Antisense targeting of decoy exons can reduce intron retention and increase protein expression in human erythroblasts

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ABSTRACT

The decoy exon model has been proposed to regulate a subset of intron retention (IR) events involving predominantly larger introns (>1 kb). Splicing reporter studies have shown that decoy splice sites are essential for activity, suggesting that decoys act by engaging intron-terminal splice sites and competing with cross-intron interactions required for intron excision. The decoy model predicts that antisense oligonucleotides may be able to block decoy splice sites in endogenous pre-mRNA, thereby reducing IR and increasing productive gene expression. Indeed, we now demonstrate that targeting a decoy 5′ splice site in the O-GlcNAc transferase (OGT) gene reduced IR from ~80% to ~20% in primary human erythroblasts, accompanied by increases in spliced OGT RNA and OGT protein expression. The remaining OGT IR was refractory to antisense treatment and might be mediated by independent mechanism(s). In contrast, other retained introns were strongly dependent on decoy function, since antisense targeting of decoy 5′ splice sites greatly reduced (SNRNP70) or nearly eliminated (SF3B1) IR in two widely expressed splicing factors, and also greatly reduced IR in transcripts encoding the erythroid-specific structural protein, α-spectrin (SPTA1). These results show that modulating decoy exon function can dramatically alter IR and suggest that dynamic regulation of decoy exons could be a mechanism to fine-tune gene expression post-transcriptionally in many cell types.

Keywords: SF3B1; alternative splicing; decoy exon; erythroid gene expression; intron retention

INTRODUCTION

Gene expression is determined not only by transcription rate, but also by post-transcriptional processes including the efficiency with which pre-mRNA is spliced into translatable mRNA. Intron retention (IR) is a form of RNA processing that selectively modulates splicing of specific introns (Braunschweig et al. 2014; Boutz et al. 2015; Mauger et al. 2016; Jacob and Smith 2017), in essence rendering them “alternative introns.” By regulating the efficiency of intron splicing, cells can alter the balance between two competing pathways: one that generates fully spliced mRNA that can be translated into protein, and a second that produces incompletely spliced “intron retention” transcripts (IR-transcripts). Most of the latter contain premature translation termination signals that preclude synthesis of full length protein. IR-transcripts that are otherwise spliced and polyadenylated can experience several fates in different cellular contexts. Such transcripts are often detained in the nucleus, where they may be degraded (Pendleton et al. 2018) or they may serve as a reservoir for new mRNA production via excision of the retained intron(s) (Ninomiya et al. 2011; Boothby et al. 2013; Mauger et al. 2016); in many cases, the fate is unknown. Alternatively, IR transcripts can be exported to the cytoplasm for degradation by nonsense-mediated decay (NMD) (Wong et al. 2013), or they may persist for translation (Rekosh and Hammarskjold 2018) or other unknown functions (Brugiolo et al. 2017). At a constant transcription rate, greater diversion of pre-mRNA into untranslated IR-transcripts should reduce output of mRNA and decrease protein synthesis. Coordinate regulation of IR can effect programmed changes in gene expression patterns during normal development as cells differentiate and respond to
environmental signals (Wong et al. 2013; Shalgi et al. 2014; Boutz et al. 2015; Edwards et al. 2016; Mauger et al. 2016; Ni et al. 2016; Pimentel et al. 2016; Braun et al. 2017; Ni et al. 2017; Pimentel et al. 2017). Conversely, aberrations in the IR program are observed in many diseases including cancers where they can adversely impact expression of many genes (Dvinge and Bradley 2015; Luisier et al. 2018; Adusumalli et al. 2019). Although mechanisms of IR are not well understood, RNA binding proteins (RBPs) (Cho et al. 2014; Pendleton et al. 2017) and factors that modify RBPs (Braunschweig et al. 2014) have been shown impact IR. In a few cases, RNA sequence elements required for regulating individual intron retention events have been identified (Park et al. 2017; Pendleton et al. 2017; Parra et al. 2018; Rekosh and Hammerskold 2018).

Analysis of RNA-seq profiles from differentiating erythroid cell populations revealed highly dynamic, global changes in the erythroid transcriptome, including changes in RNA processing of both cassette exons and retained introns, as the cells undergo extensive remodeling during the final cell divisions prior to enucleation (An et al. 2014; Pimentel et al. 2014, 2016; Edwards et al. 2016). The IR program encompasses hundreds of IR-transcripts that are polyadenylated and spliced except for selective retention of one or more introns (Edwards et al. 2016; Pimentel et al. 2016). In late erythroblasts, numerous IR transcripts are abundantly expressed, many of which comprise ≥25% of the steady state RNA from their cognate genes. Some of these are dynamically regulated during terminal erythropoiesis, while others exhibit stable IR levels, indicating multiple regulatory pathways (Pimentel et al. 2016). While the majority of erythroblast retained introns are short (<1 kb), as observed in other systems (Braunschweig et al. 2014), a subset of important erythroid genes exhibit larger retained introns having embedded decoy exon(s) that are essential for retention (Parra et al. 2018). According to the decoy model, cryptic decoy exon(s) interact nonproductively with intron-terminal splice sites, engaging them in a manner that fails to stimulate efficient splicing catalysis. By competing with cross intron interactions necessary for intron removal, decoy interactions promote IR. Supporting evidence for this model includes the ability of decoy exons to activate IR in heterologous splicing reporters; the dependence of this IR activity on intact decoy splice sites; and the enrichment of U2AF binding at 3′ splice sites of decoy exons (Parra et al. 2018).

We explored the hypothesis that antisense targeting of deep intron splice sites, here referring to those associated with decoy exons, can alter endogenous RNA processing fates in primary erythroid progenitors so as to tune gene expression. Previous studies have shown that deep intron splice sites are poorly represented in most RNA-seq data sets, yet they are critical features of recursive splicing pathways that excise selected long introns (Hatton et al. 1998; Burnette et al. 2005; Suzuki et al. 2013; Sibley et al. 2015; Joseph et al. 2018). Deep intron splice sites also have essential functions during catalysis of intermediate steps of nested intrasplicaing pathways (Parra et al. 2008, 2012). Given the ability of antisense oligonucleotides to alter RNA processing outcomes in these pathways by masking deep intron splice sites (Parra et al. 2012; Sibley et al. 2015), we used a similar strategy to test whether targeting decoy splice sites with antisense reagents can inhibit IR. New results indicate that blocking highly conserved decoy exons in three broadly expressed genes (SF3B1, OGT, and SNRNP70), and in an erythroid-specific gene (SPTA1), greatly reduces intron retention activity in endogenous transcripts, and can increase spliced RNA and protein expression. These results validate the function of decoy exons in the context of their natural endogenous transcripts, and suggest that many of the ~400 predicted decoys in differentiating human erythroblasts could be regulated to impact protein expression.

RESULTS

Decoy exon targeting strategy

Candidate decoy exons were identified in retained introns of NMD-inhibited erythroblasts by virtue of the novel splice junctions created when they splice, albeit inefficiently, to adjacent exons (Parra et al. 2018). The decoy model hypothesizes that their main function is to form early spliceosomal complexes with intron-terminal splice sites that become arrested at a precatalytic stage of assembly; catalytic splicing at decoy splice sites is inefficient and typically leads to NMD. To assess decoy function in endogenous erythroid transcripts, we reasoned that antisense oligonucleotides targeting decoy exon splice sites should interfere with IR to reduce retention efficiency. To maximize our ability to detect such changes, we selected IR-transcripts meeting the following criteria: (i) The transcript must possess a unique intron exhibiting ≥20% retention in late erythroblasts; (ii) its cognate gene must be expressed in moderate to high abundance; and (iii) the embedded decoy exons must have simple splice site architecture. The last feature served to restrict analysis to decoys that either have unique splice junctions, or have closely spaced alternative junctions that can be blocked with a single 25 nt antisense morpholino (MO). This design was expected to maximize the likelihood of blocking spliceosome assembly at the decoy. However, it eliminated from consideration strong decoys in ARGLU1 and DDX39B that possess alternative splice sites distributed over a wider range (Pimie et al. 2017; Parra et al. 2018). Finally, we targeted 5′ splice sites, because the relatively low GC content at the 3′ splice sites of these decoy exons was predicted to reduce MO affinity and effectiveness.

Figure 1A shows relevant features of the IR regions from four genes chosen for analysis. OGT intron 4 (3.3 kb),
SF3B1 intron 4 (1.8 kb), SNRNP70 intron 7 (3.2 kb), and SPTA1 intron 20 (1.8 kb) all exhibit substantial retention in erythroid progenitors at day 9 of the culture (D9) and in well-differentiated erythroblasts at day 16 (D16). Each of these introns encodes decoy exon(s), not represented in Refseq annotations, that were defined by analysis of splice junction reads (Parra et al. 2018) and are depicted in a custom reannotation track (Fig. 1A). The decoys in OGT, SF3B1, and SNRNP70 have been highly conserved from fish to mammals, while the SPTA1 decoy is conserved only among mammals. In previous assays with splicing reporters, the OGT and SF3B1 decoys exhibited strong IR activity, while the activity of the SNRNP70 decoy had weaker activity (Parra et al. 2018). The SPTA1 decoy has not been assayed previously for IR activity.

The 5′ splice site regions of decoys targeted in this study are shown in Figure 1B. The SF3B1 decoy exhibits only one 5′ splice site, while the other three decoys all have alternative 5′ splice sites located within 7–12 nt of each other. The presence of multiple splice sites could be integral to the decoy mechanism, since this appeared to be a frequent feature of decoy exons, and because it has been shown that concurrent occupancy of alternative splice sites can inhibit splicing (Chen et al. 2017). The shaded regions indicate sequences targeted by antisense MOs in the IR assays below.

**Reduction of intron retention by antisense targeting of decoy 5′ splice sites**

Primary human erythroid cultures were electroporated with antisense MOs and cultured for 2 d under standard conditions. RNA was then isolated for analysis by RT-PCR to investigate changes in the balance between IR-transcripts and spliced transcripts. We studied the effects of decoy targeting in four different genes using this approach. The targeting scheme and PCR strategy for analysis of IR in the OGT gene, which encodes O-GlcNAC transferase, is shown in Figure 2A. The decoy in OGT intron 4 exhibited strong IR activity in minigene splicing reporters (Parra et al. 2018). In endogenous OGT transcripts, we first assessed retention of the full length intron 4 by standard RT-PCR analysis under conditions that interrogate the E3–E6 region. Control cells treated with an irrelevant MO yielded two major OGT amplification products: a short product representing spliced mRNA, and a larger product corresponding to an IR transcript in which introns 3 and 5 were removed but intron 4 retained (Fig. 2B, lane 1). Cells treated with the OGT decoy-specific MO exhibited a substantial decrease in the IR isoform (Fig. 2B, lane 2). In contrast, the OGT MO did not alter retention of a heterologous decoy-containing intron in the SF3B1 gene, confirming specificity of the MO effects on IR (Parra et al. 2018). In endogenous OGT transcripts, we first assessed retention of the full length intron 4 by standard RT-PCR analysis under conditions that interrogate the E3–E6 region. Control cells treated with an irrelevant MO yielded two major

**FIGURE 1.** Intron retention and candidate decoy exons in targeted erythroid genes. (A) Annotation of key features in IR regions of four prominent erythroid genes. (Top panel) Refseq gene annotations, lacking indications of intron retention isoforms or decoy exons predicted within the introns. For SF3B1, only the four most frequently spliced decoys are shown, from a total of six (Parra et al. 2018). Lower panels show a reannotation that includes predicted decoy exons (boxed), RNA-seq data from early stage (D9) and late stage (D15) erythroblasts, and phylogenetic conservation of the relevant gene regions. (B) 5′ splice site features of targeted decoy exons. Upper case, decoy exon sequence; lower case, downstream intron sequence. Vertical bars show 5′ splice site junctions identified in RNA-seq data from erythroblasts inhibited for nonsense-mediated decay. Shaded regions indicate regions targeted by antisense MOs.
when the MO concentration was doubled (results not shown), suggesting that a component of OGT IR is modulated in a decoy-independent manner.

The next decoy-mediated IR event selected for analysis was in the SF3B1 gene (Fig. 3A). We focused on decoy exon 4e, shown previously to exhibit the strongest IR activity among several potential decoys in SF3B1 intron 4 (Parra et al. 2018). Cells treated with the SF3B1-specific MO exhibited much-reduced amounts of the IR-transcript when examined by standard RT-PCR (Fig. 3B, compare lanes 1 and 2). Quantitation by qPCR yielded a different result than was observed for OGT (Fig. 3C), since PIR in controls cells (~26%) was almost eliminated by the SF3B1 decoy 5′ splice site MO (~3%).

The two remaining targets represented decoys about which less prior information was known than for OGT and SF3B1. The predicted decoy exon in SNRNP70 is 60/72 nt, depending on alternative 5′ splice site choice, and might have unique properties since retention has been observed primarily only for downstream intron sequences (Fig. 1A). Moreover, this decoy exhibited only weak IR activity in a heterologous splicing reporter (Parra et al. 2018). For SPTA1, an 80/87 nt noncoding decoy exon mapping near the 3′ end of retained intron 20 was predicted on the basis of splice junction reads. A few RNA-seq reads spanned the SPTA1 decoy exon and linked it to both exon 19 upstream and exon 20 downstream, confirming its potential to be spliced at low frequency (data not shown). Given that SPTA1 encodes an abundant and erythroid-specific structural protein, α spectrin, control of IR could be important in regulating assembly of the erythroid membrane skeleton during terminal erythropoiesis.

Figure 4A,B show the targeting approach and PCR strategies used to test IR-promoting activity for predicted decoy exons in SNRNP70 and SPTA1. The effects of decoy-specific antisense MOs were assessed by RT-qPCR to quantitate both IR transcripts and fully spliced transcripts (Fig. 4C, D). Electroporation of human erythroid blasts with a MO against the 5′ splice site region of SNRNP70′s decoy substantially reduced the level of IR from ~35% in control erythroid cells to ~9% in MO-treated cells (Fig. 4C). Interestingly, the SPTA1 decoy-specific MO also strongly inhibited IR, from ~20% down to only ~2% (Fig. 4D). Together these results strongly support the hypothesis that decoy exons represent a novel regulatory component of the gene expression program, in which they can quantitatively modulate mRNA expression levels by tuning the splicing efficiency of key retained introns.

Impact of decoy targeting on expression of spliced RNA and protein

The dramatic reduction in IR for several genes suggests that inhibition of decoy exon function should lead to increased expression of spliced mRNA and increased capacity for protein synthesis. We explored this issue using OGT as a model, since the large MO-induced reduction in PIR would be expected to yield a significant increase in protein expression. Based on the fourfold difference in IR between control cells and cells treated with the OGT decoy-specific MO, measured at 48 h post-electroporation, one might predict a similar fourfold increase in OGT mRNA and OGT protein. Analysis of qPCR data revealed that the spliced OGT transcripts were actually increased 1.7–2.7-fold, when normalized to actin transcript expression in the same cells (Fig. 5A). To explore the reason for the modest discrepancy in expected versus observed expression,
we quantitated total OGT transcript levels (spliced plus IR transcripts) and found that overall OGT RNA expression was reduced in comparison to control cells. Therefore, the fourfold increase in splicing efficiency was partially offset by reduced steady state OGT RNA levels, presumably due to other compensatory mechanisms as part of O-GlcNAc homeostasis. Nevertheless, this result confirms that regulation of decoy-mediated IR effected a significant change in spliced OGT RNA expression.

Finally, we assessed erythroblast OGT protein expression as a function of variation in IR efficiency. Equal amounts of protein from control MO- or OGT MO-treated cells were immunoblotted with anti-OGT antibodies (Fig. 5B). Densitometric analysis of OGT expression, normalized to expression of a control protein, GAPDH, revealed ~1.4-fold increase in OGT protein levels in two independent experiments (compare control lanes to OGT MO lanes). Interestingly, it has been shown that OGT intron retention can also be reduced, and OGT protein expression elevated, by pharmacological treatment of nonerythroid cell lines with the OGT enzyme inhibitors OSMI-1 and OSMI-2 (Park et al. 2017; Tan et al. 2020). This effect has been interpreted as a compensatory mechanism for cells to respond to reduced OGT enzyme activity, that is, they modify RNA processing pathways so as to reduce OGT IR and stimulate production of more total OGT protein. We found that reducing OGT IR independently through the use of decoy-targeting MOs had a similar stimulatory effect on OGT protein expression in erythroblasts (compare Fig. 5B,C), suggesting that the effects of OSMI-1 may be mediated through the decoy exon mechanism.

**DISCUSSION**

The decoy exon model proposes that intron retention levels can be controlled by modulating the balance between two competing splice site interactions: (i) productive cross-intron interactions, involving annotated splice sites at the intron termini, that promote splicing catalysis to remove the intron, and (ii) nonproductive interactions, involving contacts between internal decoy exon(s) and intron terminal splice sites, that function mainly to block intron excision and promote intron retention. The latter are spliced inefficiently or not at all, presumably due to arrest of spliceosomal assembly at a precatalytic complex by mechanisms yet to be defined. The current study validates a major prediction of the decoy hypothesis, namely, that blocking decoy exon function in endogenous pre-mRNA should shift RNA processing in favor of better intron removal. All four genes targeted with decoy exon 5′ splice site-specific antisense MOs were shown to exhibit substantial decreases in IR. Targeting decoy 3′ splice sites should in principle also abrogate intron retention, based on experiments with minigene splicing reporters showing that 3′ splice site mutations disrupt IR (Parra et al. 2018). However, preliminary experiments with 3′ targeting MOs thus far have yielded mixed results: All three tested were less effective than the corresponding 5′-targeting reagents, likely due to lower predicted base-pairing affinity and/or undesirable self-complementarity properties of these particular decoy sequences (results not shown). Interestingly, for two genes (SF3B1 and SPTA1), blocking the decoy exon 5′ splice site essentially eliminated IR, suggesting that the decoy pathway may be the sole determinant of IR. In contrast, IR was not completely abrogated by antisense treatment in the OGT gene, consistent with the coexistence of decoy-independent IR mechanisms (Cho et al. 2014; Braun et al. 2017; Wong et al. 2017; Monteuuis et al. 2019).
Importantly, in the one case tested, OGT, decreased IR was accompanied by increases in spliced RNA and protein expression.

The IR transcripts studied here regulate expression of genes with diverse roles in erythropoiesis. Three function in general biochemical processes such as O-GlcNAc homeostasis (OGT) and pre-mRNA splicing (SF3B1 and SNRNP70), that are widely important in both erythroid and nonerythroid cells. Presumably the decoy mechanism actively regulates these genes in many different cell types. In contrast, SPTA1 functions predominantly in erythroid cells where it encodes a major structural component of the membrane skeleton that mechanically supports the eventual red cell membrane. Given the measured PIR values of 25%-75% in these genes, full inhibition of IR could lead to 1.3- to fourfold increases in protein expression, with most genes capable of less than or equal to twofold changes based on these bulk measures of IR. We speculate that the major purpose of decoy-mediated IR may be fine-tuning of expression according to the cell’s physiological needs. In fact, IR has already been shown to tune OGT expression in cancer cell lines via an intronic element (Park et al. 2017) that likely operates via the decoy mechanism. For SPTA1, decoy-mediated intron retention could function in a similar manner to balance expression of the α and β spectrin chains, two high molecular weight proteins that form an extended heterodimer that assembles into higher order structures supporting the red cell membrane. An imbalance of spectrin chains might be detrimental to human erythroblasts, and control of IR could serve to equalize the cellular content of these binding partner proteins. Finally, differentiating erythroblasts might dynamically regulate IR for splicing factor genes so that RNA splicing capacity could adapt to changes pre-mRNA abundance as thousands of genes are down-regulated during terminal erythropoiesis (An et al. 2014).

Interestingly, global comparison of RNA and protein abundance profiles in differentiating human erythroblasts has revealed discordant expression patterns that can be explained in part by IR (Gautier et al. 2016). Profiling experiments have shown that genes displaying increased RNA levels but decreased protein expression, as cells progressively differentiate into late stage erythroblasts, are enriched in IR-transcripts. In such cases, IR may function to down-regulate productive gene expression in a post-transcriptional manner. We propose that decoy-mediated IR contributes substantially to this phenomenon, since erythroblasts express an estimated 400 retained introns embedded with candidate decoy exon(s) (Parra et al. 2018). Moreover, the number of functional decoys could be greater, because many intronic U2AF binding sites detected in K562 cells do not align with splice junction-predicted erythroblast decoy exons. These U2AF sites of unknown function, perhaps regulated by novel RBP cofactors (Sutandy et al. 2018), could

![FIGURE 4.](https://example.com/figure4)

**FIGURE 4.** Antisense inhibition of SNRNP70 IR and SPTA1 IR. (A) SNRNP70 gene structure in the IR region showing retained intron 7 (thick gray line) with its major decoy exon, together with adjacent introns and exons. Position of antisense MO designed to block the 5′ splice site is shown, along with primer pairs used for RT-PCR. (B) SPTA1 gene structure in the IR region showing retained intron 20 (thick gray line) with its major decoy exon, together with adjacent introns and exons. Position of antisense MO designed to block the 5′ splice site is shown, along with primer pairs used for RT-PCR. (C,D) IR in cells treated with SNRNP70-specific (C) or SPTA1-specific (D) MO, in parallel with cells subjected to control MO treatment. IR was assessed using RT-qPCR to compare the relative amounts of IR and spliced products.
"silent" decoys that promote IR without ever being catalytically spliced. Preliminary experiments supporting this idea are under further investigation.

The discovery of decoy exons provides new evidence that many unannotated splicing elements reside in deep intron space, hundreds to many thousands of nucleotides from the regulated splice sites, and that they play essential roles in regulating proper splice patterns via several mechanisms (Ule and Blencowe 2019). The decoy model discussed here is presumably used in diverse cell types, since many decoy-containing introns are retained in a wide range of nonerythroid cell types. Another mechanism dependent on deep intron splicing elements is recursive splicing (RS). RS involves functional recognition of RS-exons, embedded deep within long introns, as critical splicing intermediates (Sibley et al. 2015; Joseph et al. 2018). Finally, a mechanism termed intrasplicing requires deep intron splicing elements, located tens of kilobases upstream of the regulated splice acceptors, to promote nested splicing reactions required for proper splice site selection in two paralogs of the protein 4.1 gene family (Parra et al. 2008, 2012). In various contexts, antisense oligonucleotides that block deep intron elements have been used to demonstrate their functional importance in splicing of endogenous pre-mRNAs (Parra et al. 2012; Lovci et al. 2013; Sibley et al. 2015).

Finally, the current results suggest new clinical applications of antisense reagents for the purpose of improving gene expression. Pioneering work more than 20 yr ago showed that antisense oligonucleotides can block aberrant splice sites to restore correct splicing of the erythroid β-globin gene in thalassemic pre-mRNA (Dominski and Kole 1993). Since that time an increasing array of antisense strategies has been utilized to increase proper gene expression by manipulating RNA processing. Antisense oligonucleotides can induce exon skipping in dystrophin pre-mRNA to restore the translational reading frame in patients with Duchenne muscular dystrophy (e.g., Aartsma-Rus and Krieg 2017); they can mask intronic splicing silencer(s) downstream from SMN2 exon 7 to improve productive splicing in patients with spinal muscular atrophy (Bennett et al. 2019); and, given increasing evidence that deep intron mutations can cause human disease (Vaz-Drago et al. 2017), antisense reagents can also improve gene expression by blocking deep intron splicing mutations that activate inclusion of cryptic noncoding exons in the breast cancer gene BRCA2 (Anczukow et al. 2012) and the deaf-blindness gene USH2A (Slijkerman et al. 2016). As mentioned in the introduction, antisense reagents can mask deep intron splice sites so as to modulate RNA processing during recursive splicing and intrasplcing (Parra et al. 2012; Sibley et al. 2015). Antisense reagents can even increase productive gene expression by inhibiting nonsense-mediated mRNA decay (Nomakuchi et al. 2016). Here we have shown that antisense oligonucleotides can increase protein expression by blocking decoy splice sites in retained introns, which could allow a functional allele to increase protein output to compensate for genetic deficiencies in various disease states. Relevant disease phenotypes may arise due to haploinsufficiency or, as in the case of hereditary spherocytosis associated with α-spectrin deficiency, they may be due to biallelic SPTA1 mutations (Chonat et al. 2019), some of which induce aberrant splicing by activating distal branch points (Gallagher et al. 2019). Since disease severity correlates with the level of α-spectrin protein in the patient’s red blood cell cytoskeleton (Chonat et al. 2019), blocking decoy-mediated IR in SPTA1 could have therapeutic value.
MATERIALS AND METHODS

Erythroblast culture

CD34+ erythroid progenitors were enriched from cord blood and cultured under conditions previously shown to support selective growth and differentiation of erythroid cells (Hu et al. 2013). For electroporation, 10^6 erythroblasts at day 11 of culture were electroporated at room temperature in supplemented P3 solution using a Lonza 4D-Nucleofector system with the ER 100 pulse code. MOs 25 nt in length were obtained from Gene Tools LLC, maintained in sterile saline solution, and added to the cells at 30 µM final concentration just prior to electroporation. After electroporation cells were incubated in culture medium at 37°C for 2 d before further processing. When RNA and protein were isolated from the same sample, ∼2.5 × 10^5 cells were used for the RNA preparation and 7.5 × 10^5 cells for protein purification. Morpholino sequences antisense to the 5′ splice sites of the targeted decoys were as follows:

- OGT, 5′-gtggcagttacaaac|ccgttac|CAT-3′;
- SPTA1, 5′-ctggctggaac|ctcttac|GTGGCTG-3′;
- SF3B1, 5′-atccggaatactag|ACTTTCGTGC-3′;
- SNRNP70, 5′-ccatgatac|aaac|CCTTATACCAAC-3′.

Sequences antisense to the intron are in lower case; sequences antisense to the exon are in upper case; EXON|intron boundaries are marked by vertical lines.

RT-PCR

Total RNA was extracted from human erythroid progenitors and analyzed by standard RT-PCR methods as previously described (Pimentel et al. 2016). For quantitative analysis, RT-qPCR was performed using an Applied Biosystems 7500 Fast Real-Time PCR System with Quanta SYBR Green Fastmix low ROX reagents. The SYBR Green buffers were supplemented with forward and reverse primers at a final concentration of 0.5 µM, and the spliced RNA or IR-RNA DNAs amplified using the following program: initial denaturation (1 cycle): 94°C for 15 min; amplification stage (40 cycles): 95°C for 10 sec, 60°C for 25 sec, and 72°C for 30 sec; final extension at 72°C for 30 sec. Size and identity of qPCR products (Table 1) were confirmed by gel electrophoresis (Supplemental Fig. S1) and by DNA sequencing. The relative expression of each gene was calculated using the comparative ΔCt method after normalizing to the ACTB control.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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### TABLE 1. Primers used for amplification of spliced and IR products

| Product | Size (nt) | Primer location and sequence | Primer location and sequence |
|---------|----------|------------------------------|------------------------------|
| SF3B1-spl | 138 | forward E4/5F | TCTTTTGATCCCTTTTGGACAATGGAGG |
| SF3B1-IR | 173 | reverse B6R | GCTGAAAATTTTCTGGTAAGCTGG |
| OGT-spl | 159 | forward E3/4F | TCTCTCTTTGAGAACATCTGATGG |
| OGT-IR | 171 | reverse E5R | AATATTCTGGACATGGCCAG |
| SNRP70-IR | 149 | forward E7/8F | AGCTGAAAATTTTCTGGTAAGCTGG |
| SPTA1-spl | 178 | reverse E2/3F | CATTTGCTGTCGTCTGGTT |
| SPTA1-IR | 166 | reverse E8/9R | AGCAATGACAGCTGGGGGCTG |
| ACTB | 187 | reverse E8/9 | ACAGACACATATGGGCTG |

Upper part of the table represents primers used to amplify small products via RT-qPCR. Lower table represents primers used for standard RT-PCR. Primer designations: E, exon; i, intron; F, forward; R, reverse. The symbol “/” indicates a primer that overlaps two exons and can only amplify spliced transcripts.
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Targeting decoy exons reduces intron retention

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