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Rbm24 controls poly(A) tail length and translation efficiency of crystallin mRNAs in the lens via cytoplasmic polyadenylation

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Running title: Rbm24 regulates cytoplasmic polyadenylation
Abstract

Lens transparency is established by abundant accumulation of crystallin proteins and loss of organelles in the fiber cells. It requires an efficient translation of lens mRNAs to overcome the progressively reduced transcriptional activity that results from denucleation. Inappropriate regulation of this process impairs lens differentiation and causes cataract formation. However, the regulatory mechanism promoting protein synthesis from lens-expressed mRNAs remains unclear. Here we show that in zebrafish, the RNA-binding protein Rbm24 is critically required for the accumulation of crystallin proteins and terminal differentiation of lens fiber cells. In the developing lens, Rbm24 binds to a wide spectrum of lens-specific mRNAs through the RNA recognition motif, and interacts with cytoplasmic polyadenylation element-binding protein (Cpeb1b) and cytoplasmic poly(A)-binding protein (Pabpc1l) through the C-terminal region. Loss of Rbm24 reduces the stability of a subset of lens mRNAs encoding heat shock proteins, and shortens the poly(A) tail length of crystallin mRNAs encoding lens structural components, thereby preventing their translation into functional proteins. This severely impairs lens transparency and results in blindness. Consistent with its highly conserved expression in differentiating lens fiber cells, the findings suggest that vertebrate Rbm24 represents a key regulator of cytoplasmic polyadenylation and plays an essential role in the post-transcriptional control of lens development.

Key words: RNA-binding protein; Rbm24; lens; crystallin; cataract; poly(A) tail; cytoplasmic polyadenylation; CPEB; PABPC; post-transcriptional regulation; zebrafish
**Significance**

Lens transparency critically requires the abundant accumulation of crystallin proteins, and deregulation of this process causes congenital cataracts in humans. Rbm24 is a RNA-binding protein with highly conserved expression in differentiating lens fiber cells among all vertebrates. We use zebrafish model to demonstrate that loss of Rbm24 function specifically impedes lens fiber cell differentiation, resulting in cataract formation and blindness. Molecular analyses reveal that Rbm24 interacts with cytoplasmic polyadenylation complex and binds to a large number of lens-expressed mRNAs to maintain their stability and protect their poly(A) tail length, thereby crucially contributing to their efficient translation into functional proteins. This work identifies an important novel mechanism by which Rbm24 post-transcriptionally controls lens gene expression to establish transparency and refraction power.
Introduction

Lens morphogenesis is temporally and spatially coordinated by a multitude of signaling pathways (1-4). The terminal differentiation of lens fiber cells is characterized by the accumulation of extremely abundant crystallin proteins, as well as the disappearance of cell nuclei and other organelles (3, 5). Crystallins belong to a large family of proteins that constitute the highly abundant soluble proteins in lens fiber cells (6-8), and can be grouped into three main classes: α-, β- and γ-crystallins. α-crystallins are small heat shock proteins that have molecular chaperone activity to ensure the correct folding of other proteins, while β- and γ-crystallins are structural components (7). These proteins play an essential role in lens transparency and refraction, with structure and function particularly conserved among vertebrates (9, 10). Mutations or dysfunctions of crystallin proteins impede lens transparency and lead to different types of cataracts (11), which are the leading cause of human blindness (12). Whereas the transcriptional regulation of lens-specific expression of crystallin genes is relatively well understood, post-transcriptional mechanisms underlying the accumulation of crystallin proteins are largely unclear, especially when transcriptional contribution slows down during denucleation of lens fiber cells.

RNA-binding proteins (RBPs) are implicated in post-transcriptional regulation of gene expression in a variety of biological processes (13-15). Through tissue-specific expression and interaction with specific cofactors, RBPs-constituted mRNA interactome controls RNA metabolism at multiple levels (16, 17). Several RBPs are linked to congenital cataracts and other lens abnormalities in humans or animal models (18-20). The highly conserved Rbm24 protein contains a single RNA recognition motif (RRM) and exhibits tissue-specific expression patterns in all vertebrate species. It plays an important role in skeletal and cardiac muscle development (21-24), by regulating mRNA stability (25, 26), pre-mRNA splicing (24, 27-30), and mRNA translation (31). Moreover, its expression in head sensory organs, particularly in the lens, is also conserved in vertebrates (23, 32). Recent studies have associated rbm24 gene with microphthalmia and anophthalmia (33, 34), highlighting its implication in vertebrate eye development.
Nevertheless, the molecular and biochemical mechanisms by which it orchestrates lens development remain elusive.

Here, we report the functional implication of Rbm24 in lens differentiation using zebrafish embryos. We present the unexpected finding that it is critically required for maintaining the stability and poly(A) tail length of lens-expressed mRNAs. The expression of rbm24 gene is restricted in differentiating lens fiber cells, and its loss of function severely compromises the accumulation of crystallin proteins in the developing lens. Moreover, Rbm24 binds to a wide spectrum of lens-specific mRNAs and interacts with key components of the cytoplasmic polyadenylation complex. Thus, we uncover an important novel mechanism by which Rbm24 post-transcriptionally controls lens gene expression through cytoplasmic polyadenylation.
Results

Zebrafish *rbm24a* is expressed in differentiating lens fiber cells

There are two *rbm24* paralogs in zebrafish, *rbm24a* and *rbm24b*. The expression patterns of *rbm24a* in the somites, heart, and lens are identical as those of *rbm24* in other vertebrates (22, 23), whereas *rbm24b* mainly shows heart and somitic expression (35). In situ hybridization analysis on ocular sections indicated that *rbm24a* expression was restricted to differentiating lens fiber cells. At 1 dpf (day post-fertilization), *rbm24a* transcripts were detected in the posterior and central regions of the lens mass, where differentiation of primary fiber cells and initial formation of secondary fiber cells take place (36, 37). They were not present in the anterior lens mass including the prospective lens epithelium (Fig. 1A). At 1.5 and 2 dpf, *rbm24a* transcripts were localized in the posterior and lateral regions, and remained absent in the lens epithelium (Fig. 1B and C). At 3 dpf, the transcripts became further restricted to differentiating fiber cells in the posterior border and the lateral zone (Fig. 1D). This highly localized expression pattern suggests that Rbm24a may be involved in the differentiation of lens fiber cells.

Zebrafish *rbm24a* mutants display severe cataract phenotype and defective lens differentiation

We used TALENs (transcription activator-like effector nucleases) to create mutation in the *rbm24a* locus, and obtained a deletion of 8 base pairs in the first exon, which caused a frameshift in the coding region of the RRM and a premature termination of translation (Fig. S1A-C). RT-PCR analysis further confirmed the mutation at mRNA level (Fig. S1D). Homozygous mutants showed complete penetrance of severe heart defects, with heart edema becoming evident at 1.5 dpf. As development proceeds, they also developed microcephaly, which was associated with bilateral microphthalmia (Fig. S2). Examination of the ocular phenotypes revealed defective lens development. In wild-type (WT) siblings, both the retina and the lens progressively increased in size from 2 to 4 dpf (Fig. 2A, C and E), and the lens developed a sleek surface, reflecting the establishment of transparency (Fig. 2A’, C’ and E’). In time-matched *rbm24a* mutants, however, the retina and the lens were strongly reduced in size (Fig. 2B, D and
and the lens was particularly coarse and highly irregular, implying a lack of transparency (Fig. 2B', D' and F'). Thus, the mutants showed a severe cataract phenotype. Heterozygous mutants developed normally, while homozygous embryos could not survive beyond 7 dpf primarily due to heart defects.

Analysis of ocular sections showed no obvious differences between WT siblings and rbm24a mutants at 2 dpf (Fig. S3A and B). However, the reduction of lens size and the retention of fiber cell nuclei were particularly evident in rbm24a mutants at 3 and 4 dpf (Fig. S3C-F). TUNEL labeling did not detect obvious increased cell death in the lens of rbm24a mutants when compared with time-matched WT siblings (Fig S4A-C), but few BrdU-positive cells were present in the lens epithelium, indicating reduced cell proliferation (Fig. S4D-F). Consistently, there were few lens epithelial cells undergoing cell cycle progression in rbm24a mutants, as revealed by anti-phospho-Histone H3 antibody (Fig. S4G-I). We then monitored lens differentiation defects by staining ocular sections with phalloidin and Hoechst. At 2 dpf, F-actin and fiber cell nuclei were present similarly in WT siblings and rbm24a mutants (Fig. 2G and H). At 3 dpf, F-actin was organized as concentric rings at the periphery of the lens, and the central lens nucleus showed weak phalloidin staining and a complete absence of fiber cell nuclei in WT siblings (Fig. 2I). By contrast, disorganized F-actin and degenerating fiber cell nuclei were present in rbm24a mutants (Fig. 2J). At 4 dpf, the lens further differentiated in WT siblings, with complete disappearance of F-actin in the central lens nucleus (Fig. 2K). However, phalloidin-stained large aggregates and Hoechst-positive fragmented nuclei were still present in rbm24a mutants (Fig. 2L). Thus, these defects lead to the development of a cataract phenotype.

Immunofluorescence staining using specific markers was performed to further examine lens differentiation defects. Pax6 transcription factor is localized in the anterior epithelium during lens differentiation (38). When compared with WT siblings from 2 to 4 dpf, this expression pattern was not affected in rbm24a mutants (Fig. S5A-F), indicating that Rbm24a is not required for the specification and maintenance of lens epithelium. This is consistent with its absence of expression in these cells. AQP0 protein marks elongating lens fiber cells (39), its staining pattern clearly revealed a defective elongation of posterior lens fiber cells in rbm24a mutants.
when compared with time-matched WT siblings (Fig. S5G-L). These analyses indicate that mutation of Rbm24a severely impairs the differentiation of lens fiber cells.

**Loss of Rbm24a causes cataract formation independently of fiber cell denucleation**

The heart valve defects in *rbm24a* mutants block blood circulation (Movie S1). Moreover, *rbm24a* is not expressed in the retina. Thus, the microphthalmia defects could be caused by the absence of blood supply. We examined this possibility by specifically expressing an *rbm24a-GFP* transgene in the heart of *rbm24a* mutants, under the control of a *cmlc2* promoter region (Fig. S6A and B). As might be expected, the transgene fully rescued heart and microphthalmia defects (Fig. S6C-H), but it did not prevent cataract formation (Fig. S6I-K). Furthermore, heterotopic parabiosis, by joining a WT embryo with an *rbm24a* mutant at 4 hpf (hours post-fertilization), restored blood circulation and prevented microphthalmia in the mutant partner (Fig. 3A and B, Movie S2). However, the mutant lens remained disorganized, lacking a regular and smooth surface (Fig. 3C and D). The opacity of the mutant lens was confirmed using confocal reflection mode. In WT embryos at 3.5 dpf, the lens showed no or weak reflected signal (Fig. 3E and K). By contrast, in unparabiosed and parabiosed *rbm24a* mutants, the lens displayed stronger reflection (Fig. 3F, G and K). We also detected the presence of large reflected granules in the lens of unparabiosed *rbm24a* mutants, which is likely due to defective fiber cell denucleation. Indeed, confocal microscopic analysis following staining with phalloidin and Hoechst indicated that nuclear degradation was severely impaired in unparabiosed *rbm24a* mutants, whereas this was only weakly delayed in parabiosed *rbm24a* mutant partners (Fig. 3H-J and L). These observations strongly suggest that both microphthalmia and altered fiber cell denucleation in *rbm24a* mutants are indirect consequence due to the absence of blood supply. Thus, loss of Rbm24a should directly impair other processes that promote lens transparency.

**The overall abundance of lens-specific transcripts is differentially affected in *rbm24a* mutants**

We then performed RNA-sequencing (RNA-seq) to analyze Rbm24a-regulated events. Indirect changes were minimized by preparing polyadenylated mRNA samples from whole embryos or
head regions at 33 hpf, when morphological and histological alterations of the eye were not apparent. Compared with previously reported muscle-specific alternative splicing events in mice (24), the RNA-seq data revealed that 60.9% (42/69) of them displayed similar alternative splicing patterns in zebrafish (Dataset S1). However, identification of splice junctions did not reveal meaningful differences in alternative splicing events for lens-specific pre-mRNAs between WT embryos and rbm24a mutants (Dataset S2). Rather, differential gene expression analysis indicated a strongly reduced expression level of a large number of lens-related genes in rbm24a mutants (Fig. 4A, Fig. S7). This was further validated by qRT-PCR analysis with cDNAs reverse-transcribed using oligo(dT) primer (Fig. 4B).

Based on the RNA-seq and qRT-PCR data, we selected 10 genes and performed in situ hybridization to compare their expression patterns between WT siblings and rbm24a mutants at 33 hpf. Unexpectedly, only a few genes that encode heat shock proteins, including cryaa, hsp70, and hsp90aa1.2, displayed strongly reduced expression in rbm24a mutants (Fig. 4C-E and C'-E'). By contrast, a large number of crystallin genes, including crygm2d1-21, crygmx cryba1b, cryba2b, cryba4, crybb1, and crybb1l1, showed similar expression between WT siblings and rbm24a mutants (Fig. 4F-L and F'-L'), suggesting that the overall abundance of their transcripts was not affected in rbm24a mutants. Because RNA-seq and qRT-PCR analyses used polyadenylated mRNAs, the results from in situ hybridization raises the possibility that loss of Rbm24a may essentially affect the polyadenylation, but not the overall abundance of those crystallin mRNAs coding for structural components.

**Rbm24a binds to a wide spectrum of lens-expressed mRNAs**

To provide direct evidence on the implication of Rbm24a in regulating crystallin gene expression, we analyzed its interaction with lens-expressed mRNAs by RNA immunoprecipitation and qPCR (RIP-qPCR). This was performed using transgenic lines specifically expressing Rbm24a-GFP or Rbm24aΔRRM-GFP fusion proteins in the lens, under the control of a cryaa promoter region (Fig. 5A and B). Confocal microscopic analysis indicated that both fusion proteins were localized in the cytoplasm. Interestingly, Rbm24a-GFP showed
granular staining at 2 dpf (Fig. 5C), and displayed a punctate pattern in elongating lens fiber cells at 3 dpf (Fig. 5D). This is likely a result of its association with RNAs. In sharp contrast, Rbm24aΔRRM-GFP was present diffusely and uniformly in lens fiber cells at different stages (Fig. 5E and F), probably due to the lack of binding with RNAs. Indeed, we found that Rbm24a-GFP bound to multiple lens-related transcripts with different efficiency, whose overall abundance was either decreased (cryaa and hsp70) or unchanged (cryba1b, cryba4, crybb1, crygm2d1, crygm2d1-21, crygmx, crybb111) in rbm24a mutants (Fig. 5G). The RRM was required for interaction with these transcripts, because no PCR products could be detected using transgenic embryos expressing Rbm24aΔRRM-GFP (Fig. 5H). Furthermore, neither Rbm24a-GFP nor Rbm24aΔRRM-GFP interacted with otx2 transcripts, which were not expressed in the lens (Fig. 5G and H). These results show that Rbm24a specifically interacts with a large number of lens-expressed mRNAs.

**Loss of Rbm24a reduces the stability and poly(A) tail length of crystallin mRNAs**

It is unclear whether the reduced overall transcript abundance of lens mRNAs in rbm24a mutants could be caused at transcriptional level. We thus generated a transgenic line by placing GFP coding region along with β-globin leader sequence and SV-40 poly(A) signal under the control of a cryaa promoter region (Fig. 6A). We reasoned that this transgene should faithfully recapitulate the transcriptional activation of the endogenous gene, but the stability and polyadenylation of the resulting transcripts should not be affected in the absence of Rbm24a. Indeed, qPCR analysis following reverse transcription using oligo(dT) primer showed a similar level of GFP expression in WT siblings and rbm24a mutants (Fig. 6B). There was also a comparable kinetics and intensity of lens-specific fluorescence in live embryos (Fig. 6C). These indicate that Rbm24a is not involved in the transcriptional control of cryaa, at least not through the promoter region used here. To determine whether loss of Rbm24a affects the stability of cryaa transcripts, head regions dissected from WT siblings and rbm24a mutants were treated with actinomycin D to inhibit de novo RNA synthesis, and the levels of cryaa transcripts at different time intervals were analyzed by qRT-PCR. We found a more rapid decline of cryaa
transcripts in \textit{rbm24a} mutants, with a half-life of about 1.86 hours compared to 2.92 hours in WT siblings (Fig. 6D). Thus, Rbm24a functions to maintain the stability of a subset of lens mRNAs.

For most other \textit{crystallin} genes, only their polyadenylated mRNAs showed a decreased expression level in \textit{rbm24a} mutants. To further validate this unexpected and potentially novel function of Rbm24a, we used poly(A) tail-length assay to examine the polyadenylation state of \textit{crygm2d1}, \textit{crybb1l1}, \textit{crygmx}, \textit{cryba1b}, and \textit{crybb1} mRNAs. Following poly(G/I) tailing and reverse transcription, the poly(A) PCR-amplified products from \textit{rbm24a} mutants showed a weak shift with respect to those from WT siblings (Fig. 6E). Furthermore, sequence analysis confirmed an overall reduction of poly(A) tails (Fig. 6F-H). As an example, the average poly(A) tail length for \textit{crygmx} transcripts in WT embryos was 86 nucleotides, whereas this was reduced to 42 nucleotides in \textit{rbm24a} mutants (Fig. 6H, Dataset S3). More strikingly, a large amount of \textit{crygmx} transcripts showed a complete absence of poly(A) tail, such that the 3’-untranslated region terminated at the polyadenylation signal (PAS), whereas in WT embryos the poly(A) tail followed the general rule by starting at about 20 nucleotides downstream of this signal (Fig. 6I). These results strongly suggest that Rbm24a is critically required for the elongation or maintenance of poly(A) tail length in a large number of \textit{crystallin} mRNAs.

\textbf{Rbm24a is required for the accumulation of crystallin proteins}

To determine the functional consequence of this reduced polyadenylation on the accumulation of crystallin proteins, we first generated a transgenic line by inserting GFP coding sequence into the third exon of \textit{crygm2d1} gene, before its stop codon (Fig. 7A). Thus, the expression of crygm2d1-GFP fusion protein under the control of \textit{cryaa} promoter should closely reproduce the actual condition with respect to \textit{crygm2d1} mRNA expression and translation. Indeed, in situ hybridization analysis using \textit{GFP} probe revealed a similar overall abundance of the fusion transcripts between WT siblings and \textit{rbm24a} mutants at 2 dpf (Fig. 7B). However, qPCR analysis following reverse transcription using oligo(dT) primer showed that the amounts of polyadenylated \textit{crygm2d1}-\textit{GFP} mRNAs were markedly reduced in \textit{rbm24a} mutants (Fig. 7C), suggesting an absence or severe shortening of the poly(A) tail. Consistently, the expression
level of the fusion protein was severely reduced in rbm24a mutants, as determined by Western blot analysis using GFP antibody (Fig. 7D and E). The decreased translational efficiency of crygm2d1-GFP fusion mRNA was also evident in live embryos. Compared with WT siblings, time-matched rbm24a mutants showed very weak or no lens-specific fluorescence (Fig. 7F). Furthermore, analysis by quantitative proteomics confirmed a global decrease of endogenous crystallin proteins in rbm24a mutants (Fig. 7G, Fig. S8, Datasets S4 and S5). Thus, these observations demonstrate a critical requirement for Rbm24a in the accumulation of crystallin proteins.

Since restoration of blood circulation in rbm24a mutants led to nearly normal degradation of fiber cell nuclei, the decreased synthesis of crystallin proteins should represent a direct and autonomous effect of Rbm24a mutation on cataract formation. This was supported by cell transplantation experiments (40, 41), in which fluorescently labeled cells were implanted in the presumptive lens placode region of stage-matched recipient embryos (Fig. S9A and B). In WT recipients receiving rbm24a mutant donor cells, the lens displayed a cataract-like phenotype at 3.5 dpf, and this caused blindness in the chimeric larvae, as they were unable to prey the paramecia (Fig. S9C-F, Movie S3). We further used the crygm2d1-GFP reporter to monitor fiber cell differentiation. Compared to WT cells, transplanted rbm24a mutant cells integrated into the WT lens failed to express the fusion protein at 1.5 dpf (Fig. S9G and H), but they were expected to undergo normal denucleation because of the presence of blood supply. By contrast, WT cells were able to express the fusion protein in rbm24a mutant recipients, regardless of the absence or presence of blood circulation (Fig. S9I and J). These results clearly show that Rbm24a regulates the accumulation of crystallins in a cell-autonomous manner.

**Rbm24a interacts with the cytoplasmic polyadenylation complex**

The critical implication of Rbm24a in maintaining poly(A) tail length to promote translational efficiency raises the question whether and how it interacts with the cytoplasmic polyadenylation complex. Among different components that participate in the formation and function of this complex, cytoplasmic polyadenylation element-binding protein (CPEB) promotes poly(A) tail
elongation when it is phosphorylated, and cytoplasmic poly(A)-binding protein (PABPC) stimulates translation and protects poly(A) tail from degradation (42, 43). By co-immunoprecipitation using full-length and truncated Rbm24a proteins (Fig. 8A), we found that GFP-tagged Rbm24a specifically bound to myc-tagged zebrafish Pabpc1l and Cpeb1b (Fig. 8B and C). In addition, the C-terminal region but not the RRM was directly involved in these interactions (Fig. 8D and E). Interestingly, the C-terminal region of Rbm24a showed a stronger affinity with Pabpc1l than with Cpeb1b, suggesting that Rbm24a may interact with these proteins via different sub-domains, and that the binding of Rbm24a C-terminal region with Cpeb1b may also need the assistance of the RRM. These analyses indicate that Rbm24a may participate in the translational activation of lens-expressed mRNAs, by regulating the elongation and the stability of the poly(A) tail through interaction with Cpeb1b and Pabpc1l, respectively (Fig. 8F). Altogether, we have revealed that vertebrate Rbm24 is an important component of the lens mRNA interactome, and demonstrated that it represents a novel regulator of cytoplasmic polyadenylation.
Discussion

The establishment of lens transparency is coordinated by the accumulation of functional crystallin proteins and degradation of organelles. The transcriptional control of lens differentiation is relatively well documented, but the post-transcriptional regulation remains largely unknown. Particularly, when transcriptional contribution declines during denucleation of differentiating lens fiber cells, it becomes important to set up a mechanism that promotes the efficient translation of lens mRNAs. However, the cytoplasmic control of protein expression mediated by RNA-binding proteins is generally less understood mechanistically (42, 43). This work reveals a critical role of vertebrate Rbm24 in regulating cytoplasmic polyadenylation during lens differentiation. We show that loss of Rbm24 impairs the stability and polyadenylation of lens-expressed mRNAs encoding regulatory and structural components in lens fiber cells, thus preventing the accumulation and correct folding of crystallins, which eventually leads to cataract formation.

Due to its multiple expression sites, constitutive knockout of rbm24a gene in zebrafish produces pleiotropic morphological alterations, including impaired blood circulation that delays eye development and lens denucleation. Our results establish that the reduced eye size and impaired fiber cell nuclear degradation are indirect consequences due to the absence of blood supply. Restoration of blood circulation in rbm24a mutants by transgenic correction of heart defects or by fusion of the circulatory systems through heterotypic parabiosis completely rescues eye size and leads to essentially normal denucleation in lens fiber cells, but this does not prevent the occurrence of the cataract phenotype. A direct consequence of rbm24a mutation in producing lens opacity is unequivocally proved by transplantation of rbm24a mutant cells into the developing lens of WT recipients, which causes cataract formation and leads to blindness. These observations strongly suggest that Rbm24a functions in an autonomous manner to regulate lens differentiation, independently of organelle degradation. Indeed, it is known that lens fiber cell denucleation is controlled by regulated transcription and activity of acidic DNase (44-46). Thus, it will be of interest to investigate how blood supply and DNase activity cooperate in this process.
The lens refractive power is ensured by an extremely high concentration of crystallins (8). Our RNA-seq data indicate that the expression level, but not the alternative splicing of lens-specific genes, is affected in rbm24a mutants. Furthermore, Rbm24a binds to a large number of lens-specific mRNAs and regulates their translation efficiency via distinct mechanisms, but with the same outcome for the accumulation of crystallin proteins. First, Rbm24a is required for maintaining the stability of a subset of lens mRNAs, such as cryaa, hsp70, and hsp90aa1.2. Thus, their rapid decay in rbm24a mutants affects the accumulation of the corresponding proteins that have molecular chaperone activity to ensure the correct folding and solubility of other lens proteins. Second, loss of Rbm24a causes the shortening of poly(A) tail length in those crystallin mRNAs encoding structural components, without affecting the overall RNA abundance. This markedly impairs their translation and clearly represents a novel mechanism by which Rbm24a regulates mRNA fate during lens fiber cell differentiation. It is well established that one of the important functions of poly(A) tails is the regulation of translational state through interaction with the cytoplasmic polyadenylation complex and the translation initiation complex (43, 47). During lens differentiation, the rapid degradation of fiber cell nuclei may preclude a continuous transcription of crystallin genes. As a compensatory mechanism, the transcribed mRNAs should be either stabilized and/or processed for efficient translation into proteins. Rbm24a may function as a major regulator mediating these critical processes. In the absence of Rbm24a function, both the reduced stability and shortened poly(A) tail length in lens mRNAs prevents the accumulation of crystallin proteins, and consequently decrease lens refractive power. Thus, our finding emphasizes a predominant role of post-transcriptional control during lens differentiation, and supports previous observations showing the involvement of translation initiation machinery in lens development (48).

It has become increasingly evident that cytoplasmic polyadenylation regulates poly(A) tail elongation of nuclear exported mRNAs and plays an important role in translation activation during oogenesis and early embryonic development (49). This process is mediated by cytoplasmic poly(A) polymerases through interaction with specific RNA-binding proteins, such as CPEB that recognizes U-rich cytoplasmic polyadenylation element present in target mRNAs.
In addition, PABPC interacts with other components of the cytoplasmic polyadenylation complex, and promotes binding of the complex to newly produced poly(A) tails. Thus, it can activate translation and prevent mRNAs from deadenylation (50). Based on the requirement of its RRM in binding to a large number of crystallin mRNAs, and its C-terminal domain in interaction with CPEB and PABPC, we could postulate that Rbm24a may function as a novel component in the cytoplasmic polyadenylation machinery to promote efficient translation of lens-specific mRNAs. Although the precise mechanism awaits further investigation, it is possible that the interaction between Rbm24a and CPEB may activate the cytoplasmic polyadenylation complex for the elongation of poly(A) tails, this is consistent with an overall shortening of poly(A) tails in crystallin mRNAs following loss of Rbm24a. In addition, Rbm24a may interact with PABPC to protect lens mRNAs from deadenylation, this is supported by the complete disappearance of poly(A) tail in crygmx transcripts when Rbm24a function is absent. Therefore, we unveil a critical role of vertebrate Rbm24 in cytoplasmic polyadenylation. This unprecedented function of Rbm24 may be not limited to lens differentiation, it may also operate in other tissues or cells where Rbm24 is localized in the cytoplasm, such as inner ear, nasal epithelium, and myoblasts (23, 32).

Our recent work shows that mouse Rbm24 protein is first localized in the cytoplasm of lens fiber cells at the posterior region of the lens vesicle, and then in the equatorial epithelial cells, which will give rise to the secondary lens fibers (32). As these cells differentiate, they terminally leave the cell cycle, start to elongate, and express numerous genes that help to establish lens transparency (51, 52). Zebrafish rbm24a mRNA is also restricted to differentiating and elongating lens fiber cells, and Rbm24a protein is localized in the cytoplasm as a punctate pattern, reminiscent of binding with RNAs. Therefore, the conserved expression patterns and the cataract phenotype in rbm24a mutants are fully consistent with a critical role of vertebrate Rbm24 in controlling important events related to lens fiber differentiation. Taken together, evidence now available clearly shows that Rbm24 represents a major post-transcriptional regulator of embryonic cell lineage differentiation. Its cell-type specific expression in different tissues is fully consistent with a critical role in orchestrating the initiation and maintenance of cell
differentiation. Moreover, Rbm24 can act as a multi-functional RBP that governs tissue-specific, and also stage-specific regulation of gene expression. These functions are likely accomplished by its dynamic subcellular localization, and interaction with specific partners. Thus, it will be important in the future to determine the regulated changes of Rbm24 cellular compartmentalization, and its tissue- and stage-specific functions.

In summary, we show that vertebrate Rbm24 plays a key role in controlling the translation efficiency of tissue-specific mRNAs. This may likely represent a general function of Rbm24 in regulating various differentiation processes. Because the implication of RBPs in eye development and disease (53), and the similarities in structure and regulation between zebrafish and human lenses (54), understanding the post-transcriptional mechanisms that help to produce high levels of transparent proteins in the lens could open novel perspectives for the prevention and treatment of cataracts. Furthermore, given the highly conserved expression of vertebrate Rbm24 in the lens, it will be particularly interesting to identify human mutations affecting the cis-regulatory elements that disrupt its lens-specific expression and lead to congenital cataracts.
Materials and Methods

Zebrafish and *rbm24a* mutants. Adults of the AB strain were maintained at 28.5°C in standard housing systems (Haisheng, Shanghai). TALEN repeat variable di-residues targeting the first exon of *rbm24a* gene were engineered through the Golden Gate Assembly method (55). The constructs were assembled in the modified pCS2+KKR and pCS2+ELD vectors (56). Screening of F0 founder and F1 offspring were performed by sequencing PCR products amplified from tail fin genomic DNA (57). Identified *rbm24a* heterozygous fish were used for different experiments. Additional materials and methods are available in SI Materials and Methods.

Data availability

The sequencing data reported in this paper have been deposited in National Center for Biotechnology Information Gene Expression Omnibus (GEO) database, https://www.ncbi.nlm.nih.gov/geo (accession no. GEO136006 and GEO136003).
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Author contributions

M.S. performed molecular cloning, RNA-seq, RIP-qPCR, poly(A) tail assay, transgenic analysis, histology, micromanipulation and confocal analysis; T.L., C.Z. and Y.Z.Z. performed molecular cloning, genotyping, in situ hybridization, co-immunoprecipitation, Western blot, TUNEL assay and immunofluorescence staining; S.H.K. initiated characterization of rbm24a mutants; M.S. and D.L.S. designed the experiments, analyzed the data, obtained grant supports, and wrote the paper.

Competing interests

The authors declare no competing interest.
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Fig. 1. Expression of zebrafish *rbm24a* transcripts during lens development. (A) At 1 dpf, the expression of *rbm24a* transcripts (purple) is localized in differentiating fiber cells at the posterior and central regions of the lens mass. (B-D) From 1.5 to 3 dpf, the expression of *rbm24a* becomes progressively restricted to the posterior lens border and lateral zone. The transcripts were not present in the anterior lens epithelium (Le) at all stages of lens differentiation. Arrows indicate the anterior limit of *rbm24a* expression, corresponding to the equator. The sections were counterstained with eosin. Scale bar, 50 µm.

Fig. 2. Cataract phenotype and lens differentiation defects in *rbm24a* mutants. (A-F) Live images compares eye phenotypes between WT siblings and time-matched *rbm24a* mutants from 2 to 4 dpf. Scale bar, 100 µm. (A'-F') Differential interference contrast images illustrate the cataract phenotype in surgically dissected lenses. In WT siblings, the lens progressively increases in size, with fiber cells forming regular and refractive concentric layers. In *rbm24a* mutants, the lens shows no growth, with an irregular surface and a granular aspect. Scale bar, 50 µm. (G-L) Staining of lens sections with phalloidin and Hoechst 33342 at the level of optic nerve compares fiber cell organization and denucleation between WT siblings and *rbm24a* mutants. Scale bar, 50 µm.

Fig. 3. Rescue of lens fiber cell denucleation but not lens transparency in *rbm24a* mutants by heterotypic parabiosis. (A) Parabiosis by surgically joining an unlabeled WT embryo and an RLDx-labeled *rbm24a* mutant (red). Scale bar, 200 µm. (B) Parabiotic WT and *rbm24a* mutant partners at 3.5 dpf, with eyes outlined by dashed circles. Scale bar, 100 µm. (C and D) Differential interference contrast images show the lens phenotypes in parabiotic WT and mutant partners. Scale bar, 50 µm. (E-G) Lens transparency determined by confocal reflection mode, with vertical color bar indicating colors of 10 focal planes at a depth of 30 µm. Scale bar, 50 µm. (H-J) Staining of lens sections with phalloidin and Hoechst 33342 shows lens fiber cell denucleation in WT, unparabiosed and parabiosed *rbm24a* mutants. Scale bar, 50 µm. (K) Scatter plot compares lens transparency by analyzing the integrated density of the scanned lens region in the maximum projection image obtained under confocal reflection mode. Horizontal
lines and error bars represent the mean value ± s.d. from three independent lenses (*, p < 0.05; ***, p < 0.001, Student’s t-test). (L) Statistical analysis of lens fiber cell denucleation by examining the number of fiber cell nuclei present within 2/3 radius of a lens section at the level of optic nerve (outlined by dashed yellow circles in H-J). This area shows complete denucleation in WT embryos at 3.5 dpf. The results were obtained from three independent lens sections (**, p < 0.01, Student’s t-test).

**Fig. 4. Reduced expression level of polyadenylated crystallin mRNAs in rbm24a mutants.** (A) RNA-seq analysis of whole embryos at 33 hpf. Volcano plot shows the down-regulation of lens-related genes coding for heat shock proteins and other crystallins in rbm24a mutants (represented by green dots). Red dots show up-regulated genes, with Padj less than 0.05 and fold change more than 1.5; blue dots represent down-regulated genes, with Padj less than 0.05 and fold change less than 0.67. (B) qRT-PCR analysis compares crystallin mRNA levels in WT siblings and rbm24a mutants, following reverse transcription using oligo(dT) primer. The results were obtained from three independent experiments, using different batches of embryos. (C-L) In situ hybridization analysis of crystallin gene expression patterns at 33 hpf. Representative images show strongly reduced expression of cryaa, hsp70, and hsp90aa1.2 in rbm24a mutants. The overall transcript abundance for other crystallin genes does not change. Due to the extremely high level of nucleotide sequence identity, the crygm2d1-21 probe recognizes transcripts of all 21 types of γ-crystallins. Scale bar, 100 μm.

**Fig. 5. Rbm24a protein is localized in the cytoplasm of lens fiber cells and specifically binds to lens-expressed mRNAs.** (A and B) Transgenic embryos express Rbm24a-GFP or Rbm24aΔRRM-GFP fusion proteins in the lens under the control of a cryaa promoter region. (C-F) Lens sections show differential patterns of cytoplasmic localization of the two fusion proteins at 2 and 3 dpf. Scale bar, 20 μm. (G) RIP-qPCR analysis shows the interaction between Rbm24a and lens-expressed mRNAs. For each RIP reaction, 10% of the supernatant was used as input, and the value obtained after PCR amplification was normalized as 1. The remaining supernatant was processed for RNA purification in the presence of anti-GFP antibody (IP+) or
anti-myc antibody (IP-). Non-specific binding of mRNAs to antibody-conjugated beads was controlled by amplification of *otx2*. The results were obtained from three independent experiments, using different batches of embryos.  

(F) RIP-qPCR analysis shows absence of interaction between Rbm24aΔRRM and different mRNAs.

**Fig. 6. Reduced stability and poly(A) tail length of crystallin mRNAs in *rbm24a* mutants.**  
(A) Schematic representation of the GFP reporter under the control of *cryaa* promoter. (B) qRT-PCR analysis compares the expression of GFP transcripts in WT embryos and *rbm24a* mutants. Bars represent the mean value ± s.d. from three independent experiments (NS, not significant; Student’s *t*-test). (C) Live images show lens-specific fluorescence in WT siblings and *rbm24a* mutants. Scale bar, 100 µm. (D) Rapid decay of endogenous *cryaa* mRNA in *rbm24a* mutants. The initial mRNA levels in WT siblings and *rbm24a* mutants were normalized as 1 in triplicate experiments. (E) Analysis of poly(A) tail length in crystallin mRNAs. Smear bands from poly(A) PCR products in WT embryos are indicated by asterisks. The corresponding bands from *rbm24a* mutants (-/-) show a shift in gel mobility. RT+, reverse transcription; RT-, no reverse transcription. (F-H) Scatter plots show differences of poly(A) tail length in three crystallin mRNAs between WT embryos and *rbm24a* mutants. Horizontal lines and error bars represent the mean value ± s.d. from all sequenced clones (***, *p* < 0.001; ****, *p* < 0.0001, Student’s *t*-test). (I) Sequencing chromatograms show the complete absence of polyadenylation of *crygmx* transcripts in *rbm24a* mutants.

**Fig. 7. Reduced translation efficiency of crystallin mRNAs in *rbm24a* mutants.** (A) Schematic representation of *crygm2d1-GFP* transgene. The three exons (E1, E2, and E3) of *crygm2d1* gene are shown. Dark blue represents 5′- and 3′-untranslated regions that will be present in the fusion transcripts. (B) In situ hybridization analysis using GFP probe detects a similar expression pattern of the fusion transcripts in WT embryos and *rbm24a* mutants at 2 dpf. Scale bar, 100 µm. (C) qPCR analysis following reverse transcription using oligo(dT) primer shows a residual level of polyadenylated *crygm2d1-GFP* mRNAs in *rbm24a* mutants at 2 dpf. The data were obtained from three independent samples of different batches of embryos, and
the expression level in WT embryos is set as 1 after normalization to glyceraldehyde-3-phosphate dehydrogenase (***, p < 0.001; Student’s t-test). (D) Western blot analysis using anti-GFP antibody shows a trace amount of crygm2d1-GFP protein in rbm24a mutants at 2 dpf. β-actin is an input control. (E) Quantification of protein bands after normalization to β-actin using three independent Western blots (***, p < 0.001; Student’s t-test). (F) Live images show marked differences in the lens-specific expression of crygm2d1-GFP between WT embryos and rbm24a mutants at 1.5 and 2.5 dpf. Scale bar, 100 µm. (G) Analysis by quantitative proteomics shows a globally decreased accumulation of endogenous crystallin proteins in rbm24a mutants at 2 dpf.

Fig. 8. Rbm24a physically interacts with Pabpc1l and Cpeb1b through the C-terminal region. (A) Schematic representation of GFP-tagged full-length and truncated Rbm24a proteins. (B and C) The full-length Rbm24a interacts with Pabpc1l and Cpeb1b (arrowheads). (D and E) The C-terminal region of Rbm24a co-immunoprecipitates with Pabpc1l and Cpeb1b (arrowheads), but the interaction with Cpeb1b is weak. (F) Possible function of vertebrate Rbm24 in poly(A) tail elongation and stabilization. In the lens, Rbm24 directly associates with CPEB and promotes the activity of PAPD4/Gld2 poly(A) polymerase in poly(A) tail elongation; it also interacts with PABPC to facilitate loop formation and stabilize poly(A) tail. The RNA motif that recruits Rbm24 remains to be determined. EIFC, eukaryotic translation initiation factor complex; PBE, PUM2-binding element; CPE, cytoplasmic polyadenylation element; PAS, polyadenylation signal; N, Rbm24 N-terminal region; C, Rbm24 C-terminal region.
