Rbm20 regulates titin alternative splicing as a splicing repressor

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Received August 29, 2012; Revised November 20, 2012; Accepted December 7, 2012

ABSTRACT

Titin, a sarcomeric protein expressed primarily in striated muscles, is responsible for maintaining the structure and biomechanical properties of muscle cells. Cardiac titin undergoes developmental size reduction from 3.7 megadaltons in neonates to primarily 2.97 megadaltons in the adult. This size reduction results from gradually increased exon skipping between exons 50 and 219 of titin mRNA. Our previous study reported that Rbm20 is the splicing factor responsible for this process. In this work, we investigated its molecular mechanism. We demonstrate that Rbm20 mediates exon skipping by binding to titin pre-mRNA to repress the splicing of some regions; the exons/introns in these Rbm20-repressed regions are ultimately skipped. Rbm20 was also found to mediate intron retention and exon shuffling. The two Rbm20 speckles found in nuclei from muscle tissues were identified as aggregates of Rbm20 protein on the partially processed titin pre-mRNAs. Cooperative repression and alternative 3’ splice site selection were found to be used by Rbm20 to skip different subsets of titin exons, and the splicing pathway selected depended on the ratio of Rbm20 to other splicing factors that vary with tissue type and developmental age.

INTRODUCTION

Titin, the gene containing the largest number of exons (363 exons in human), encodes the largest polypeptide in nature [2.97–3.7 megadaltons (MDa)] (1,2). The titin molecule is elastic, with a size ~1 μm long and 3–4 nm wide (3–5). One molecule spans half of the sarcomere with the amino terminus located in Z-line and the carboxyl terminus in the M-line (2,6–8). The elasticity of titin mainly comes from the folding and extending of polymeric immunoglobulin regions (middle Ig) and the PEVK region [rich in proline (P), glutamate (E), valine (V) and lysine (K)] (2,9). The giant size and the specialized structure enables titin to play a mechanical role in maintaining sarcomere length and structure integrity: it accounts for most of the passive tension of striated muscles in the physiological extension range to restore the sarcomere to normal length after stretch and reposition the thick and thin filaments (5,10–13). Besides its mechanical function, titin also plays important roles in many other physiological processes. Titin acts as a scaffold for myofibrillar assembly during muscle development, and it interacts with many structural proteins (14–17).

Titin undergoes developmental isoform transition from large to small in both cardiac and skeletal muscles (1,2,18–21). Diverse titin isoforms result from the alternative splicing of titin mRNA in the regions corresponding to the middle Ig (exons 50–96) and PEVK regions (exons 115–225) (1,2,9). In heart, the titin-based passive tension determines the stiffness of the myocardial wall during ventricular filling, so it is important to maintain the proper isoform ratios; abnormal titin isoform expression has been associated with heart disease (22–27). Many classes of titin isoforms (such as N2A, N2B and N2BA) and their splicing pathways have been characterized (2,9,18,21), but the mechanism underlying these splicing pathways remains unknown.

We found a mutant rat lacking in titin alternative splicing (19,21,28) and identified the mutation as a nearly complete deletion of the Rbm20 gene (29). Rbm20 protein is a putative RNA-binding protein with one RNA recognition motif and one arginine-serine rich domain. Thus far, mechanistic studies on RBM20 are lacking. Only a few articles have reported mutations in the human RBM20 gene, and these were associated with human dilated cardiomyopathy (DCM), with cell function descriptions lacking (30–34). Our previous study found that the mutant rat with Rbm20 deficiency had a similar pathological phenotype as found in...
human DCM, and we also found titin splicing was altered in a human DCM subject with an RBM20 mutation. These observations suggest that the deficiency in Rbm20-regulated titin alternative splicing may be an underlying cause for DCM (29). The current work reports investigations on the mechanism of Rbm20 in regulating titin alternative splicing. We demonstrate that Rbm20 mediates intron retention, exon skipping and exon shuffling of titin mRNA, forms microscopically identified aggregates with partially processed titin pre-mRNAs in the nucleus and uses different splicing pathways to skip different subsets of titin exons to form different isoforms. We found muscle cells use a relative simple system, by controlling the Rbm20/splicing factors ratio expression level, to switch splicing pathways and regulate the extremely complex titin alternative splicing process.

MATERIALS AND METHODS

RT–PCR analysis

RNA was purified from the indicated tissues with TRIzol regent (Invitrogen, 15596026) and further treated with RQ1 RNase-free DNase (Promega) to remove genomic DNA contamination. One microgram of total RNA was reverse transcribed with ImProm-II Reverse Transcription System (Promega) using Random primers (Promega). The RT reaction was used as template for PCR to characterize intron retention, exon skipping and exon shuffling in titin mRNA. The primers are listed in Supplementary Table S1. The PCR products were analyzed on ethidium bromide agarose gels. The bands of interest were gel purified and sequenced to confirm their identity.

Antibodies

Polyclonal antibodies directed against full-length Rbm20 were produced from chicken and rabbit and affinity purified (29). For immunohistochemistry, the concentration of chicken anti-Rbm20 was adjusted to 1 mg/ml and used at 1:500 dilution, and rabbit polyclonal anti-SP1C1 (sc-84117, Santa Cruz), mouse monoclonal anti-SC-35 (S4045, Sigma) and mouse monoclonal anti-fibrillarin (ab45661, Abcam) were used at 1:200 dilution. The secondary antibody for chicken anti-Rbm20 was Alexa Fluor 594 goat anti-chicken IgG (Invitrogen), used at 1:200 dilution. The secondary antibodies for SP1C1, SC-35 and fibrillarin were Alexa Fluor 488–conjugated goat anti-rabbit and goat anti-mouse IgG (Invitrogen), used at 1:200 dilution. For western blots, the concentration of rabbit anti-Rbm20 was adjusted to 1 mg/ml and used at 1:5000 dilution, and mouse monoclonal anti-SFRS1 (sc-73026, Santa Cruz), rabbit polyclonal anti-U2AF65 (sc-48804, Santa Cruz) and rabbit polyclonal anti-hnRNPL (sc-28726, Santa Cruz) were used at 1:3000 dilution. The secondary antibodies for western blots were horseradish peroxidase–conjugated ECL donkey anti-rabbit IgG (NA934V, GE Healthcare) and horseradish peroxidase–conjugated ECL sheep anti-mouse IgG (NA931V, GE Healthcare) used at 1:5000 dilution.

Immunofluorescence and mRNA fluorescence

in situ hybridization

The procedure for immunohistochemistry in HL-1 cells is described previously (29). For the immunohistochemistry in samples from living tissue, homogenized muscle was fixed in fresh 4% formaldehyde in phosphate-buffered saline (PBS) for 15 min, spread on a cover slip and allowed to attach by air drying. The tissues were then permeabilized for 10 min in ice-cold PBS with 0.1% Triton X-100. After three washes in PBS, the homogenized tissues attached on cover slips were immunostained using the same procedures as used for staining HL-1 cell. The procedures for combined immunofluorescence and mRNA fluorescence in situ hybridization (FISH) were described previously (35). Titin exons 1–50 were amplified by four pairs of primers listed in Supplementary Table S1 and cloned into four pGEM-T easy vectors, respectively (Promega). The biotin-labeled cDNA probe against titin exons 1–50 was prepared by nick translation (Roche, 11745824910) using 0.5 μg of each of these plasmids. Chicken anti-Rbm20 and Alexa Fluor 594 goat anti-chicken IgG were used to immunostain Rbm20; fluorescein isothiocyanate–conjugated streptavidin (Sigma, S3762) was used to detect biotin-labeled cDNA probe hybridized with titin mRNA.

RNA immunoprecipitation

The RNA immunoprecipitation (RIP) procedure described previously (36) was followed with some modifications. Nuclei from a 150-day-old wild-type (Wt) rat left ventricle (LV) were enriched first (37); the nuclei were suspended in 1 ml of FA lysis buffer containing 1 mM ethylenediaminetetraacetic acid, 50 mM (4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid) (HEPES)–KOH (pH 7.5), 140 mM NaCl, 0.1% sodium deoxycholate (w/v), 1% Triton X-100 (v/v) and 1× protease inhibitor cocktail (Sigma, P8340). The lysis buffer is supplemented with 400 U/ml RNAsin inhibitor (Promega, N2511), 10 mM vanadyl ribonucleoside complex (New England Biolabs, S1402S), 10 U/ml RNase-free DNase (Promega, M6101) and 0.5 μg/ml yeast tRNA (Invitrogen, 15401) during RIP. Cross-linking protein with the RNA is not necessary in this study. Sonication was adopted to break down the nuclei and the ribonucleoprotein complex. After centrifugation at 15 000g for 10 min at 4°C, the supernatant was collected. The supernatant was first pre-cleaned by incubation with 200 μl of Dynabeads® Protein G (Invitrogen, 10004D) for 30 min at room temperature and then incubation with antibody-coated Dynabeads® Protein G for 1 h at room temperature. After incubation, the supernatant was discarded, and the beads were washed five times with FA lysis buffer containing 10 mM vanadyl ribonucleoside complex and submitted for protein extraction and RNA extraction. The precipitated protein was extracted with urea–thioura sodium dodecyl sulphate buffer as described previously (38). The precipitated RNA was extracted with TRIzol reagent. To precipitate Rbm20 protein and the associated RNA, 8 μg of rabbit anti-Rbm20 coated on 200 μl of Dynabeads® Protein G was
used. As a parallel control, 8 μg of rabbit anti-spectrin α coated on 200 μl of Dynabeads® Protein G was used.

**Plasmid construction**

Full-length Rbm20 was in-frame cloned into pcMS-EGFP vector (Clontech) using SalI and NotI restriction sites. MS2–Rbm20 fusion construct was made by inserting MS2 coat sequence before Rbm20 in PCMS-EGFP using the restriction sites NheI and MluI. The sequence coding MS2 coat protein was amplified from the pMAL-MS2-MBP construct (provided by Zhaolan Zhou, University of Pennsylvania). Titin minigene 64–70, which spans from −120 bp upstream of titin exon 64 to 75 bp downstream of titin exon 70, was cloned into PCDNA 3 vector (Invitrogen) using the restriction sites EcoRV and XhoI. The fragment containing three MS2 hairpins was amplified from the pCR2.1 MS2 construct (provided by Zhaolan Zhou, University of Pennsylvania). MS2–exons 64–70 construct was made by inserting MS2 stem–loop sequence before exon 64 of the titin minigenic in pcDNA 3 using restriction sites BamHI and EcoRI. MS2–exons 64–70 construct was made by inserting MS2 stem–loop sequence after exon 70 of the titin minigene in pcDNA 3 using restriction sites XhoI and ApaI. Rbm20 without MS2 coat protein is cloned into PCMS-EGFP, and titin minigene expressing exons 25–27, 35–37, 52–58, 56–62, 60–66 and 64–70 without MS2 stem–loop sequence is constructed into PCDNA 3 vector (for primers, see Supplementary Table S1). All expression plasmid and minigene structures were confirmed by sequencing.

**Mammalian cell culture, transfection and transcription inhibition**

HL-1 cells were cultured on fibronectin-coated coverslips as previously described (29). Transcription in HL-1 cells was inhibited by adding 1 μg/ml of actinomycin D to the medium for a period of 2.5 h before immunofluorescence staining. HeLa cells were cultured in minimum essential medium (Invitrogen, 11095080) in standard conditions. HeLa cells (10^5) were plated in each well of 24-well plates and grown for 24 h before transfection. Cells are co-transfected with 1 μg of MS2-Rbm20-pcMS-EGFP and 0.2 μg of each plasmid expressing titin minigene using FuGENE HD transfection reagent (Promega, E2311). An equivalent amount of empty pcMS-EGFP was co-transfected with titin minigenes to serve as a parallel control.

**Titin isoform characterization**

Titin isoforms were analyzed using the vertical sodium dodecyl sulfate–agarose gel electrophoresis system (38).

**RESULTS**

**Rbm20 represses titin mRNA splicing**

Our previous study showed that Rbm20 is required for the alternative splicing of titin mRNA. In Wt (Rbm20+/+/) rats, the cardiac titin mRNA undergoes extensive exon skipping within the regions corresponding to the middle Ig (from exons 50 to 96) and the PEVK (from exons 115 to 225) regions. These exon-skipping events are inhibited in homozygous mutant (Hm, Rbm20−/−) rats where 13 of the 14 coding exons of Rbm20 gene are deleted (29). Theoretically, if RT-PCR is used to amplify the cardiac titin mRNA for the middle Ig and PEVK regions, we should expect smaller PCR products in the Wt than in the Hm. This is true when the primer pairs are designed to span large number of exons (e.g. >10 exons) (21). However, when using primers spanning just two or three exons, we amplified larger PCR products in the Wt than in the Hm for many primer pairs (Figure 1 and Supplementary Figure S1A–E). Sequencing these larger PCR products revealed retained intronic sequences in titin mRNA. Intron retention in Wt titin mRNA occurs between exons 50 and 70, 80 and 88, 117 and 118, 120 and 122, 124 and 129, 130 and 136, 138 and 145 and 146 and 218, but it never occurs in the Hm. A no-RT control was used to exclude the possibility of genomic contamination (Supplementary Figure S1F). The fact that introns cannot be spliced out in the Wt but can be normally removed in the Hm directly suggests that Rbm20 functions to repress the splicing of titin mRNA.

This repression varies in extent along the titin mRNA. For the region between exons 50 and 70, splicing patterns corresponding to retained introns and constitutive splicing co-exist, but the retained intron pattern dominates (Figure 1). This means the splicing between exons 50 and 70 is repressed in the majority of titin mRNAs. The pattern is reversed for the region between exons 80 and 88 so that retained intron and constitutive splicing patterns co-exist but the constitutive splicing pattern dominates (Figure 1), indicating the splicing is repressed only in the minority of titin mRNAs. The retained intron is the only splicing pattern detected from most of the primer pairs for the PEVK region between exons 124 and 218 (Figure 1). This demonstrates the splicing in much of the PEVK region is completely inhibited in all titin mRNAs.

**Rbm20 repression is associated with exon skipping and exon shuffling**

The intron retention pattern along the cardiac titin mRNA in Wt rats is summarized in Figure 2A. The middle Ig and PEVK regions of titin mRNA (exons 50–225) can be further divided into regions repressed by Rbm20 and regions not repressed. We then investigated whether the repression of Rbm20 on splicing is related to titin exon skipping. The exon skipping in Wt is repressed in the majority of titin mRNAs. The retained intron is the only splicing pattern detected from most of the primer pairs for the PEVK region between exons 124 and 218 (Figure 1). This demonstrates the splicing in much of the PEVK region is completely inhibited in all titin mRNAs.
Figure 1. Splicing of titin mRNA is inhibited in the wild type (Wt, Rbm20^+/+) but not in the homozygote mutant (Hm, Rbm20^-/-). RT-PCR was used with primer pairs spanning two or three exons to amplify cardiac titin mRNA from Wt and Hm left ventricle (LV). Usually three bands were obtained in the Wt for the middle Ig region between exons 50 and 70 and exons 80 and 88, but only one band in the Hm. As exemplified by exons 68–70 in the top panel, the lower Wt band has the same size as the adjoining Hm band and is the product of constitutive splicing; the upper band contains product where two introns for the primer-spanned region are both retained; the middle band has one of the two spanned introns retained. The splicing in the PEVK region (exons 124–218) is repressed to a greater extent in the Wt. Under most circumstances (as exemplified by exons 124–126), PEVK introns cannot be removed from Wt titin mRNA so that only one band, representing the retained intron pattern with a larger size than the adjoining band in the Hm, can be detected for most primer pairs. No retained introns can be detected in the Hm. The size of marker is indicated on the left, and the expected size of the PCR product for each primer pairs are listed in the Supplementary Table S2. The PCR products match their predicted size for all the primer pairs.
correspondingly, no exon skipping occurs in these regions (Figure 2B). Our results suggest that the exon skipping of titin mRNA is closely related with the repression of Rbm20 on splicing (Supplementary Figure S2G). No intron retention can be detected where Rbm20 is absent, and correspondingly all titin exons between exons 50 and 225 are constitutively spliced in the Hm (21).

Exon shuffling was found with cardiac titin mRNA in Wt rats. When we amplified titin mRNA using a forward primer set on exon 137 and reverse primer set on multiple upstream exons ranging from exon 24 to exon 96, multiple PCR products were obtained (Supplementary Figure S3A). Sequencing these PCR products revealed exon shuffling in titin mRNA: downstream exons were spliced before upstream exons. We then tested many primer combinations and identified exons 84, 116, 124, 138, 146 and 175 as additional donor exons and exons 70, 82, 87, 88, 124, 129 and 136 as acceptor exons for exon shuffling (Figure 2C). Each donor exon is able to splice with multiple acceptor exons and vice versa, demonstrating that the splicing between downstream donor exons and upstream acceptor exons was somewhat random. Exon shuffling is significantly inhibited in the Hm (Supplementary Figure S3B), indicating this event is also regulated by...
Rbm20. Exon shuffling can barely be detected from polyadenylated titin mRNA; this means most of them are not from mature titin mRNA (Supplementary Figure S3C). The exon shuffling with titin mRNA is conserved between species. Many exon-shuffling events were detected with human cardiac titin mRNA, and human exons mostly orthologous to those in rat mediated exon shuffling (Figure 2C). Exon shuffling in rat and human titin mRNA is detailed in Supplementary Figure S4.

Nuclear Rbm20 speckles are titin pre-mRNA processing sites

Immunofluorescence staining always revealed two Rbm20 speckles per nucleus with tissue obtained from living muscles (Supplementary Figure S5A). Co-staining of Rbm20 speckles with a nucleolus marker (fibrillarin) showed that Rbm20 speckles have a size similar to nucleoli (0.6 μm in diameter), but the two are not co-localized (Figure 3A). Rbm20 was not co-localized with other nuclear bodies (Supplementary Figure S5B). We suspected that the Rbm20 speckles are aggregates of Rbm20 proteins on titin mRNAs. Immunofluorescence combined with RNA FISH demonstrated that titin mRNAs co-localized with the Rbm20 speckles in nuclei from in vivo muscle (Figure 3B). The two Rbm20/titin mRNA speckles are possibly formed by recruiting Rbm20 protein onto the newly transcribed titin mRNAs near the two titin transcription loci on sister chromosome 3s. Based on this hypothesis, the formation of Rbm20 speckles should depend on transcription. As expected, the Rbm20 speckles dissociated after transcription is inhibited (Figure 3C), indicating that the Rbm20/titin mRNA speckles are formed on newly synthesized titin mRNAs. We isolated Rbm20-bound titin mRNA by RIP. Results showed that Rbm20 protein can be specifically precipitated (Figure 3D), and that the titin mRNAs co-precipitated with Rbm20 are the titin mRNAs with retained introns. In contrast, the constitutively spliced titin mRNAs are barely precipitated (Figure 3E). This result directly shows that Rbm20 binding represses titin splicing. The repression of Rbm20 on titin splicing in the repressed region remains even after the splicing on the rest of the regions is finished: introns 68 and 69 remain retained even after their downstream introns are successfully spliced out (Figure 3F). These results demonstrate that the titin mRNAs bound by Rbm20 are partially processed titin pre-mRNAs; they are not spliced in the Rbm20-repressed regions but are normally spliced in the rest, suggesting that the intron retention we characterized in Figure 1 is from this kind of titin mRNAs. The partially processed titin pre-mRNAs are restricted in the Rbm20 speckle regions in the nucleus instead of being exported.

Rbm20 cooperatively represses the splicing between exons 52 and 68

We amplified the Wt cardiac titin mRNA for the region between exons 52 and 68 using primer pairs with partially overlapped spanning regions. We unexpectedly found constitutive splicing and all-intron retention are the two major splicing patterns for all the primer pairs (Figure 4A). This result illustrates that the all-intron–retained patterns are joined together and constitutive splicing patterns are joined together for the entire region between exons 52 and 68 so that the splicing on this region is cooperative: either all repressed or not repressed. In homozygote mutants (Ht, Rbm20<sup>+/−</sup>), where the Rbm20 expression level is only half of the Wt, the splicing patterns are the same as in the Wt (Figure 4A). The only difference between Wt and Ht was that the all-intron–retained pattern accounts for the majority in the Wt but only for minority in the Ht. Thus, when the Rbm20 concentration is lowered from Wt to Ht, less titin mRNAs are repressed in splicing, but the repression is still cooperative: all or none. To test whether Rbm20 can repress the splicing of titin mRNA on this region cooperatively in other cell types, we constructed minigene expressing titin pre-mRNA from exons 64 to 70 and transfected this minigene into Hela cells with and without Rbm20. To strengthen the interaction between Rbm20 and titin pre-mRNA in Hela cells, we ligated the MS2 stem–loop sequence on either upstream or downstream of titin minigene and fused MS2 coat protein on the N terminus of Rbm20. The splicing assay on the mini-titin pre-mRNA shows that when Rbm20 is artificially recruited to either upstream or downstream positions of titin mRNA, the splicing of all the exons in the mini-titin pre-mRNA is repressed at a higher level (Figure 4B). Therefore, the presence of Rbm20 at a certain position can repress the splicing of not only its adjacent exons but also the exons >2 kb away. In Hela cells, Rbm20 can repress the splicing of mini-titin pre-mRNAs expressing exons/introns in the repressed regions but not those expressing exons/introns outside the repressed regions (Supplementary Figure S6A–C), indicating there should be some native Rbm20-binding sites within the Rbm20-repressed regions.

Rbm20-dependent 3′ alternative splice site selection

Thus far, we found the exon skipping in titin mRNA is associated with the repression of Rbm20 on the splicing in the repressed regions (Figure 2A and B). However, some titin-splicing pathways show skipping of exons in regions not repressed by Rbm20. The N2B-splicing pathway is such an example, in which exon 50 directly splices with exon 219 and skips all the exons in between, including the regions between exons 70 and 79 and exons 88 and 116, even though the splicing in these regions is not repressed by Rbm20. The N2B-splicing pathway is predicted to encode the 2.97-MDa N2B titin isoform (9,18), which is the dominating titin isoform in Wt adult heart (Figure 5A). Based on our intron retention data, we found that exon 50 and exon 219 are located at the 5′ and 3′ end of Rbm20-repressed regions, respectively (Figure 2A), so we have reason to hypothesize that other exons located at the border of Rbm20-repressed regions can be involved in similar alternative splicing pathways. As expected, we found alternative 3′ splice site (ss) pathways in which exon 50 can directly splice with many other exons along titin mRNA besides exon 219, and most acceptor exons involved in the alternative 3′ ss pathway.
**Figure 3.** Rbm20 aggregated on newly synthesized titin mRNAs forms Rbm20 speckles and represses the splicing of titin mRNA.

(A) Immunofluorescence shows two Rbm20 speckles in the nucleus of cardiomyocytes from LV. The size of Rbm20 speckles was similar to nucleoli; Rbm20 is stained red, and nucleoli are stained green with antibody against fibrillarin. 4',6-diamidino-2-phenylindole (DAPI)-stained nuclei are blue. Black scale bar: 2 microns. (B) Immunostaining of Rbm20 protein combined with titin mRNA FISH shows the co-localization of titin mRNA with Rbm20 speckles. Rbm20 is stained red; titin mRNA is stained green; and the overlap of red and green yields yellow. (C) Rbm20 speckles are transcription dependent. When the transcription of HL-1 cell is inhibited, Rbm20 speckles (red) dissociate. The behavior of paraspeckles (continued)
are located at the 3’ borders of Rbm20-repressed regions such as exons 70, 82, 88, 124, 136, 145 and 219; exons 130, 137, 138 and 146 are one or two exons downstream of an Rbm20-repressed region but can also be involved in this splicing pathway (Figure 5B). Whether the unrepressed regions between exons 70 and 79 and exons 88 and 116 would be skipped depends on which acceptor exon is spliced with exon 50 (Figure 5B).

Based on the molecular mass of different cardiac titin isoforms and the splicing pathways we have characterized thus far, the splicing pathways corresponding to the five major cardiac titin isoforms in Hm, Ht and Wt rat from large to small should be as follows: (i) constitutive splicing of all the exons between exons 50 and 225, (ii) skipping of the completely repressed PEVK exons, (iii) additional skipping of the strongly repressed middle Ig exons 50–70, (iv) additional skipping of the mildly repressed middle Ig exons 80–87 plus the unrepressed exons between exons 70 and 79, and (v) additional skipping of unrepressed exons between exons 88 and 116. All these splicing isoforms have been verified by sequencing (Figure 5B).

**Titin isoform transformation during development as well as between muscles is related to the alteration of Rbm20/splicing factors ratio**

Titin undergoes developmental transition from larger neonatal isoforms to smaller adult isoforms, but the development-associated mechanism remains unknown. Because Rbm20 is the protein that regulates titin alternative splicing, we investigated whether developmental changes in Rbm20 lead to the developmental titin isoform transitions. Because Rbm20 is involved in pre-mRNA processing, we tested whether Rbm20 undergoes
developmental changes in parallel with other splicing factors: SFRS1 (SF2/ASF), U2AF65 and hnRNP L. Results using LV showed that during development, the ratio between the latter three splicing factors remained essentially constant, but the ratio of Rbm20 to these splicing factors increased with development. This increased ratio coincides with the progressive reduction in titin size (Figure 6A). Similar results were also found in the skeletal muscle tibialis anterior (TA) (Figure 6B). However, if GAPDH was used as a control, we got exactly the opposite conclusion; namely, the ratio of Rbm20 to GAPDH decreased with age (Figure 6A and B). Because titin isoforms vary between muscles, we could test the relationship between Rbm20 expression and titin size between different muscles. LV, TA and longissimus dorsi (LD) with titin size from small to large were selected for this study. Results show that Rbm20/splicing factors ratio in LV, TA and LD is inversely related with the titin size (Figure 6C). The Rbm20 in vivo always forms two Rbm20 speckles in each nucleus from cardiac and skeletal muscles, and the size of the Rbm20 speckles remains relative constant during development as well as between muscles (Figure 6D), indicating the number and size of Rbm20 speckles are determined by newly transcribed titin mRNA instead of the Rbm20 expression level. The fact that the cellular localization of Rbm20 remains the same during development and between different muscles also suggests that the localization of Rbm20 in the muscle cell is not the factor that affects titin alternative splicing.

Diverse titin isoforms expressed in different muscles suggested different splicing pathways, which allows us to do the correlation analysis on these protein and splicing isoforms. With increasing Rbm20/splicing factors ratio in Wt LD, Wt TA, Ht LV and Wt LV, cooperative skipping of internal exons between exons 50 and 70 starts from Wt TA, which leads to the direct splicing of exon 50 or 51 with exon 70 (Figure 6E). In Ht LV and Wt LV, where Rbm20/splicing factors ratio is at still higher levels, the alternative 3'0 ss pathway, shown by direct splicing of exon 50 to exon 219, can be detected, and corresponding to it, N2B titin isoform is found in these two tissues (Figure 6E). Interestingly, the intron retention between exons 68 and 70 can only be detected in Ht LV and Wt LV (Figure 6E bottom panel), which indicates that when exons 50–70 is cooperatively repressed, the 3' ss of exon 70 could either splice with 5' ss of exon 50 or 51 or remain repressed; the repressed 3' ss of exon 70 is coincident with the appearance of alternative 3' ss pathway. In Wt TA
where the Rbm20/splicing factors ratio is at a lower level, cooperative skipping of exons 51–70 can be detected, but the repression on the 3’ ss of exon 70 cannot; correspondingly, neither alternative 3’ ss pathway nor the titin isoform similar to cardiac N2B can be detected in this tissue (Figure 6E). The cooperative skipping of internal exons between exons 51 and 70 in Wt TA correlates with titin isoforms with discrete sizes. In Wt LD where the Rbm20/splicing factors ratio is too low to initiate the cooperative repression between exons 50 and 70, titin does not show isoforms with discrete size differences (Figure 6E).

We found that the border of the Rbm20-repressed region is not fixed but somewhat flexible. As the relative Rbm20/splicing factors ratio increases from Wt TA to Wt LV, the 5’ border of the Rbm20-repressed region can be extended from exon 51 to 50 (Figure 6E and F). Flexible 5’ and 3’ borders are common for almost all the Rbm20-repressed regions along the titin mRNA: the variable 5’ border includes exon 50 or 51, exon 116 or 117 or 118 and exon 137 or 138; the variable 3’ border includes exon 87 or 88, exons 129 or 130 or 131, exons 136 or 137 and exons 145 or 146 (Figure 2B). Higher Rbm20/splicing factors ratio favors the extension of Rbm20-repressed regions (Figure 6F).

**Full-length titin PEVK exon characterization**

Previous studies with human and rat cardiac titin cDNA have resulted in incomplete amplification of the full set of PEVK exons. This is presumably due to the early embryonic appearance and influence of Rbm20 on titin splicing (Figure 6A). The Rbm20-deficient rat model allowed us to amplify the complete PEVK exon set (Supplementary Table S3). Although most of the exons were similar to the predicted ones in humans, a few individual exons were inserted, and a 14-exon group between exons 175 and 190 (old numbering system to align with human) was missing in the rat. These exons do not appear to be expressed in the human sequences reported to date.
DISCUSSION

After the donor and acceptor exons involved in exon skipping and exon shuffling were located to the partially processed titin pre-mRNA, we found the donor exons (84, 116, 124, 138, 146 and 175) and acceptor exons (70, 82, 87, 88, 124, 129 and 136) involved in exon shuffling also serve as donor and acceptor exons in exon skipping (Figure 7A). Moreover, all these donor and acceptor exons were located at the border of Rbm20-repressed regions, with a donor exon at the 5’ end and an acceptor exon at the 3’ end (Figure 7A). Considering that the aggregation of Rbm20 protein on titin mRNA repressed splicing, we propose that Rbm20-regulated titin alternative splicing occurs as follows: Rbm20 binds to certain regions of the newly transcribed titin pre-mRNA before the splicing in these regions is initiated, and the binding of Rbm20 on titin pre-mRNA blocks splicing so that the introns in these regions cannot be spliced out. At the same time, the rest of titin pre-mRNA can be normally spliced, leading to partially processed titin pre-mRNAs. The partially processed titin pre-mRNAs are retained in the nucleus, awaiting further processing, instead of being exported. Finally, the 5’ ss and 3’ ss on the exons flanking the Rbm20-repressed regions splice with each other to skip the repressed internal exons/introns (Figure 7B). A similar two-step sequential splicing model was recently reported in which the spliced tumor susceptibility gene 101 (TSG101) and fragile histidine triad (FHIT) mRNA can undergo a second round of splicing to mediate extra exon skipping (39).

The 2.97-MDa N2B titin isoform dominates in Wt LV and results from the exon 50–involved alternative 3′ ss pathway that skips most exons between exons 50 and 219 (9,18), including exons 70–79 and 88–116, which are normally spliced in the partially processed titin pre-mRNA. Our correlation study showed the coincidence of alternative 3′ ss pathway with the repressed 3′ ss on exon 70 (Figure 6E), suggesting the use of downstream alternative 3′ ss may result from the unavailability of upstream 3′ ss in the partially processed titin pre-mRNA (Figure 7C). The repression of Rbm20 on the border exons of different repressed regions should be dynamic: the temporal repression of upstream 3′ ss will lead to the use of downstream 3′ ss to induce large-scale exon skipping and the formation of lariats composed of repressed exons/introns and unrepressed exons (Figure 7D). The temporal relief of Rbm20 on border exons within the lariat will lead to re-splicing within the lariat and cause exon shuffling. The model shown in Figure 7D may be the main mechanisms for exon shuffling in Wt titin mRNA, considering the alternative 3′ ss pathway that produce lariats with different exon/intron combinations is the main splicing pathway in Wt LV (Figure 5A and B). No matter what exon/intron combination the lariat has, the exon shuffling should always occur between the border exons, which are consistent with our result. However, we also found some exon-shuffling events in polyA RNA (Supplementary Figure S3C); we think this may result from trans splicing between independent titin messages (Figure 7E). Thus far, we do not have enough data to rule out this possibility. Our previous transcriptomic analysis identified 31 Rbm20-dependent spliced genes that were shared between rats and humans (29). However, there were not a corresponding number of Rbm20 speckles in the nucleus; instead, there are only two Rbm20 speckles per nucleus. These speckles were co-localizing with newly transcribed titin mRNA as determined by combined immunofluorescence and in situ hybridization. Several pieces of evidence suggest that titin mRNA is the major target of Rbm20 regulation. First, titin is the third most abundant protein in the muscle after actin and myosin, so the titin mRNA is probably more abundant than the other regulated genes. Second, the number of potential Rbm20-binding sites in titin pre-mRNA is likely much larger than for any of the other Rbm20-regulated genes. The titin pre-mRNA is ~250,000 nucleotides long, with the Rbm20-repressed region spanning ~100 kb, so there may be a considerable amount of Rbm20 protein bound on each titin pre-mRNA. The apparent absence of speckles associated with other genes could be due to either lower message abundance or fewer Rbm20-binding sites per message.

An interesting question one might ask is whether the exon skipping events shift the open reading frame (ORF) of titin mRNA. The answer is no. After sequencing all the exons in the middle Ig and PEVK region from the Hm (Supplementary Table S3), we found that the nucleotide number of all the other exons from 50 to 225 is a multiple of 3 except for exons 103, 104 and 106. Thus, no matter which exons or combination thereof are skipped, the ORF of titin mRNA will remain the same. The sum of the nucleotide number for exons 103, 104 and 106 is a multiple of 3, and these exons are located in a constitutively spliced region from exon 88 to 116. Therefore, these three exons will either be co-retained or co-skipped during splicing with the other exons of this region, and the ORF of titin mRNA will remain unchanged in either situation.

This is the first mechanistic description of Rbm20 and also the first regarding the mechanism of titin alternative splicing. Our data suggested that Rbm20 proteins aggregate on titin pre-mRNA to regulate alternative splicing by repressing the splicing of certain regions. Different regions of titin pre-mRNA are differentially sensitive to Rbm20 repression, and higher Rbm20/splicing factors ratio can favor the repression on the regions with less sensitivity. However, we do not know whether these Rbm20-repressed blocks on titin pre-mRNA are repressed independently or synergistically. From the data we have characterized thus far, the only factor that is correlated with titin isoform transition is Rbm20/splicing factors ratio, which is fine tuned during development as well as between tissues by totally unknown mechanisms. The undetermined issues also include the mechanism for the cooperative repression of Rbm20 on the region between exons 50 and 70. These questions deserve further efforts to elucidate the molecular mechanism of Rbm20-regulated titin alternative splicing. Our current work provides a foundation for future studies.
Figure 7. Mechanisms of Rbm20-regulated alternative splicing. (A) Schematic diagram depicts the partially processed titin pre-mRNA and the location of the donor and acceptor exons involved in exon skipping. The same donor and acceptor exons are also involved in exon shuffling. The numbers of donor exons and acceptor exons are colored in green and blue, respectively. The thin lines connecting the donor exons with acceptor exons illustrate different splicing pathways. (B) Hypothesized model showing the process of Rbm20-regulated exon skipping as exemplified by exon skipping between exons 50 and 70. The exons are indicated with numbered boxes, and introns with black lines. The unrepressed exons are colored in red, and the Rbm20-repressed exons in black. SnRNPs are indicated with colored circles. U1 defines the 5' ss and U2 defines the 3' ss on titin exons. The association of U4/5/6 to U1 and U2 fulfills the splicing. Splicing between exons 50 and 70 skips the internal exons. (C) Hypothesized model related the upstream 3' ss repression to the use of downstream alternative 3' ss. (D) Hypothesized model showing the process of exon shuffling induced by re-splicing within the lariat spliced out during alternative 3' ss pathway. Here we only show one example, many other alternative 3' ss pathways and exon-shuffling events have been detected from titin mRNA (refer to Figure 5B and Figure 2C). (E) Hypothesized model for trans splicing-induced exon shuffling. Active 5' ss on exon 84 splice with active 3' ss on exon 70 from different titin messages to cause exon shuffling.
SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online: Supplementary Tables 1–3 and Supplementary Figures 1–6.

ACKNOWLEDGEMENTS

The authors thank Zhaolan Zhou (University of Pennsylvania, Philadelphia, PA) for kindly providing the pMAL-MS2-MBP plasmid and pCR2.1 MS 2 construct.

FUNDING

National Institutes of Health (NIH) [HL77196 to M.L.G., and R01HG005232 to C.N.D.]; College of Agricultural and Life Sciences, University of Wisconsin-Madison with support from the National Institute of Food and Agriculture, United States Department of Agriculture [ID number WIS01556 to M.L.G.]. Funding for open access charge: University funds.

Conflict of interest statement. None declared.

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