derived B-cells and the wild-type SBDS control cells do not express endogenous C/EBPα. Therefore, we examined the biological effects of reduced SBDS function in the promyelocytic leukaemia cell line HL-60 in response to shRNA mediated SBDS knockdown because this cell line expresses both C/EBPα and -β endogenously. Knockdown of SBDS delayed increase in viable cells counts compared to control shRNA-expressing cells, indicating that cell proliferation was attenuated in the SBDS knockdown cells. Attenuation of cell proliferation correlated with the decreased expression of the proto-oncogene MYC both at the mRNA and protein level in response to SBDS knockdown. The effect on MYC protein levels seems to be much stronger than observed for mRNA levels (Figure 4, A, C and E), suggesting that additional regulation at the post-transcriptional level may be involved. It has been shown earlier that MYC is a transcriptional target of both C/EBPα and -β and that the full-length isoforms act as repressors of MYC transcription (25,32). Furthermore, alteration of the C/EBPα isoform ratio through treatment of HL-60 cells with the drug CDDO that downregulates p30 expression strongly reduces MYC mRNA levels and inhibits proliferation (41), similar to what we observed in response to the reduction of p30 expression through mTOR inhibition. Therefore, we hypothesize that the altered C/EBPα ratio in response to SBDS knockdown leads to the inhibition of MYC expression, which then results in the attenuation of proliferation. This hypothesis is supported by the observation that constitutive expression of the p30flag isoform restores both MYC expression and proliferation in SBDS knockdown cells. Interestingly, decreased MYC mRNA levels were also observed in embryonic brain, liver, cartilage and bone tissue from an SDS mouse model (compound heterozygotes for SBDS null and SBDS-R126T alleles) and in the pancreas of a pancreas-specific version of this SMD mouse model. Decreased MYC expression in the pancreas was, however, shown to depend on the upregulation of p53 upon loss of SBDS function (49). HL-60 cells don’t express functional p53 indicating that impairment of SBDS function might result in the downregulation of MYC and inhibition of proliferation by a different mechanism, which as we propose is the changed C/EBP isoform ratio. Both p42 and LAP have been shown to inhibit cell proliferation through repression of E2F target genes, while p30 and LIP counteract this function and thereby support proliferation (31,50). Therefore, apart from the downregulation of MYC, the reduced expression of other E2F target genes in response to the changed C/EBPα/β isoform ratio is probably involved in slowing down cell proliferation upon SBDS deficiency.

A prominent feature of SDS patients is bone marrow failure associated with neutropenia. Such a neutropenia could be caused by a failure of progenitor cells to either differentiate or to proliferate because the latter situation would strongly reduce the number of cells that undergo differentiation. Contradicting results have been published concerning the cause of neutropenia. Studies using shRNA-mediated downregulation of SBDS expression in murine hematopoietic progenitors or using an sbds deficient mouse model with deletion of SBDS predominantly in the myeloid lineage suggest that loss of SBDS interferes with terminal differentiation of neutrophil (51,52). Although the exact mechanism of this differentiation block is not known, it is accompanied with reduced expression of the myeloid transcription factor retinoid acid receptor α mRNA, upregulation of p53 and induction of apoptosis (52). On the other hand, it was shown that haematopoietic progenitors derived from SDS patients have proliferation defects (42). Similarly, SBDS knockdown
in the 32D myeloblast cell line resulted in reduced proliferation without affecting differentiation (44). The observation that SBDS expression is high in undifferentiated 32D cells but decreases upon differentiation (44) also speaks in favour for a function in proliferating cells.

C/EBPa is a decisive factor for neutrophil differentiation that at the same time is involved in proliferation control. Its deletion in mice results in a block in granulocyte differentiation and enhanced self-renewal capacity of haematopoietic stem cells (53). The p42 isoform of C/EBPα is a decisive factor for neutrophil differentiation and enhanced self-renewal capacity of haematopoietic stem cells (53). The p42 isoform of C/EBPα induces the transcription of CSF3R, C/EBPa and NE (neutrophil elastase), which are decisive for differentiation (24,54). On the other hand, mutations of C/EBPa have been detected in AML patients, some of the mutations resulting in expression of only the truncated isoform and preventing expression of the p42 isoform (55). Knockin mice expressing only p30 isoform develop leukaemia and it has been suggested that p30 acts as a negative inhibitor of p42 by blocking the expression of differentiation related genes and by stimulating proliferation (26).

Interestingly, during the differentiation of HL-60 cell, we observed downregulation of SBDS expression similarly to what has been shown in 32D cells (44) with high levels of SBDS correlating with high levels of p30 and high levels of MYC in the early phase where the cells still proliferate. However, at later stages of the differentiation process when cells stop proliferating, SBDS was downregulated concomitantly with p30 and MYC. Thus, our results support the idea that in SDS patients the reduced levels of SBDS protein preferentially affect cell proliferation and thus expansion of the early progenitor pool rather than the differentiation process per se.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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REFERENCES


