

Birc7: A Late Fiber Gene of the Crystalline Lens

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PURPOSE. A distinct subset of genes, so-called “late fiber genes,” is expressed in cells bordering the central, organelle-free zone (OFZ) of the lens. The purpose of this study was to identify additional members of this group.

METHODS. Fiber cells were harvested from various layers of the lens by laser micro-dissection and subjected to microarray, in situ hybridization, and Western blot analysis.

RESULTS. Expression of Livin, a member of the inhibitor of apoptosis protein (IAP) family encoded by *Birc7*, was strongly upregulated in deep cortical fiber cells. The depth-dependent distribution of Livin mRNA was confirmed by quantitative PCR and in situ hybridization. The onset of Livin expression coincided with loss of organelles from primary fiber cells. Livin expression peaked at 1 month but was sustained even in aged lenses. Antibodies raised against mouse Livin labeled multiple bands on immunoblots, reflecting progressive proteolysis of the parent molecule during differentiation. Mice harboring a floxed *Birc7* allele were generated and used to conditionally delete *Birc7* in lens. Lenses from knockout mice grew normally and retained their transparency, suggesting that Livin does not have an indispensable role in fiber cell differentiation.

CONCLUSIONS. *Birc7* is a late fiber gene of the mouse lens. In tumor cells, Livin acts as an antiapoptotic protein, but its function in the lens is enigmatic. Livin is a RING domain protein with putative E3 ubiquitin ligase activity. Its expression in cells bordering the OFZ is consistent with a role in organelle degradation, a process in which the ubiquitin proteasome pathway has been implicated previously.

Keywords: fiber cell differentiation, organelle degradation, apoptosis, hypoxia, Livin

Lens fiber cells are formed continuously by differentiation of cells at the edge of the lens epithelium. Newly formed cells are aligned on top of existing fiber cells, stacked like planks of wood in a lumber yard. The cell stacks (also referred to as radial cell columns¹) arise deep in the tissue and extend to the surface of the lens. There is no cell turnover in the fiber cell compartment. Consequently, each stack contains a complete cellular record, with the oldest fibers at the base and the youngest on the top.

All fiber cells are transparent and share an elongated prismatic shape. This has sometimes led to the assumption that they are a homogeneous population. In fact, depending on the relative positions of fibers in the cell stack, differences in biochemical or cell biological properties can be stark.² For example, the basal tips of superficial fiber cells make direct contact with the posterior lens capsule. In contrast, older cells, located farther down the stack, have detached from the capsule and instead form connections with other fiber cells, in a region called the lens suture. Deep in the stack, fiber cells are deprived of oxygen, due to consumption by overlying cells.³ Consequently, the proportion of cellular ATP derived from oxidative phosphorylation differs markedly between the hypoxic inner cell layers and the relatively oxygenated outer fibers.⁴ Finally, a few hundred micrometers below the lens surface, fibers undergo programmed organelle destruction, whereby the nucleus and other membranous organelles are rapidly and synchronously degraded.⁵ The dissolution of the organelles ensures the removal of light-scattering structures and results in the formation of the central, organelle-free zone (OFZ) of the

lens.⁶ Unlike the synthetically active surface cells, fiber cells within the OFZ lack the capacity for de novo protein synthesis.⁷ Based on such considerations it is likely that the biochemical differences between fibers located at different depths in the cell stack are as profound as those that exist between fiber cells and epithelial cells.

Once fiber cells have begun the process of terminal differentiation, gene expression is adjusted continuously, so that the cell biology of a fiber is appropriate to its position in the stack. Of particular interest are genes expressed in cells bordering the OFZ. This set of “late fiber genes” is likely to include genes associated with the organelle degradation process or stabilizing the fiber cell phenotype. Founder members of this group of late fiber genes include *Hopx*,⁸ *Capn3*,⁹ *Dnase2b* (*DLAD*),^{10,11} and *lengsin* (*Igsn*).¹² *Hopx* is expressed in fiber cells that have lost their direct connection to the lens capsule.⁸ In keratinocytes, *Hopx* is expressed in the suprabasal layer, where it may control expression of differentiation-dependent genes. *Capn3* encodes a calcium-activated endopeptidase, involved in remodeling of the spectrin cytoskeleton.¹³ *Dnase2b* is a lysosomal nuclease.¹¹ It has been shown to mediate chromatin breakdown during organelle loss and its absence results in cortical cataract due to incomplete DNA removal.¹⁰ *Lengsin* is expressed in cells during the membrane remodeling phase and interacts directly with the intermediate filament proteins Cp49 and vimentin, perhaps facilitating cytoskeletal reorganization.¹²

To identify other late fiber genes, we used laser microdissection to harvest cells from various layers of the lens for

comparative transcriptional analysis. We sought to identify genes that were expressed immediately prior to organelle breakdown. In this report, we characterized the expression of one such transcript, Livin (encoded by *Birc7*), which was strongly expressed in cells bordering the OFZ. Livin is an antiapoptotic protein (IAP) with suspected roles in progression of melanoma¹⁴ and other malignancies.¹⁵

METHODS

Mice

Wild-type (C57BL/6J) mice were obtained from a commercial supplier (Jackson Lab, Bar Harbor, ME, USA). Mice harboring a floxed *Birc7* allele were generated by homologous recombination (Supplementary Fig. S1). Two independent transgenic lines, LeCre¹⁶ and MLR10,¹⁷ expressing Cre recombinase in lens, were used for tissue-specific inactivation of *Birc7*. A germline knockout of *Birc7*¹⁸ (generated while these studies were in progress) was obtained and included in the analysis. To examine the effect of *Birc7* inactivation on lens cell architecture, *Birc7*-null mice were crossed with mT/mG reporter mice ([B6.129(Cg)-*Gt(ROSA)26Sor*^{tm4(ACTB-tdTomato,-EGFP)Luo}/J]tdTomato). The mT/mG mice (Jackson Lab) express tandem dimer Tomato (tdTomato) in cell membranes throughout the body,¹⁹ allowing cell morphology to be imaged in living intact lenses. All procedures were approved by the Washington University Animal Studies Committee and performed in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research.

Laser Microdissection and Microarray Analysis

Three tissue samples were isolated from 1-month-old lenses: an epithelial sample (EP; obtained by manually dissecting the epithelium from the lens), nucleated outer cortical fiber cells (OC; comprising fiber cells located 0–100 μ m beneath the lens surface), and nucleated inner cortical cells (IC; comprising cells located 100–200 μ m beneath the surface). The inner cortical cell sample directly abutted the OFZ, the border of which, in 1-month-old mice, is located approximately 200 μ m beneath the equatorial lens surface. The samples of OC and IC were obtained by laser microdissection. For that purpose, enucleated globes were embedded in optimal cutting temperature (OCT) medium (TissueTek; Miles, Elkhart, IN, USA) and frozen in 2-methylbutane over dry ice. Sections (20- μ m thick) were cut on a cryostat and mounted on glass RNase-free PEN-Membrane Slides (Leica Microsystems, Buffalo Grove, IL, USA). Sections were washed in RNase-free water for 30 seconds, fixed in 70% ethanol for 1 minute, and air dried for 10 minutes. The samples of OC and IC were laser-microdissected (LMD6000, Leica Microsystems) and collected directly into the lysis buffer supplied with the RNA isolation kit (RNeasy Micro Kit; Qiagen, Valencia, CA, USA). We quantified RNA using a commercial assay (Quant-iT RNA; Invitrogen, Grand Island, NY, USA). Synthesis of cDNA was accomplished using an amplification system (NuGEN Ovation Pico WTA System; NuGEN Technologies, Inc., San Carlos, CA, USA). Independent quadruplicate replicates were applied to bead microarrays according to manufacturer instructions (Illumina Mouse 6; Illumina, San Diego, CA, USA). These experiments were modeled on published methodology.²⁰

Endpoint PCR

Two splice variants (α and β) of Livin have been described, differing by a stretch of 18 amino acids located between the BIR and RING domains.²¹ Transcripts corresponding to the α and β forms were differentiated by endpoint PCR using the following primers:

Forward: 5' TCCTCTGGGGCTGCCTTCCTT 3' (349-369)

Reverse: 5' AGGCACCCTGCAGATAGGGCA 3' (930-949)

The predicted amplicon included exon 6, which contains the alternatively spliced region.

RT-qPCR

For real-time quantitative PCR (RT-qPCR), cDNA was synthesized with 1- μ g lens RNA as a template using a reverse transcription supermix for RT-qPCR (iScript; BioRad, Hercules, CA, USA). The following primers were used:

Gapdh-F 5' CCTCGTCCCGTAGACAAAATG 3'

Gapdh-R 5' TGAAGGGGTCGTTGATGGC 3'

Ppia-F 5' GCGGCAGGTCCATCTACG 3'

Ppia-R 5' GCCATCCAGCCATTCAGTC 3'

Birc7-F 5' GGGGAGCCTGCCTTCCTCTGA 3'

Birc7-R 5' AGGGGTCATCCCCACGCTCC 3'

Experiments with RT-qPCR were designed according to the MIQE guidelines,²² using a supermix according to manufacturer instructions (SsoFast EvaGreen; BioRad). Each biological sample was obtained from a pool of five different animals, unless otherwise specified, and three technical replicates were performed per sample. Reference genes (*Gapdh* and *Ppia*) were selected using Normfinder²³ software and verified by determining the M value. Efficiency of RT-qPCR was determined for all primers and was within 93% to 102% range.

Northern Blot

The size and distribution of Livin mRNA transcripts were evaluated by Northern blotting, as described.⁷ Total RNA (1 μ g) was separated on denaturing gels and transferred to nitrocellulose. Blots were probed with digoxigen-labeled sense and antisense riboprobes (spanning region 579–811 of the Livin transcript) prepared using Dig RNA labeling mix (Roche Diagnostics, Indianapolis, IN, USA).

In Situ Hybridization

In situ hybridization was performed as described.²⁴ Briefly, 4- μ m thick, midsagittal sections of paraffin-embedded lens tissue were prepared. A target probe set raised against the Livin transcript was obtained from Advanced Cell Diagnostics (ACD, Hayward, CA, USA). The probe set consisted of 20 pairs of oligonucleotides spanning a \approx 1-kb contiguous region of the target mRNA transcript (NM_001163247.1: 2-1141). In situ hybridization was performed using a multiplex assay (RNAscope 2.0; ACD). Following proprietary preamplification and amplification steps, target probes were detected using an alkaline-phosphatase-conjugated label probe with Fast Red as substrate and Gill's hematoxylin as counterstain. As negative controls, some sections were hybridized with target probe against *DapB*, a bacterial gene. Alternatively, the Livin probe set was hybridized to lens sections from *Birc7*-null mice.

Antibodies and Immunoblotting

Two independently generated antibodies against mouse Livin were used for immunofluorescence and Western blotting applications. A recombinant protein corresponding to amino acid positions 22 through 285 of mouse Livin and containing an N-terminal histidine tag was generated in *Escherichia coli* and used to immunize rabbits (PrimmBiotech, West Roxbury, MA, USA). Anti-Livin was purified from the antiserum using CNBr-sepharose affinity column chromatography. The second antibody (clone 7H5.1.1-IgG2a) was raised against mouse Livin and characterized in an earlier study.¹⁸ Both antibodies

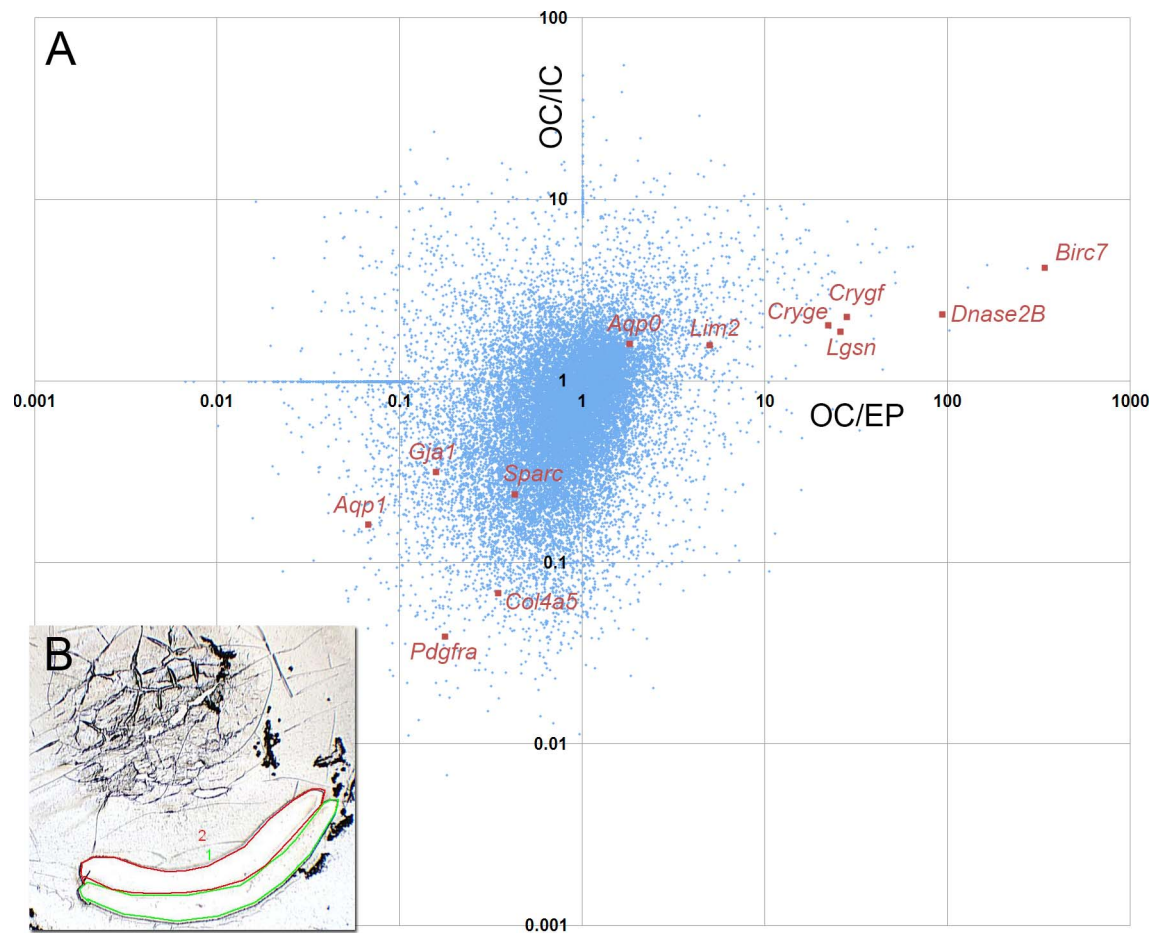


FIGURE 1. Microarray analysis of depth-dependent gene expression in 1-month-old mouse lenses. **(A)** Scatterplot of microarray data showing relative levels of gene expression in EP cells, OC and IC fiber cells. The quadrants define four distinct patterns of lens gene expression. The *lower left quadrant*, for example, contains transcripts that are highly expressed in the epithelium (with respect to either of the fiber samples). Epithelial specific transcripts, *Aqp1*, *Pdgfra*, *Gja1*, *Sparc*, and *Col4a5* are located in this quadrant. Transcripts that show upregulated expression in both the early and late stages of fiber differentiation are shown in the *upper right quadrant*. Members of this group include genes known to be expressed predominantly in fiber cells, such as *Aqp0*, *Lim2*, and the gamma crystallins *Cryge* and *Crygf*. Among strongly upregulated genes in this quadrant are the late fiber genes *Dnase2B* and *Lgsn*. The most upregulated transcript, with expression levels several hundred-fold higher in the inner cortical fibers than in the epithelium, is *Livin*. The other two quadrants contain transcripts that are transiently upregulated (*lower right quadrant*) or downregulated (*upper left quadrant*) during early fiber cell differentiation. **(B)** The *inset* shows an example of a frozen lens section from which the OC (*green*) and IC (*red*) samples were collected by laser microdissection.

recognized recombinant *Livin* and endogenous *Livin* on Western blot and exhibited little or no immunoreactivity on lens samples from *Birc7*-null mice. Western blotting was performed as described previously.²⁵ *Livin* antibodies were used at 1:500 dilution. To examine *Livin* expression in various strata of the tissue, lenses were progressively solubilized and fractions collected for analysis, as described.⁹

Immunofluorescence

To visualize the expression of *Livin* protein, P1 mouse lenses were fixed for 2 hours in 4% PBS, washed and sectioned at 150 μ m on a tissue slicer (Vibratome series 1000; TPI, Inc., St. Louis, MO, USA). Lens slices were permeabilized in 0.1% Triton X-100/PBS for 30 minutes and blocked with 1% bovine serum albumin/10% normal goat serum in PBS. Slices were incubated overnight in anti-*Livin*, washed in PBS, and incubated for 2 hours in secondary antibody. Draq5 (Invitrogen) was included in the washing solution as a nuclear counterstain. Slices were coverslipped and examined using a confocal microscope (LSM510, Carl Zeiss, Inc., Thornwood, NY, USA). To visualize the fate of cytoplasmic organelles, paraffin sections were

incubated with an antibody against protein disulfide isomerase (PDI; an ER resident protein), as described.²⁶

Multiphoton Imaging of Fiber Cell Structure

Wild-type or *Birc7*-null mice were crossed with a reporter strain (mT/mG) expressing membrane targeted TdTomato. Lenses were dissected from the eye and incubated in warm artificial aqueous humor solution.³ Image stacks were collected using a multiphoton microscope (Fluoview1000 MPE; Olympus, Center Valley, PA, USA) using an excitation wavelength of 785 nm and an XLPlan N \times 25 water immersion objective lens (1.05 NA).

RESULTS

Tissue samples from the lens epithelium (EP), outer cortex (OC), and inner cortex (IC) were harvested manually (EP) or by laser microdissection (OC, IC) and relative levels of gene expression in the three samples were determined by microarray analysis (Fig. 1). On this analysis, mRNA products of

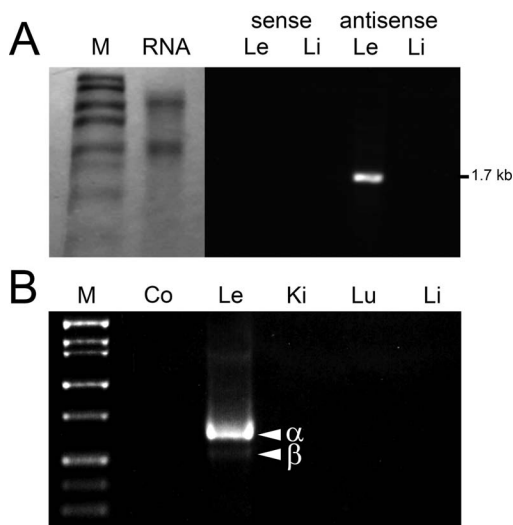


FIGURE 2. Expression of Livin mRNA in mouse tissues. **(A)** Northern blot analysis identifies a 1.7-kb transcript in total RNA extracted from lens (Le) but not liver (Li). Sense control probe does not hybridize to RNA from either tissue. **(B)** Two amplicons (α and β) are detected following endpoint PCR amplification of lens cDNA with Livin-specific primers. The larger and more abundant amplicon corresponds to Livin α and the smaller amplicon to Livin β . The identity of the amplicons was confirmed by sequencing. Co, negative control (lacking template DNA); Ki, kidney; Lu, lung; M, marker.

known late fiber genes (such as *Dnase2B* and *Igsn*) segregated with transcripts (*Cryge* and *Crygf*) characterized by enhanced expression in the OC and IC samples. Surprisingly, the most upregulated of this group was Livin, which was undetectable

in the EP sample, but expressed strongly in the OC sample and even more so in the IC sample. Livin (encoded by *Birc7*) is a member of the IAP family.²⁷ Previous studies have shown Livin to be expressed in certain types of melanoma cells as well as some fetal tissues.²⁸

Expression of Livin mRNA in lenses from 1-month-old mice was evaluated by Northern blot (Fig. 2). A prominent 1.7-kb RNA transcript was detected in RNA extracted from lens but not liver (Fig. 2A). Two alternative transcripts (α and β) of Livin have been described²¹ and, by endpoint PCR, both were detected in lens (Fig. 2B). Livin α , the larger of the transcripts, was the more abundant. Endpoint PCR failed to detect Livin expression in kidney, lung, or liver samples.

We used RT-qPCR to quantify Livin expression in lens and other tissues (Fig. 3). Livin was strongly expressed in lens but not liver, lung, or retina (Fig. 3A). The mouse lens forms at \approx day 11 of embryonic development (E11), but Livin expression was first detected at E16.5 (Fig. 3B), the stage at which organelle degradation commences in the innermost fiber cells.²⁹ Livin expression increased in the lens during the early postnatal period, peaking in 1-month-old lenses (Fig. 3B). Expression was sustained in the maturing lens and the Livin transcript was readily detected even in lenses from 18-month-old mice (the oldest examined). In laser microdissected samples from 1-month-old lenses, Livin expression was undetectable in epithelial samples (Fig. 3C) and \approx 4-fold greater in inner cortical cells than outer cortical cells, substantiating the microarray results (see Fig. 1).

To visualize the expression of Livin mRNA at the single cell level within the developing mouse lens, paraffin sections were hybridized with a target probe set raised against a 1-kb region of the Livin mRNA transcript (Fig. 4). At the first stage examined, E12.5, the lens consisted of a hollow ball of cells (the lens vesicle). The posterior wall of the vesicle comprised

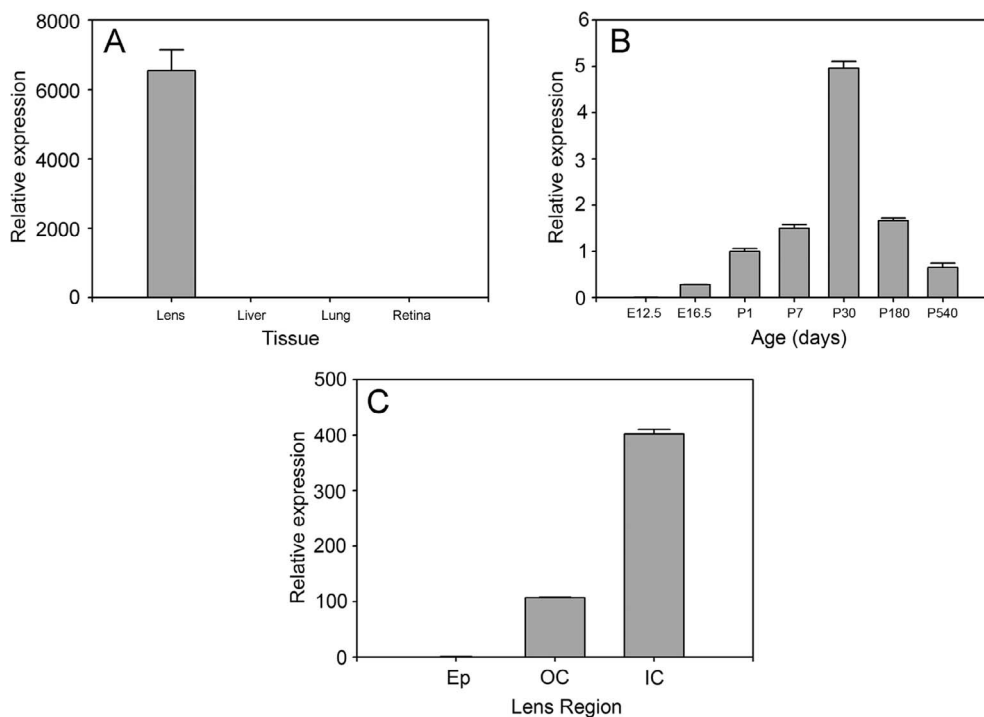


FIGURE 3. Quantification by RT-qPCR of Livin expression in mouse tissue. **(A)** In 1-month-old mice, Livin is strongly expressed in lens compared with other tissues (liver, lung, retina) examined. **(B)** Livin expression in the lens commences between E12.5 and E16.5. Expression increases throughout the early postnatal period, peaking at postnatal (P) day 30. Although expression levels are somewhat reduced in older animals, Livin is an abundant transcript at all ages. **(C)** Within the lens, Livin expression is undetectable in the epithelium, and strongest in the inner cortical fiber cells. Data represent mean and standard deviation of at least three determinations in each case.

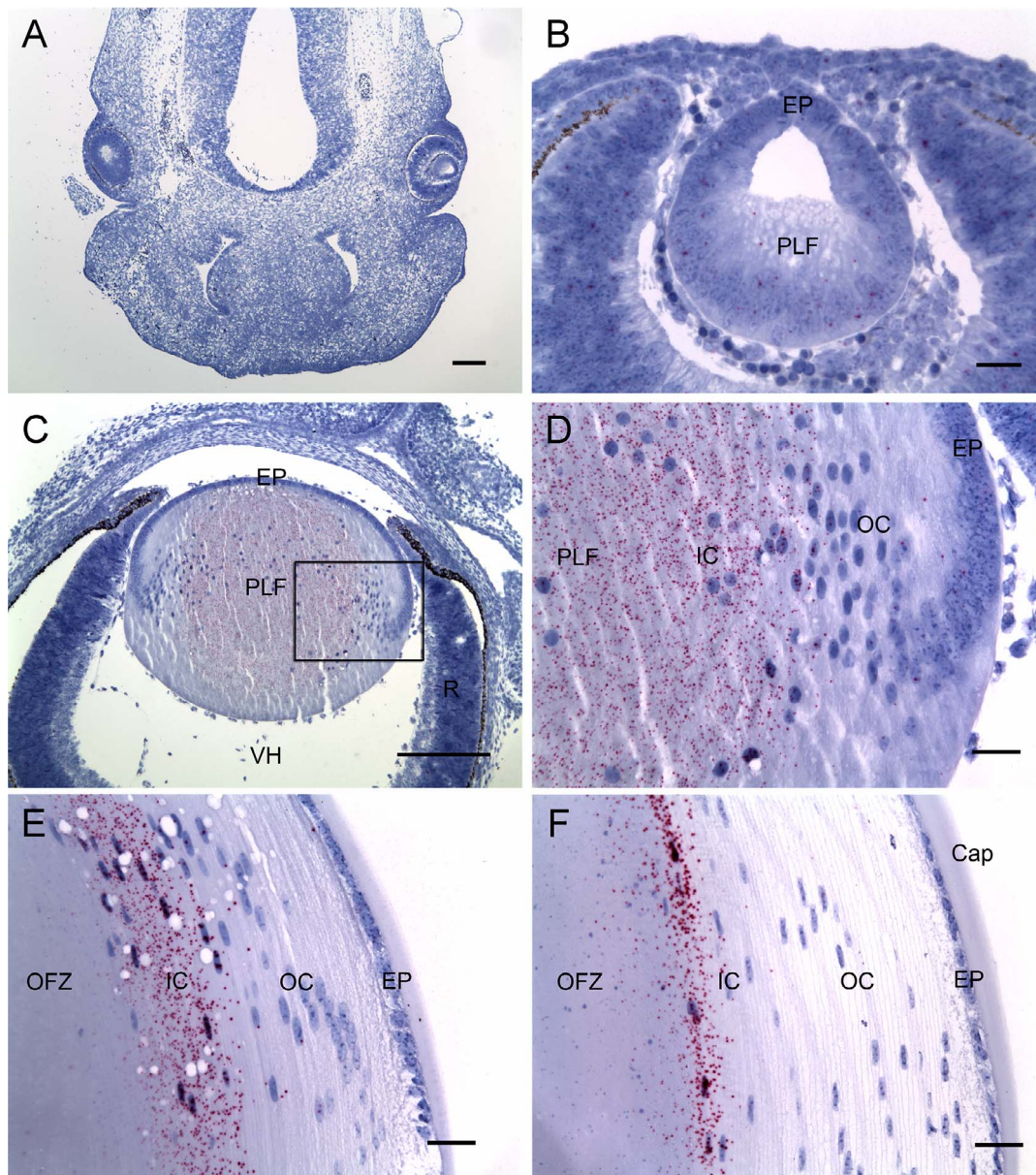


FIGURE 4. In situ hybridization analysis of Livin mRNA expression in the developing mouse eye. (A) At day 12.5 of embryonic development, Livin expression (*red dots*) is undetectable in the embryonic head. (B) The lens, at this stage a hollow vesicle, is composed of epithelial cells (EP) and primary lens fibers (PLF) in the process of elongation. Livin expression in the E12.5 lens is barely above background. (C, D) At day 16.5 of embryonic development, Livin is expressed strongly by primary fiber cells located in the center of the lens. The boxed region in (C) is shown at higher magnification in (D). Livin expression is undetectable elsewhere in the eye. (E) At postnatal day 30, Livin expression is restricted to nucleated IC fiber cells bordering the central OFZ. (F) At age 6 months, Livin expression is restricted to a thin layer of fiber cells adjacent to the OFZ. Cap, capsule; R, retina; VH, vitreous humor. Scale bars: (A, C) 200 μ m, (B, D-F) 25 μ m.

the primary lens fibers, which at that stage were in the process of elongating. At E12.5, Livin expression was low throughout the embryonic head, including the developing eye and lens (Figs. 4A, 4B). However, by E16.5, Livin mRNA was readily detected in nucleated primary fiber cells located in the center of the lens (Figs. 4C, 4D). Thus, expression of Livin immediately preceded organelle breakdown, which commences in the central fiber cells during the late embryonic period.²⁹ Elimination of organelles from the innermost fiber cells leads to the formation of the OFZ.⁵ The process of organelle degradation continues postnatally. At age 1 or 6 months, Livin was expressed by a contingent of cortical fiber located adjacent to the OFZ (Figs. 4E, 4F). Thus, at all ages, Livin

was expressed by fiber cells shortly before organelle degradation began. Transcripts were not detected within the anucleated cells of the OFZ nor in the anterior lens epithelium.

Livin was originally cloned from a melanoma cell line.²⁸ To compare the pattern of Livin expression in lens and melanoma, histopathology samples of human uveal melanoma were examined. Livin mRNA distribution within the tumor was visualized by in situ hybridization, using a probe set raised against human Livin (Fig. 5). Livin was detected in four of six melanoma specimens examined. Significantly, in Livin-positive tumors, expression was restricted to the core of the tumor. Melanoma cells in the superficial region of the tumor (Fig. 5A) or regions adjacent to vascular elements (Fig. 5B) did not

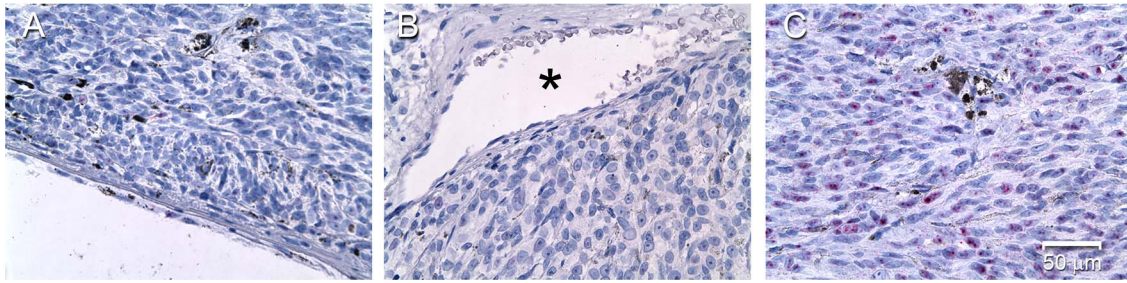


FIGURE 5. In situ hybridization analysis of Livin expression in human uveal melanoma. Expression is undetectable in the superficial (A) region of the tumor (at the vitreous surface) or around vascular (*) elements (B). Marked Livin expression (red dots) is detected only in the core of the tumor (C). Scale bar: 50 μ m.

express Livin. Thus, expression of Livin in melanoma samples recapitulated its expression pattern in lens, where the Livin transcript was restricted to the inner cell layers.

To visualize the expression of Livin protein, antibodies were raised against recombinant mouse Livin and used to probe Western blots of whole lens samples prepared from wild-type or *MLR10^{tg/-};Birc7^{fllox/fllox}* mice (the latter serving as a negative control). Several immunopositive bands were detected in lens samples. The largest and most prominent band had an apparent molecular mass of \approx 41 kDa (Fig. 6A) and likely corresponds to full-length Livin. However, the presence of additional immunopositive bands with apparent masses of \approx 37, \approx 34, and \approx 22 kDa was noted and a diffuse band of \approx 30 kDa was also present. Immunopositive bands were not detected in lens samples prepared from *MLR10^{tg/-};Birc7^{fllox/fllox}* mice, implying that antibody labeling was specific and that the 22- to 41-kDa bands corresponded to authentic Livin splice variants or posttranslationally modified species. To follow the fate of Livin protein during fiber cell differentiation and aging, lenses were progressively solubilized (Fig. 6B). This semiquan-

titative technique allows fractions to be collected from progressively deeper layers of the lens, although the precise spatial relationships between the fractions are uncertain (it is unlikely, e.g., that fractions are derived from strata of equal thickness). The multiple immunopositive bands observed in whole-lens lysates (Fig. 6A), were evident even in the outermost fractions (containing the youngest fiber cells), suggesting that the lower molecular weight bands were not simply proteolytic fragments generated during cellular aging. However, depth-dependent changes in Livin expression were noted. For example, the putative full length (41 kDa) form was not detected in the deeper fractions and no immunopositive bands were detected in the innermost fiber cells.

To better localize Livin protein expression, we used confocal immunofluorescence microscopy of vibratome sections prepared from early postnatal lenses (Fig. 7). Analysis of sagittal sections from P1 lenses revealed that Livin expression was first evident in fiber cells that had recently detached from the lens epithelium (Fig. 7A). In equatorial sections, Livin immunofluorescence formed an annular pattern (Fig. 7B). Livin

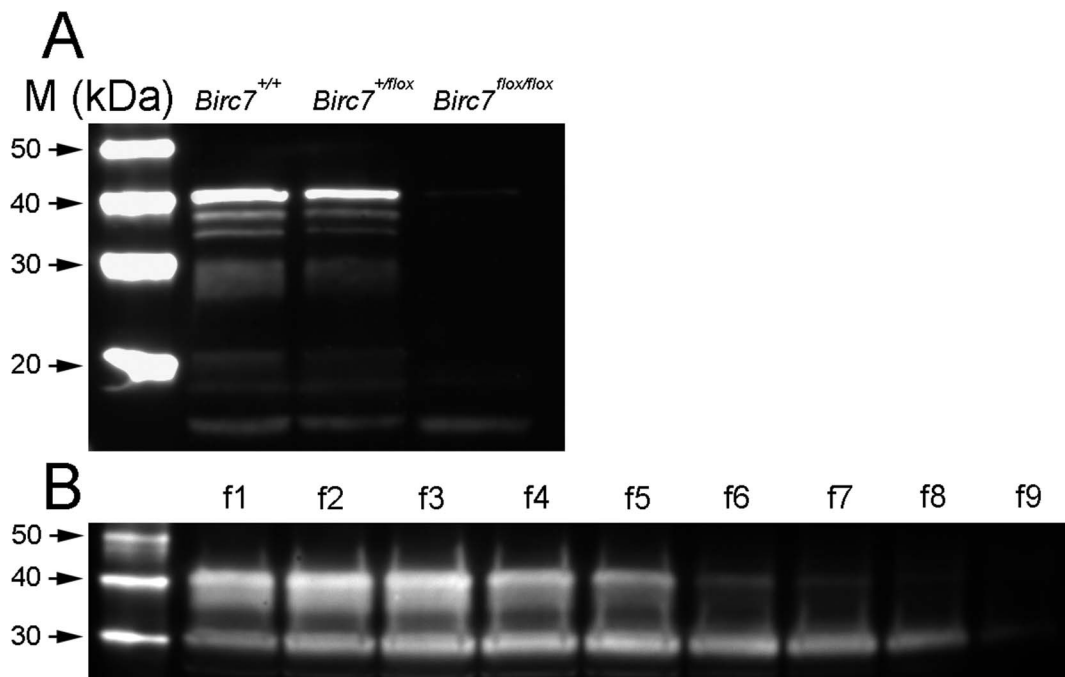


FIGURE 6. Western blot analysis of Livin protein expression in the mouse lens. (A) Several immunopositive bands are present in lens samples from wild-type mice, or mice heterozygous for the floxed allele. To conditionally delete *Birc7* in the lens, *Birc7^{fllox/fllox}* animals were crossed with *MLR10* mice. No immunopositive bands were detected in the conditional null lenses. (B) Progressive solubilization of lenses allowed the collection of nine fractions, corresponding with various strata of the lens. In the outmost fractions (f1–f5) two immunopositive bands (at \approx 40 and \approx 30 kDa) are present. The larger band is absent in fractions from the inner fiber cells (f6–f8). Neither band is detected in the innermost fraction (f9).

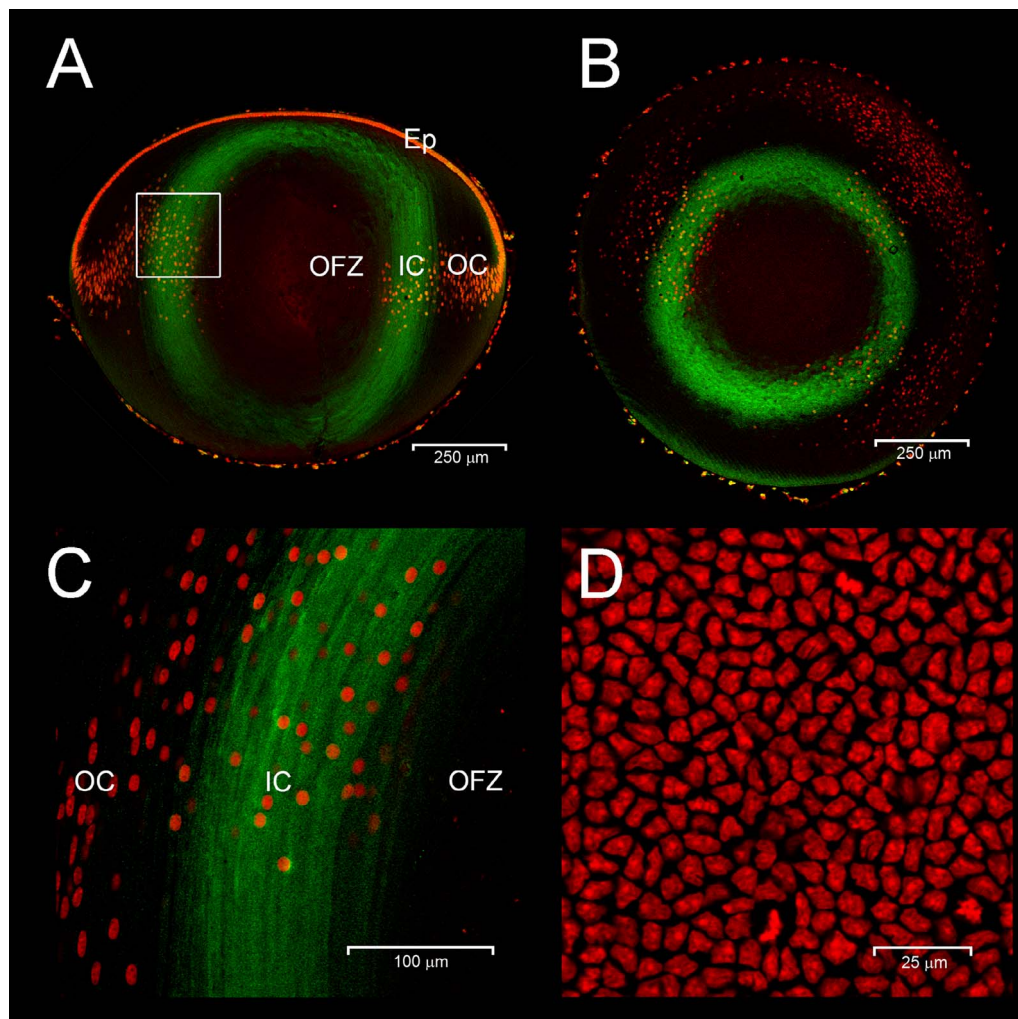


FIGURE 7. Livin expression in P1 mouse lens. Sagittal (A) and equatorial sections (B) show Livin expression (green) in fiber cells in the IC region. The boxed region in (A) is shown at higher magnification in (C). The disappearance of fiber cell nuclei (red) marks the border of the OFZ. Note that Livin is expressed by fiber cells immediately prior to denucleation. In expressing fiber cells, Livin immunofluorescence is distributed throughout the cytoplasm. En face views of the lens epithelium (D) demonstrate the absence of Livin expression in this cell population. Scale bars: (A) 250 μm , (B) 250 μm , (C) 100 μm , (D) 25 μm .

protein was detected in the deeper layers of nucleated IC fiber cells, located just outside the central OFZ (Fig. 7C). In expressing cells, Livin protein was diffusely distributed throughout the cytoplasm (Fig. 7C). Expression was undetectable in the overlying epithelium (Fig. 7D) consistent with the results of RT-qPCR analysis (Fig. 3C).

Fiber cells are among the longest lived cells in the body. Based on its expression pattern and putative antiapoptotic role in other systems, we hypothesized that Livin might contribute to lens cell longevity by suppressing cell death in the hypoxic core of the tissue. Alternatively, Livin, which is expressed by cells as they approach the OFZ (Fig. 7C), might have a role in organelle degradation. To test these hypotheses directly, we generated mice carrying a floxed *Birc7* allele (see Supplementary Fig. S1), *Birc7^{fllox/fllox}* mice were crossed with either *LeCre¹⁶* or *MLR10¹⁷* mice to conditionally delete *Birc7* in the lens. The *LeCre* and *MLR10* strains express Cre recombinase in lens and have been widely used to conditionally delete genes in this tissue. The two strains differ primarily in the timing of the onset of lens Cre expression (E8.75 for *LeCre* and E10.5 for *MLR10*). Since our data suggested that Livin expression does not commence until after E12.5 (Fig. 3B), we anticipated that

the efficacy of the two lines (and, therefore, any resulting phenotypes) should be broadly similar. Unexpectedly, there was a striking qualitative difference in the lens phenotypes between *LeCre¹⁶;⁻Birc7^{fllox/fllox}* and *MLR10¹⁶;⁻Birc7^{fllox/fllox}* animals. The eyes of *LeCre¹⁶;⁻Birc7^{fllox/fllox}* mice were invariably (16/16) smaller than wild type (Fig. 8). Histologic analysis revealed that the microphthalmic phenotype was associated with the presence of a small and severely disrupted lens (Supplementary Fig. S2). Fiber cells in the *LeCre¹⁶;⁻Birc7^{fllox/fllox}* mice were heavily vacuolated and cell death assays revealed the presence of numerous apoptotic cells in the central lens fibers, as early as E16 (data not shown). These data were consistent with the hypothesis that Livin expression actively suppressed cell death in the hypoxic lens interior. The majority (12/16 or 75%) of lenses from mice heterozygous for the floxed allele exhibited a similar lens phenotype. Surprisingly, however, analysis of lenses from *LeCre¹⁶;⁻Birc7^{+/+}* littermate controls revealed a relatively high incidence (12/32 or 37%) of microphthalmia. In affected animals, the lens phenotype was often indistinguishable from that observed in the conditional knockouts. Thus, at least on the mixed genetic background used here, expression of the Cre transgene in the *LeCre* strain

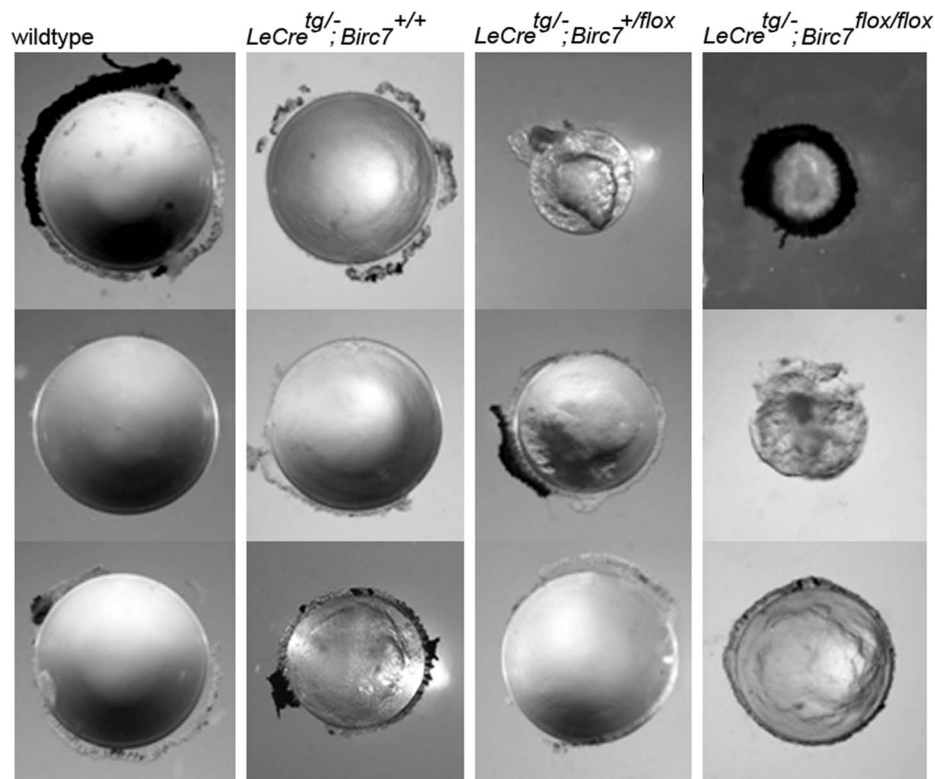


FIGURE 8. Conditional deletion of Livin in the lens using the *LeCre* transgene results in small and disrupted lenses. Compared with wild-type lenses from 1-month old animals (*left column*), lenses from mice homozygous for the floxed *Birc7* allele (*right column*) were severely disrupted, with 100% penetrance. However, a similar lens phenotype was also observed in 37% of control (*LeCre*^{tg/-};*Birc7*^{+/+}) littermates. A phenotype was observed in 75% of lenses from mice that were heterozygous for the floxed allele and hemizygous for the *LeCre* transgene.

produced a phenotype independent of the presence of the floxed *Birc7* allele. In contrast, no overt lens phenotype was observed in *MLR10*^{tg/-};*Birc7*^{flox/flox} animals (Fig. 9). We confirmed that *Birc7* was efficiently disrupted (as evidenced by absence of Livin protein) in both *LeCre*^{tg/-};*Birc7*^{flox/flox} and *MLR10*^{tg/-};*Birc7*^{flox/flox} mice (Supplementary Fig. S3).

We examined lenses from *MLR10*^{tg/-};*Birc7*^{flox/flox} mice for the presence of cataracts or growth defects. Histological analysis indicated that the lens cellular architecture was indistinguishable from wild type (Supplementary Fig. S4). Lenses retained their clarity at all ages examined (up to 12 months; data not shown) and careful measurements of lens size failed to detect any significant growth delays (Fig. 10). To substantiate findings obtained with the *MLR10* mice, we also examined lenses from germline *Birc7*-null mice¹⁸ that became available in the course of our studies. Lenses from *Birc7*-null animals were indistinguishable from *MLR10*^{tg/-};*Birc7*^{flox/flox} lenses (data not shown).

In wild-type lenses, the observation that Livin was expressed in cells near the border of the OFZ (Fig. 7), suggested a potential role for Livin in organelle degradation. To test this hypothesis, lenses from *MLR10*^{tg/-};*Birc7*^{flox/flox} mice and control littermates were processed for immunofluorescence and labeled with antibodies against PDI, a marker for the endoplasmic reticulum (ER). Previous studies have shown that the ER is rapidly degraded at the edge of the OFZ.²⁶ The disappearance of PDI staining at the border of the wild-type OFZ was confirmed in the current study. The OFZ also formed normally in *MLR10*^{tg/-};*Birc7*^{flox/flox} mice (Fig. 11), indicating that Livin is not necessary for the removal of the endoplasmic reticulum (nor the fiber cell nuclei; see Supplementary Fig. S4).

Previous studies have shown that the lens can be surprisingly resilient to the inactivation of even relatively

abundant genes. Although deletion of membrane³⁰ or cytoskeletal proteins³¹ does not necessarily result in loss of transparency it often disrupts the organization of the fiber cell stacks. To examine whether the absence of Livin affected the

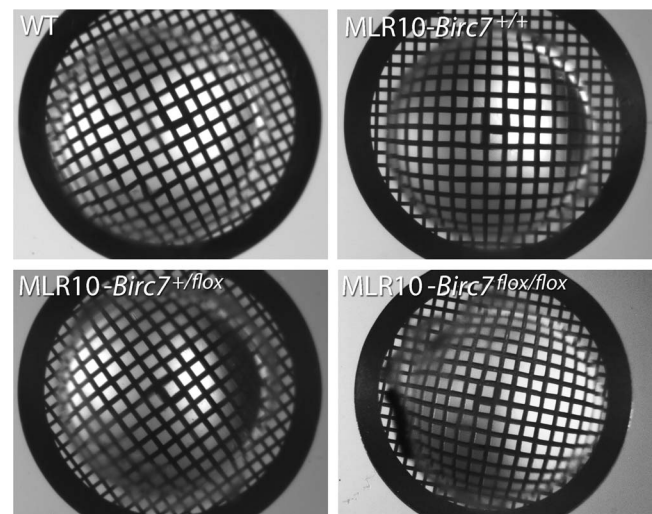


FIGURE 9. Lens clarity and refractive properties are unaffected in Livin knockout lenses. We used *MLR10* to conditionally delete *Birc7* in the lens. Lenses from 1-month-old wild-type or conditional knockout animals were photographed against a grid pattern. The optical properties of control animals (wild-type or mice hemizygous for the *MLR10* transgene) did not differ noticeably from mice in which the *Birc7* locus was partially (*Birc7*^{+/flox}) or completely (*Birc7*^{flox/flox}) inactivated.

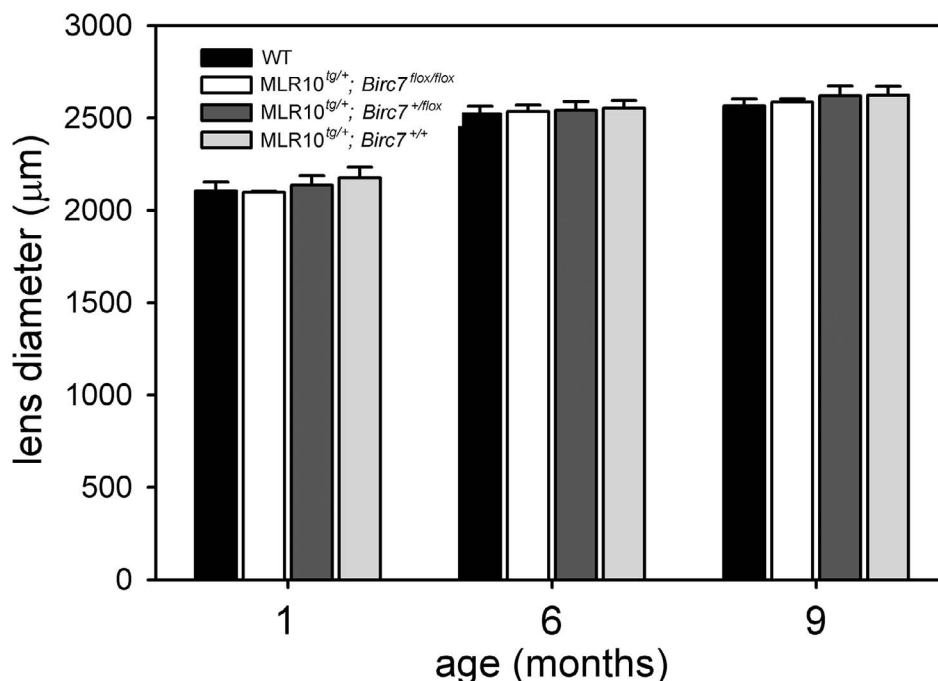


FIGURE 10. Lens growth is not affected by the absence of Livin. No significant differences in the equatorial diameter of the lens were noted among the various genotypes. Data represent mean values \pm SD ($n = 4$ in each case).

cytoarchitecture of the lens, we used a novel multiphoton imaging technique. For this application, we utilized *Birc7*-null mice,¹⁸ which were crossed with the mT/mG reporter strain. In mT/mG mice, membrane-targeted TdTomato is expressed in cells throughout the body, including the lens. Endogenous expression of TdTomato in the plasma membranes of all fiber cells allowed fiber morphology to be imaged in three dimensions in intact, living lenses. Using a long working distance water immersion lens, lens fiber cells were visualized at depths of up to 500 μ m below the equatorial lens surface. This allowed cells to be inspected in the EP, OC, IC, and OFZ regions (i.e., before, during, and after the period of Livin expression in wild-type lenses). Image stacks collected from 1-month-old wild-type or *Birc7*-knockout lenses were indistinguishable (Fig. 12). In each case, fiber cells were organized into radial cell columns and the cells had characteristic shapes that varied with depth, as described previously.⁶ Thus, the absence of Livin was not associated with a gross disturbance in the cellular organization of cortical lens fiber cells.

DISCUSSION

Microarray analysis of laser microdissected lens tissue was used to screen for genes expressed late in the fiber cell differentiation program. A similar strategy was employed previously by Ivanov et al.,²⁰ who identified several genes with modulated expression in mature fibers. In the current work, we confirmed the expression of previously identified late fiber genes, including *Igsn* and *Dnase2B*. However, in our study, the most upregulated transcript in inner cortical fibers was Livin, encoded by the *Birc7* gene. Livin expression has been noted previously in human lens samples, as part of the expressed sequence tag NEI bank project,³² and in a recent RNA-Seq study,³³ but a detailed analysis of its expression and function in the lens has not been reported.

Livin is a member of the inhibitor of apoptosis family of proteins. These proteins are characterized by the presence of one or more BIR motifs (baculovirus inhibitor of apoptosis protein repeat) that interact directly with caspases to suppress

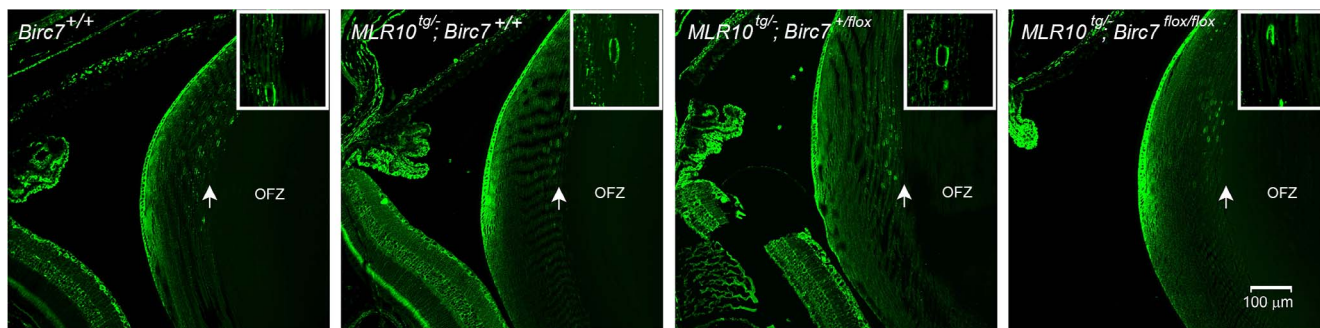


FIGURE 11. Degradation of the endoplasmic reticulum occurs normally in the absence of Livin. The distribution and fate of the endoplasmic reticulum was visualized with anti-PDI. In wild-type lenses, PDI immunofluorescence is strong in the lens epithelium and outer cortical fibers. The abrupt disappearance of PDI staining marks the boundary of the OFZ. The loss of PDI immunoreactivity in cells bordering the OFZ was indistinguishable in wild-type or conditional knockout lenses.

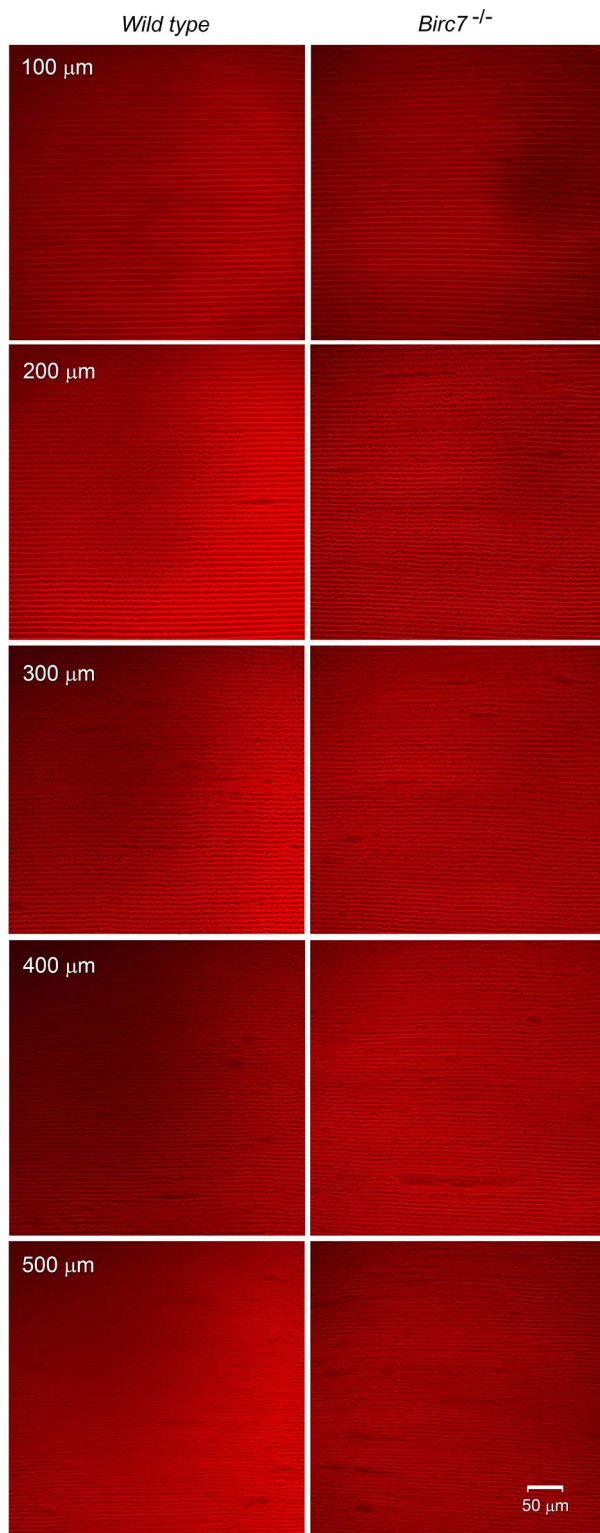


FIGURE 12. Multiphoton microscopic analysis of cellular organization in lenses from wild-type or *Birc7*-null animals. Crosses were established with mT/mG mice (a line in which the fiber cell membranes are endogenously labeled with TdTomato) to allow the morphology of individual cells located at various depths below the lens surface to be visualized. Individual optical sections are shown. Note that the border of the OFZ is located approximately 200 μm below the lens equatorial surface. There was no obvious difference in cell morphology or packing arrangement on either side of the OFZ border.

apoptotic cell death. Livin contains a single BIR domain and a RING-domain with putative E3 ubiquitin ligase activity. Livin was first isolated from malignant melanoma cells and its expression level in most nontransformed cell types is believed to be very low.³⁴ Its antiapoptotic action is believed to be mediated via inhibition of caspase-3, -7, and -9³⁴ and by its ability to promote the degradation of Smac/Diablo, an endogenous regulator of the IAP family.³⁵

Two splice variants (α and β) of Livin have been described.²¹ In the lens, the α variant dominates. In apoptotic cells, Livin is cleaved by caspases, generating a truncated version (tLivin) that redistributes to the Golgi apparatus.³⁶ Paradoxically, under some circumstances tLivin may promote cell death.³⁷ In the lens, Livin was evenly distributed throughout the cytoplasm. Several Livin species were observed by immunoblotting. Presumably the multiple bands represented full length and truncated α and β variants. Immunopositive bands were not detected in lens samples from *MLR10^{tg/-}*; *Birc7^{fllox/fllox}* mice, indicating that the wild-type labeling pattern represents authentic Livin immunoreactivity rather than nonspecific antibody binding. The identity of the various species and their functional significance, if any, has yet to be determined. Lens cells are retained throughout life and by isolating successive layers of tissue, it was possible to follow the fate of Livin protein during the fiber differentiation process. Such analysis indicated that, over time, full-length Livin disappeared and that the inner cortical fiber cells contained only the 30-kDa isoform. The progressive cleavage of lens proteins is a well-established phenomenon. For example, Aqp0, the most abundant integral membrane protein of the lens is extensively cleaved in the lens core³⁸ and the cytoskeletal protein alpha-spectrin, is cleaved to a 150-kDa fragment in the inner cortical region.³⁹ The protease(s) responsible for degrading full-length Livin in differentiating lens fiber cells is unknown but several proteases are activated in cells bordering the OFZ, including the calcium-activated protease, calpain 3.⁹

Livin expression began in primary lens fiber cells sometime between E12.5 and E16.5. It is noteworthy that elimination of cytoplasmic organelles in the innermost fiber cells begins in the period E15 to E18.²⁹ The strict spatial/temporal relationship between organelle loss and Livin expression was maintained in the secondary fiber cells of postnatal lenses, where Livin expression in the inner cortex immediately preceded organelle degradation. Although Livin expression is not essential for formation of the OFZ, it is possible that Livin has a role in organelle degradation. The ubiquitin proteasome pathway has been implicated in organelle degradation during formation of the OFZ.^{40,41} In the ubiquitin proteasome pathway, protein substrates are recognized by E3 ubiquitin ligases which mediate the transfer of one or more ubiquitin molecules to the substrate, thereby targeting it for degradation via the proteasome. Although the human genome encodes >600 RING domain-containing E3 ligases,⁴² the identities of the E3 ligases operating during organelle degradation have yet to be determined. The specific expression of Livin (a putative ubiquitin E3 ligase) in cells bordering the OFZ suggests that it may function in the process.

Livin was expressed specifically by the innermost nucleated fiber cells. In human uveal melanoma samples, the expression of Livin followed a broadly similar pattern, with low-level expression in the outer tissue layers and enhanced expression in the inner layers. Recent studies suggest that Livin is induced in the hypoxic interior of solid tumors through binding of hypoxia-inducible factor 1 α (Hif-1 α) to the Livin promoter.⁴³ A similar mechanism may operate in the avascular lens. The concentration of oxygen in the central region of the lens is extremely low, due to consumption by the overlying fiber

cells.³ Thus, during terminal differentiation, fiber cells necessarily traverse a steep standing gradient of oxygen. The inner cortical cells are the most hypoxic of all the nucleated cells in the lens. The concentration of oxygen in the interior of the mouse lens is unknown; but in bovine lens the partial pressure of oxygen (pO₂) in the central region is 1 to 2 mm Hg³ and in rabbit lenses ≈10 mm Hg.⁴⁴ For comparison, mean pO₂ values in human melanoma xenografts typically range from 1 to 10 mm Hg.⁴⁵ The role of Hif-1 α in the lens is not well defined, but conditional deletion of Hif-1 α is associated with widespread lens cell death.⁴⁶

To test the physiological role of Livin, we conditionally inactivated the *Birc7* locus by crossing mice carrying a floxed *Birc7* allele with either LeCre or MLR10 mice, two strains that express Cre recombinase in the lens. Unexpectedly, the resulting lens phenotypes depended on the Cre line that was used. *LeCre^{tg/-};Birc7^{fllox/fllox}* mice showed a highly penetrant lens phenotype, with vacuolated fiber cells and widespread apoptosis in the lens. However, a similar lens phenotype was also observed in many (37%) control (*LeCre^{tg/-};Birc7^{+/+}*) animals. Thus, on the mixed background used here, mice hemizygous for the *LeCre* transgene exhibited ocular phenotypes that were independent of the floxed *Birc7* allele. This finding is consistent with a recent report in which, depending on the genetic background, hemizygous LeCre mice developed severe eye abnormalities even in the absence of LoxP sites.⁴⁷ The precise mechanism remains to be determined but, presumably, reflects an interaction between the *LeCre* transgene and modifier genes present in certain backgrounds. The LeCre mouse has been widely used in developmental studies of the lens. The current findings, together with those of Dora et al.,⁴⁷ indicate that conclusions of earlier studies utilizing the LeCre strain should be interpreted with caution.

To control for the confounding effect of the *LeCre* transgene, we generated conditional knockouts using MLR10 mice and we also examined the lens phenotype of germline *Birc7* knockout mice. There was no discernible lens phenotype in either case. The absence of a phenotype indicated that despite its strong expression in lens fiber cells, Livin is not required for normal lens growth, clarity, or cellular organization. It is possible that expression of other Birc family members could compensate for the loss of Livin in the *Birc7* knockouts. However, our microarray data showed that the expression of other family members is usually low in the lens relative to *Birc7* (data not shown) and, in all cases, expression was found to diminish during fiber cell differentiation in wild type lenses.

In conclusion, we have identified a new member of the late fiber cell gene grouping. Livin, known previously for its expression in melanoma and other solid tumors, is strongly expressed by lens fiber cells as they approach the OFZ. To the best of our knowledge, this is the only site in the body where constitutively high level expression of Livin has been reported. The role of Livin in the lens is enigmatic, since conditional inactivation of the *Birc7* locus did not produce an overt lens phenotype. It is possible that Livin serves to block apoptosis under conditions of cellular stress or, alternatively, contributes to organelle breakdown by targeting organelle components for proteolysis. Future experiments will test these possibilities explicitly.

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