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RNA-binding protein RBM20 represses splicing to orchestrate cardiac pre-mRNA processing

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Mutations in the gene encoding the RNA-binding protein RBM20 have been implicated in dilated cardiomyopathy (DCM), a major cause of chronic heart failure, presumably through altering cardiac RNA splicing. Here, we combined transcriptome-wide crosslinking immunoprecipitation (CLIP-seq), RNA-seq, and quantitative proteomics in cell culture and rat and human hearts to examine how RBM20 regulates alternative splicing in the heart. Our analyses revealed the presence of a distinct RBM20 RNA-recognition element that is predominantly found within intronic binding sites and linked to repression of exon splicing with RBM20 binding near 3′ and 5′ splice sites. Proteomic analysis determined that RBM20 interacts with both U1 and U2 small nuclear ribonucleic particles (snRNPs) and suggested that RBM20-dependent splicing repression occurs through spliceosome stalling at complex A. Direct RBM20 targets included several genes previously shown to be involved in DCM as well as genes not typically associated with this disease. In failing human hearts, reduced expression of RBM20 affected alternative splicing of several direct targets, indicating that differences in RBM20 expression may affect cardiac function. Together, these findings identify RBM20-regulated targets and provide insight into the pathogenesis of human heart failure.

Introduction

Alternative RNA processing is determined by RNA-binding splicing factors to generate different mRNA isoforms in a tissue-specific and developmentally-regulated manner (1). Alterations in splicing factors or mutations in their target sequences result in misspliced gene products, which is the underlying cause of several human genetic diseases (2–4).

RBM20 encodes the RNA-binding motif protein 20 with highly conserved functional domains: an RNA-recognition motif 1 (RRM-1) and an arginine/serine-rich (RS-rich) region. These structural features are characteristic of a family of SR and SR-related RNA-binding proteins (RBP) that assemble in the spliceosome and orchestrate the splicing of pre-mRNA. RBM20 is predominantly expressed in striated muscle, with the highest levels in the heart (5). Recently, mutations in RBM20 have been shown to cause human dilated cardiomyopathy (DCM) (6, 7), a major cause of heart failure and significant source of mortality and morbidity worldwide (8). Given the role of RBM20 in cardiomyopathy, efforts toward the identification of its RNA targets have been employed (5, 9) with the goal that their discovery would shed light upon human heart failure mechanisms and provide options for molecular therapy. No precise RNA recognition element (RRE) has been defined for RBM20, and with titin (Ttn), only a single bona fide mRNA target has been confirmed (5, 7). No protein interaction partners have been identified for RBM20, despite the fact that previously reported RBM20 mutations associated with cardiomyopathy cluster in the RS-rich region (7) that is predicted to mediate protein-protein interactions (7, 10–12).

Here, we reveal the previously elusive RBM20 RRE at nucleotide resolution using photoactivatable ribonucleoside–enhanced crosslinking and immunoprecipitation (PAR-CLIP) (13, 14) and transcriptome-wide RBM20-binding sites in heart-specific transcripts using high-throughput sequencing of RNA isolated by crosslinking immunoprecipitation (HITS-CLIP) (15). To identify direct regulatory targets of RBM20, we integrated the crosslinking immunoprecipitation (CLIP-seq) data with the analysis of transcriptomes of heart tissue obtained from WT rats and rats deficient for RBM20 protein as well as human heart failure patients. These data yielded a transcriptome-wide RNA map for RBM20-dependent splicing regulation in cardiac tissue. Using quantitative SILAC-based proteomics (where SILAC indicates stable isotope labeling by amino acids in cell culture) (16), we discovered RBM20 protein-binding partners. We revealed that RBM20 recognizes splicing silencers at defined intronic positions and interacts with components of the spliceosomal complex A but not complex B. Importantly, patients with severe heart failure and low levels of RBM20 gene expression in heart tissue show a loss of splicing...
mapped to the human genome with up to 1 mismatch. A hallmark of PAR-CLIP is the occurrence of diagnostic nucleotide transitions of crosslinked thio-nucleosides during reverse transcription. The PAR-CLIP reads aligned with T to C mismatches in 4SU, and G to A in 6SG libraries, respectively, indicating specific mRNA-RBM20 crosslinking (Supplemental Figure 1C). To investigate only those binding sites that were supported by reads from at least 2 out of 3 experiments, we computationally analyzed all 3 libraries jointly (see Methods; ref. 14) with an estimated false discovery rate (FDR) below 5% (Supplemental Figure 1, D and E). Within target transcripts (Supplemental Table 2), the resulting consensus read clusters mapped most frequently to introns (Figure 1A).

Identification of the RBM20 RRE

The diagnostic nucleotide transitions of PAR-CLIP experiments have been shown to occur at crosslinking sites and reflect the immediate positions of cellular RBP-RNA interactions (13). To elucidate the sequence preference of RBM20, we performed 5-mer counting in crosslink-centered regions (CCRs) derived from 4SU and 6SG experiments. The most abundant element was the UCUUUA 5-mer, followed by variations of this element, all containing a UCUU core (Figure 1B). These UCUU-containing 5-mers were most strongly enriched not only in the 4SU but also in the 6SG experiment, demonstrat-
and 6SG residues as expected (13). We performed EMSA using immunoprecipitated full-length epitope-tagged RBM20 to show that RBM20 binds an intronic RRE in the ryanodine receptor 2 (Ryr2) transcript (Figure 1E), validating the motif prediction. We compared different degrees of UCUU core mutations in Ryr2. We found that not only mutation of the entire core but also mutation of 2 core nucleotides led to a complete inhibition of RBM20 binding (Figure 1E), while a single nucleotide change within the core already led to a marked reduction in RBM20-RNA interaction (Supplemental Figure 2C).

Analysis of natural selection in RBM20-binding sites. To corroborate the functional significance of the RBM20 RRE in humans, we analyzed population genetics data for signatures of negative selection (17). Using SNP data from the 1000 Genomes Project (http://www.1000genomes.org/), we first analyzed the density of SNPs in RBM20 RREs or CLIP clusters in HEK293 cells. We found a significant decrease of SNPs/kb in clusters compared with introns and exons, with a further decrease in RREs compared with clusters in all 4 populations of the 1000 Genomes Project (Supplemental Figure 1D). Since this result might be affected by heterogeneity of mutation rates, we compared the distributions of derived

Figure 2. HITS-CLIP in rat cardiomyocytes. (A) Transcriptome-wide distribution of HITS-CLIP–derived RBM20 target sites in cardiomyocytes. The ratio of HITS-CLIP to cardiac RNA-seq percentage coverage in each region is displayed. (B) Binding motif enrichment in cardiomyocyte Rbm20 clusters. The log\_ frequencies of 5-mers in clusters correlated with frequencies of 5-mers in control sequence. (C) Sequence logo for the RBM20 RRE in cardiomyocytes was computed from the top 1000 intronic binding sites using MEME. (D) Confirmation of alternative splicing of genes with high ΔPSI values as determined by RNA-seq. It indicates isoforms expressed at higher levels in WT than in mutant rats and i2 those expressed at higher levels in mutant than in WT rats. Rbm20 was amplified as control and verifies the indicated Rbm20 genotype of the samples tested. (E) ΔPSI values > [0.01] of RBM20-regulated exons in WT versus Rbm20-deficient rat hearts. RBM20 acts predominantly as a splicing repressor in cardiomyocytes. (F) RBM20-regulated versus expected cardiac splicing events. MXE, mutually exclusive exon; CE, cassette exon; CNE, constitutive exon.
Table 1. Direct RBM20-regulated genes

<table>
<thead>
<tr>
<th>Gene ID</th>
<th>Gene symbol</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>ENSRNOG00000011589</td>
<td>Camk2d</td>
<td>Calcium/calmodulin-dependent protein kinase II delta</td>
</tr>
<tr>
<td>ENSRNOG00000012207</td>
<td>Dst</td>
<td>Dystonin</td>
</tr>
<tr>
<td>ENSRNOG0000003934</td>
<td>Enah</td>
<td>Enabled homolog (Drosophila)</td>
</tr>
<tr>
<td>ENSRNOG00000090979</td>
<td>Immt</td>
<td>Inner membrane protein, mitochondrial</td>
</tr>
<tr>
<td>ENSRNOG00000011343</td>
<td>Ldb3</td>
<td>LIM domain binding 3</td>
</tr>
<tr>
<td>ENSRNOG00000097072</td>
<td>Lmo7</td>
<td>LIM domain only protein 7</td>
</tr>
<tr>
<td>ENSRNOG00000055934</td>
<td>Mlcp</td>
<td>Muscular-enriched A-type laminin-interacting protein</td>
</tr>
<tr>
<td>ENSRNOG00000019892</td>
<td>Lxrp1</td>
<td>Leucine-rich repeat (in FLI) interacting protein 1</td>
</tr>
<tr>
<td>ENSRNOG00000016983</td>
<td>Myh7</td>
<td>Myosin heavy chain 7</td>
</tr>
<tr>
<td>ENSRNOG00000026700</td>
<td>Myom1</td>
<td>Myomesin 1</td>
</tr>
<tr>
<td>ENSRNOG00000025152</td>
<td>Nex</td>
<td>Nexilin</td>
</tr>
<tr>
<td>ENSRNOG00000022790</td>
<td>Olsn</td>
<td>Obscurin</td>
</tr>
<tr>
<td>ENSRNOG00000026558</td>
<td>Pallm3</td>
<td>PDZ and LIM domain 3</td>
</tr>
<tr>
<td>ENSRNOG00000004621</td>
<td>Rtn4</td>
<td>Reticulin 4</td>
</tr>
<tr>
<td>ENSRNOG00000017600</td>
<td>Ryp2</td>
<td>Ryanodine receptor 2</td>
</tr>
<tr>
<td>ENSRNOG000000115658</td>
<td>Sorbs1</td>
<td>Sorbin and SH3 domain containing 1</td>
</tr>
<tr>
<td>ENSRNOG00000033734</td>
<td>Tmnt2</td>
<td>Tropomin T type 2</td>
</tr>
<tr>
<td>ENSRNOG00000022637</td>
<td>Ttn</td>
<td>Titin</td>
</tr>
</tbody>
</table>

The listed direct targets show Rbm20-dependent differential splicing and are directly bound by RBM20.
with 138 proteins being frequently identified in multiple spliceosome preparations (20). In particular, we found all 7 Sm and nearly all U1 and U2 small nuclear ribonucleic particle (snRNP) proteins (except SNRPA, SNRPB2, SF3B5, and PHF5A) to precipitate with RBM20 (Figure 4C). In contrast, only few of the U4/U6 and U5 snRNP proteins were identified in the RBM20 pull-down. In addition, while we identified 45% of complex A–specific proteins, including U2AF65 and U2AF35, we found only 18% and 11% of complex B– or complex C–specific proteins, respectively (Supplemental Figure 4).

In particular, only 3 out of 20 proteins of the Prp19 complex and related proteins that are crucial during catalytic activation and join the spliceosome during complex B formation were found. Together, these data suggest that RBM20 interacts with U1 and U2 in early stages of spliceosome assembly, while it is not present in catalytically active complexes. It has been shown that the RS-rich region of for RBM20 binding did not affect reporter activity. This is further supported by mobility shift assays, in which we document that RNA containing the cluster-binding site downstream of PEVK8 interacted with the RRM domain of RBM20 (RNA I), while RNA containing UCUU elements that do not correspond to the cluster (RNA II and III) did not bind to RBM20 (Figure 3E).

**Figure 3. RBM20 binds to intronic splicing silencers upstream and downstream of repressed exons.** (A–C) ΔPSIs are shown as black lines (WT compared with homozygous rats). Elevated versus decreased ΔPSIs show inclusion or exclusion of exons, respectively. Red ticks indicate mapped RBM20 cluster positions. (A) Ttn is predominantly spliced in the elastic region. Mapped RBM20-binding sites coincide with differential splicing. (B) Camk2d exons 14–16 are mutually exclusive and regulated by RBM20. Mapped RBM20-binding sites are located upstream of exon 14 and downstream of exon 15. (C) Ldb3 undergoes an exon switch affecting exon 4 versus exons 5–7. Mapped binding sites are upstream of exon 5 and downstream of exon 8. (D) Mean CLIP density near RBM20 activated (red) and repressed exons (blue). Dotted line indicates density around exons not regulated by RBM20. Gray shows 90% CIs. (E) Right panel: specific activity of RBM20 on Ttn RNA from the PEVK region in a splice-reporter assay. Rbm20 expression leads to exclusion of the firefly luciferase–containing (Fluc) exon. Ratio of Fluc to Renilla luciferase (Rluc) activity (downstream exon) reflects splice activity. Reporter activity depends on mutation of 2 UCUU elements in cluster CID016226, but not outside the cluster. Mutations are indicated as X. White boxes only show native reporter. Controls: PEVK-construct cotransfected with empty vector (–). Rbm20 cotransfected with a Ttn M-band region construct not affected by RBM20 (M). ***P < 0.001 compared with native reporter. Left: EMSA evaluating RBM20 binding to Ttn PEVK-derived RNAs. White arrows indicate input RNA; black arrows indicate RNA-protein complex.
RBM20 is a hot spot for mutations that cause DCM. We therefore determined the impact of a previously reported RBM20 S635A mutation located in the RS-rich region (5) on protein binding by comparing proteins precipitating with WT RBM20 versus S635A-RBM20. The mutation did not abrogate interactions with core spliceosomal proteins, but for 38 of the alternative spliceosomal proteins associated with WT RBM20, interactions were significantly reduced when the RS-rich region of RBM20 was mutated (Figure 5A). To distinguish RNase insensitive interactions indicating binders not dependent on RNA as a connector, we repeated the experiment with an RNase digestion step after immunoprecipitation and compared the results with the nondigested data set. This revealed 13 of the WT-specific spliceosomal protein interactions as RNase insensitive (Figure 5B). Using a label-free quantification, 10 of these RNase insensitive WT-specific proteins identified in HEK cells were also found to precipitate with endogenous-expressed RBM20 in cardiomyocytes (Figure 5B). Overall, the set of interaction partners in cardiomyocytes encompassed the majority of splicing factors identified in the SILAC experiment, including U2AF65 and U2AF35, but only 1 protein of the Prp19 complex (Supplemental Table 7), corroborating our findings from HEK cells in cardiomyocytes.

**Figure 4. Identification of RBM20 protein-protein interaction partners.** (A) q-AP-MS using SILAC. Differentially SILAC-labeled HEK293 cells (light, medium, and heavy) were transfected with FLAG-tagged control vector, FLAG-tagged RBM20 WT, or S635A mutant. RBM20 protein complexes were enriched by affinity purification and analyzed by quantitative mass spectrometry. (B) The log_2 ratios of protein-protein interactions for WT-RBM20 versus control are shown on the left. Red dots indicate known splicing factors. Validation of q-AP-MS hits by Western blotting is shown on the right. (C) U1-, U2-, and U4/U6.U5-specific proteins are grouped according to snRNP association and highlighted in blue when identified as interacting with RBM20.

**Regulatory impact of RBM20 on cardiomyopathy-associated and novel candidate genes.** NCBI Medical Subject Heading (MeSH) analyses of all 18 directly RBM20-regulated targets revealed that the set was significantly enriched for the heading “dilated cardiomyopathy” ($P = 1.33 \times 10^{-13}$) (Supplemental Table 8). Further significantly enriched disease-related MeSH terms were “hypertrophic cardiomyopathy” ($P = 1.8 \times 10^{-09}$) and “heart failure” ($P = 8.48 \times 10^{-05}$). The identification of direct RBM20 target genes and their regulated exons allowed us to specifically test their relevance in a large cohort of heart failure patients, including a single patient that we reported previously with a heterozygous S635A mutation in RBM20 (5).

We compared the rat cardiac-splicing patterns of direct targets to those of the RBM20 S635A individual and 5 control subjects with DCM, but without any nonsynonymous changes in their RBM20-coding sequence (Supplemental Table 9). This analysis revealed conserved differential splicing of orthologous direct targets at the exon level for Camk2d, Ldb3, LIM domain only protein 7 (Lmo7), PDZ and LIM domain 3 (Pdlim3), reticulin 4 (Rtn4), Ryr2, and Ttn (Figure 6 and Supplemental Figure 5). As expected from our analysis, the differentially spliced exons were flanked
Figure 5. Impact of RBM20 S635A mutation on protein-protein interaction. (A) WT versus S635A comparison. RBM20 was identified as highly and equally abundant in WT- and S635A-RBM20 samples, indicating successful immunoprecipitation. Red circles indicate spliceosomal interaction partners not affected by the mutation. Labeled red dots indicate spliceosomal interaction partners specific to WT- (log 2 ratio > 0.4) or S635A-RBM20 (log 2 ratio < -0.4). (B) Comparison of RNase-digested versus nondigested q-AP-MS identifies RNase-insensitive RBM20 interactors. Upper right quadrant contains WT- and lower left quadrant mutant-specific interactors enriched in RNase-digested and nondigested samples. Spliceosomal interaction partners are marked by red dots and labeled when WT- or mutant-specific in both protein-protein interaction screens. Bold labeled proteins were identified as also interacting with RBM20 in cardiomyocytes.
patients with end-stage heart failure, we observed that cardiac RBM20 expression greatly varies among individuals (Supplemental Figure 6). We compared the splicing patterns of RBM20-regulated exons in 10 individuals, each with the highest versus lowest RBM20 expression (Figure 7 and Supplemental Table 9). The difference in endogenous RBM20 expression (high vs. low) significantly affected the splicing of orthologous exons for CAMK2d, LDB3, RYR2, and TTN and mirrors our observations from the S635A mutant patient and Rbm20-deficient rats. Given the fundamental role of these targets for proper heart function, this may suggest that endogenous RBM20 expression levels are important for the modulation of cardiac function.

Moreover, we tested the impact of differential expression levels of RBM20 on the splicing of its target exons in cardiac tissue from a human heart-failure cohort. In heart tissue from 148 patients with end-stage heart failure, we observed that cardiac RBM20 expression greatly varies among individuals (Supplemental Figure 6). We compared the splicing patterns of RBM20-regulated exons in 10 individuals, each with the highest versus lowest RBM20 expression (Figure 7 and Supplemental Table 9). The difference in endogenous RBM20 expression (high vs. low) significantly affected the splicing of orthologous exons for CAMK2d, LDB3, RYR2, and TTN and mirrors our observations from the S635A mutant patient and Rbm20-deficient rats. Given the fundamental role of these targets for proper heart function, this may suggest that endogenous RBM20 expression levels are important for the modulation of cardiac function.
The combination of the splicing analysis and determination of direct RNA-binding sites in the heart revealed several directly regulated RBM20 targets unknown to date. We discovered DCM-related and several other genes implicated in cardiac disorders and validated representatives in human heart failure patients. Interestingly, the identified targets allow further insights and reveal new RBM20-related disease mechanisms: RYR2 is found primarily in cardiac muscle and is a component of the major Ca\(^{2+}\) release channel for excitation-contraction coupling in the sarcoplasmic reticulum membrane. A 24-bp exon in RYR2 is upregulated in Rbm20-deficient rats and human cardiomyopathy patients (Figure 6A). This exon is usually present at low levels in hearts and shifts RYR2 from the ER to the intranuclear Golgi apparatus, with profound effects on intracellular Ca\(^{2+}\) signaling (27). Mutations in N-terminal domains of RYR2 have been associated with arrhythmogenic right ventricular tachycardia and sudden death (28). Cardiac arrhythmia and sudden death are both signs of profound effects on intracellular Ca\(^{2+}\) signaling (27). Mutations in N-terminal domains of RYR2 have been associated with arrhythmogenic right ventricular tachycardia and sudden death (28). Cardiac arrhythmia and sudden death are both signs of...
function may be associated not only with cardiac arrhythmias, but also cardiomyopathies (29, 30). Here, we were able to show that the 24-bp exon is also regulated in human heart failure patients with high and low RBM20 expression.

LMO7 is a transcription factor that previously has not been associated with DCM. In LMO7, RBM20 deficiency resulted in retention of exons 9 and 10 across species (Figure 6B). These exons encode an F-box domain present only in a brain-specific isoform of LMO7 (31). Its function in regulating expression of genes that are important for skeletal muscle and heart function (32) makes it an interesting modulator in RBM20-related DCM.

RTN4 protein, also known as neurite outgrowth inhibitor (NOGO), localizes predominantly in the endoplasmic reticulum. It is well studied in the nervous system, but its function in the heart remains largely unknown. In RBM20-mutant rats and human, exons 3 and 4 of RTN4 are retained (Figure 6C). This isoform is most strongly expressed in the brain and only weakly in the heart (33). It is increased in left ventricular tissue from DCM and ischemic hearts and has been proposed as a potential indicator of heart failure (34–36). Here, we demonstrate that RBM20 represses the predominantly neuronal isoform in favor of the heart-specific exon usage.

PDLIM3 binds α-actinin 2 in the Z-discs of myofibers. Two different major isoforms of PDLIM3 are expressed in heart and skeletal muscle tissue that include either exon 4 or exons 5 and 6. The predominant isoform in the heart includes exon 4 (37). In RBM20-deficient rats and human, the ratio of these 2 isoforms is shifted toward the skeletal muscle form (Figure 6D). Within exon 4, a Glu106Ala mutation has been linked to hypertrophic cardiomyopathy, indicating the importance of the sequence encoded by this exon for proper heart functioning (37).

Considering the fundamental role of RBM20 for proper splicing of cardiac transcripts, it was unexpected that its endogenous expression levels in heart failure patients varied drastically. On the other hand, this could also be an indication that differences in RBM20 expression levels contribute to the modulation of cardiac function under pathophysiological conditions. With the set of direct RBM20 target genes defined in this study, it was possible to investigate the impact of RBM20 expression on differential exon usage in its targets in cardiac tissue from end-stage heart failure patients. Despite the heterogeneous nature of the human heart failure samples, we could demonstrate the regulation of differential exon usage in 4 direct RBM20 target genes in 10 patients with high versus 10 patients with low RBM20 gene expression levels (CAMK2D, LDB3, RYR2, TTN). These 4 genes showed the strongest differences in RBM20-regulated exon usage across species. It is therefore conceivable that we have missed other targets with smaller effects in our analysis due to the heterogeneous nature of the patient samples. The validity of our findings, on the other hand, is supported by the fact that the same exons within the 4 genes were also regulated in a patient that carried an RBM20 nonsynonymous mutation (S635A) in the SR-rich region. Given the suggested impact of RBM20 expression levels on exon usage of its direct targets, we paid careful attention to comparing the S635A patient to human cardiomyopathy controls that were matched in their expression levels but did not have any nonsynonymous variants in their RBM20-coding sequence.

In our study, we describe the function of RBM20 as a molecular switch that orchestrates isoform transitions of a network of genes with essential cardiac functions. Not only genetic variation, but also the modulation of RBM20 gene expression levels, is operative in the regulation of this network of direct targets in the human heart. This is of interest because the modulation of RBM20 levels may be an attractive target for therapeutic intervention. Moreover, the abundance of RBM20 in heart tissue could play a role in the pathogenesis of more common forms of heart failure irrespective of the RBM20 genotype. The clinical significance of splicing controlled by RBM20 expression levels during the pathogenesis of common forms of heart failure will be an interesting avenue to explore in future studies.

In summary, our analyses tie together RBM20-regulated direct targets, the underlying mechanism of splicing repression, and molecular consequences in clinical samples. Collectively this provides insights that are extensible into the pathophysiology of human cardiac function.

Methods

PAR-CLIP. HEK293 cells with tetracycline-inducible expression of FLAG/HA-tagged RBM20 have been described previously (5). For PAR-CLIP, we essentially followed the protocol previously described (13). The libraries were sequenced on an Illumina Genome Analyzer GAI1 using the 50-bp single-read protocol.

HITS-CLIP. Neonatal rat ventricular myocytes (NRVMs) were isolated from SD rats using a modification of the Kasten’s technique, as described (38). At 48 hours after isolation, cardiomyocytes were irradiated 1 time at 400 mJ/cm and an additional 200 mJ/cm with 254 nm UV light and lysed in NP40 lysis buffer. After DNase and RNase T1 treatment, the lysate was immunoprecipitated using RBM20 antibody (5) covalently coupled to Protein G Dynabeads (Invitrogen). Radiolabeling of immunoprecipitated RNA, protein-RNA complex separation, sequencing library preparation, and sequencing were done following the PAR-CLIP protocol.

Computational analysis of PAR-CLIP and HITS-CLIP data. We used a computational pipeline that performed all steps of the analysis from raw reads to cluster sets and target genes, essentially as previously described (14), with emphasis on stringent filtering and controlling the false-positive rate in the identification of binding sites (Supplemental Methods).

Motif analysis. Word counting was performed on CCRs for PAR-CLIP and complete read clusters for HITS-CLIP. The log, word enrichment was computed for intronic-binding sites as the log ratio of the relative frequency of each 5-mer to the background frequency in RefSeq introns. Words with less than 10 occurrences inside binding sites were discarded from the analysis.

Within PAR-CLIP CCRs, we searched for the UCUU that was closest to the anchor position and selected it as the new center. CCRs without UCUU were discarded. The remaining motif-centered regions (MCRs) were used to build an average profile of nucleotide conversion events, indicative of crosslinking, around the motif.

Human studies. We obtained cardiac tissue from 147 subjects with end-stage heart failure. All subjects were transplanted at the Royal Brompton and Harefield NHS Foundation Trust (London, United Kingdom). On this occasion, cardiac samples were obtained and immediately frozen in liquid nitrogen. The S635A patient was described earlier (5).
RNA-seq and splicing analyses. Total RNA was extracted using TRIzol reagent and RNA-seq libraries prepared with the TruSeq RNA Sample Preparation Kit (Illumina). Barcoded CDNA fragments of poly(A)+ RNA were then sequenced on a HiSeq 2000 instrument from Illumina with 2 × 100 bp PE chemistry. Reads were mapped against the reference genome GRCh37 using TopHat version 1.3.1 (39) supplied with the gene annotation of the Ensembl database (40). We used DEXSeq (18) to test all exons for differential usage and to reanalyze previously published RNA-seq data of Rbm20-deficient mice. We only counted the start positions of reads, and the average of control sets of the same size that were randomly sampled from each sample applies a generalized additive model with integrated smoothness estimation. We smoothed the coverage profiles using the ggplot2 R package, which is available as described in Supplemental Methods. Counts across samples were normalized using a quantile-based scaling method (41). Gene expression levels were adjusted for available covariates as described in Supplemental Methods.

Gene expression analysis in the heart failure cohort. Gene expression in the heart failure cohort individuals was measured using RNA-seq. For each protein coding gene from the NCBI annotation, version 66, the overlapping TopHat aligned reads were counted using the HTSeq package. Counts across samples were normalized using a quantile-based scaling method (41). Gene expression levels were adjusted for available covariates as described in Supplemental Methods.

EMSA. For EMSA, FLAG/HA-tagged full-length RBM20 or GST-tagged Rbm20 RR domain was used (see Supplemental Methods). γ-32P-labeled RNA was incubated with immunoprecipitated RBM20 protein and 100 ng tRNA in 20 μl binding buffer (10 mM HEPES-KOH, pH 7.9, 50 mM KCl, 3 mM MgCl₂, 1 mM DTT, and 5% glycerol). The resulting complexes were resolved on native 5% acrylamide gels run at 4°C at 200 V for 2 hours in 0.5× TBE and exposed using a PhosphorImager.

Enrichment analyses. To assess functional and disease-related association of RBM20 splice substrates, enrichment analysis was performed with EGAN (42). To estimate the nature of transcript-processing events leading to differential usage of Rbm20-regulated exons, overlapping transcript events as annotated by Ensembl were assigned to exonic regions. The significant enrichments of GO terms and MeSH and those for certain event types of exons in direct Rbm20 targets when compared with all exons expressed in the heart were evaluated with the hypergeometric test, and significant enrichments were adjusted for multiple testing using Bonferroni’s method.

Construction of RNA maps. We determined the coverage profiles of HITS-CLIP reads for all differential used exons as defined above. We split the exons into activated (ΔPSI > 0.01) or repressed (ΔPSI < −0.01) exons and compared the coverage profiles of both sets with the read coverage of control sets of the same size that were randomly sampled from nondifferential exons. We only counted the start positions of reads, and duplicated identical reads were only counted once. We investigated regions from 400 bp upstream of the 5′ splice site to 50 bp into the exon and from 50 bp upstream of the 5′ splice site to 400 bp into the downstream intron of differentially used exons and their adjacent neighbors. We smoothed the coverage profiles using the ggplot2 R package, which applies a generalized additive model with integrated smoothness estimation and plots the smoothed coverage along with its 95% CI.

Splice reporter assay. We quantified RMB20 activity in a dual luciferase splice reporter assay (5) using a mini-gene representing titin PEVK exons 4 through 13 and a RNA destabilization motif for increased signal amplitude. Several mutations were introduced into the reporter construct, as described in Supplemental Methods. HEK293 cells (2500 cells/well on 96-well plates) were transfected with the reporter-plasmids and CMV-driven RBM20 or control plasmid (pcDNA3.1) using Polyethylenimine 40 kDa (Polysciences Europe GmbH). At 72 hours after transfection, reporter activity was measured using the Dual-Luciferase Reporter Assay System (Promega), following the manufacturer’s instructions, and an infinite M200 Pro Plate Reader (TECAN). Ratios of firefly to Renilla luciferase activity were normalized to the RBM20-negative control. Results were confirmed in 3 independent experiments. Based on the half-life of the resulting protein, we found higher sensitivity of the RNA-based splice analysis as compared with the luciferase readout (Supplemental Figure 3, E and F).

SILAC experiments. SILAC labeling of HEK293T cells, immunoprecipitation, and LC-MS/MS analysis were essentially performed as previously described (16) (for details see Supplemental Methods). Label swap experiments were performed to filter out contaminants. Experiments were performed in quadruplicate. Biological replicates were grouped and enriched proteins determined by 1-sample t test (ratio cut off: 1.3).

Accession numbers. PAR-CLIP, HITS-CLIP, HEK transcriptome, and human cardiac transcriptome sequencing data are available at the Sequencing Archive of the European Nucleotide Archive (ERP001301 and E-MTAB-2572).

Statistics. Data are presented as mean ± SD unless otherwise indicated. Differences between experimental groups were analyzed by 2-tailed Student’s t test unless otherwise specified. P < 0.05 was considered statistically significant.

Study approval. All human studies were approved by the institutional review board of the Cardiovascular Biomedical Research Unit, Royal Brompton and Harefield NHS Trust, and individuals gave informed consent. All experiments involving animals were performed in accordance with institutional and NIH guidelines and approved by the Landesamtes für Gesundheit und Soziales (LaGeSo), Berlin, Germany.

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