Maize Rough Endosperm3 Encodes an RNA Splicing Factor Required for Endosperm Cell Differentiation and Has a Nonautonomous Effect on Embryo Development

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Endosperm and embryo development are coordinated via epigenetic regulation and signaling between these tissues. In maize (Zea mays), the endosperm–embryo signals are not known, but endosperm cellularization is a key event for embryos to form shoots and roots. We screened seed mutants for nonautonomous functions in endosperm and embryo development with genetically nonconcordant seeds and identified the recessive mutant rough endosperm3 (rgh3). The wild-type Rgh3 allele is required in the endosperm for embryos to develop and has an autonomous role in embryo and seedling development. Endosperm cell differentiation is defective in rgh3. Results from endosperm cell culture indicate that rgh3 mutants remain in a proliferative state through mid-seed development. Rgh3 encodes the maize U2AF35 Related Protein (URP), an RNA splicing factor involved in both U2 and U12 splicing. The Rgh3 allele produces at least 19 alternative splice variants with only one isoform encoding a full-length ortholog to URP. The full-length RGH3α isoform localizes to the nucleolus and displays a speckled pattern within the nucleoplasm, and RGH3α colocalizes with U2AF65. A survey of alternatively spliced transcripts found that, in the rgh3 mutant, a fraction of noncanonical splicing events are altered. Our findings suggest that differentiation of maize endosperm cell types is necessary for embryos to develop. The molecular cloning of Rgh3 suggests that alternative RNA splicing is needed for cell differentiation, development, and plant viability.

INTRODUCTION

Angiosperm seeds develop an embryo and endosperm from double fertilization of the egg and central cell of the megagametophyte (De Smet et al., 2010; Linkies et al., 2010). The endosperm is a nutritive tissue for the embryo, and in grasses, it is a persistent storage tissue that is used upon germination (Sabelli and Larkins, 2009). The evolutionary conservation of the endosperm in angiosperms and the coordinated development of endosperm and embryo suggest that these tissues have interacting developmental pathways. In Arabidopsis thaliana, some aspects of embryo development are controlled nonautonomously. Endosperm-to-embryo signaling is supported by two endosperm-specific genes, ABNORMAL LEAF SHAPE1 (ALE1) and RETARDED GROWTH OF EMBRYO1/ZHOUPI (RGE1), which are required for embryos to develop a cuticle and separate from the endosperm (Tanaka et al., 2001; Kondou et al., 2008; Yang et al., 2008). Embryo patterning is not affected in ale1 and rge1 mutants, but ablation of the endosperm with tissue-specific expression of diphtheria toxin causes embryo patterning defects and seed abortion (Weijers et al., 2003).

Early endosperm development is controlled through imprinted genes and genome-wide epigenetic modification of maternal and paternal genomes (Berger and Chaudhury, 2009; Springer, 2009). Maternal and paternal genome dosage is important for seed development. In Arabidopsis interploidy crosses, increased paternal genome copies lead to greater endosperm growth and larger final seed size (Scott et al., 1998). The haku2 and mini-seed3 mutations reduce endosperm growth and seed size by inducing precocious endosperm cellularization (Luo et al., 2005). Thus, endosperm–embryo growth interactions can be uncoupled from interactions influencing patterning. In maize (Zea mays), epigenetic control of the endosperm has a stronger influence on embryo viability with an endosperm requirement for a 2:1 ratio of maternal to paternal genomes for both tissues to develop normally (Lin, 1984).

Genetically nonconcordant seeds, in which the endosperm and embryo have different genotypes, provide a powerful tool to identify nonautonomous genes underlying endosperm–embryo interactions. In maize, nonconcordant endosperm and embryo genotypes can be generated with B-A translocations between the supernumerary B chromosome and standard A chromosomes (Beckett, 1994a). B centromeres undergo nondisjunction during the second pollen mitosis, producing gametophytes with mixed sperm cells (Roman, 1947). When crossed as a male, B-A
translocations generate seeds carrying a deletion of the paternal A chromosome arm in either the embryo or endosperm. B-A translocation crosses will uncover (reveal) recessive defective kernel (dek) mutant phenotypes when the dek maps to the translocated segment. These nonconcordant uncovered kernels can determine if the dek has a tissue autonomous function in growth or development (Sheridan and Neuffer, 1982; Neuffer et al., 1986; Chang and Neuffer, 1994; Scanlon and Myers, 1998; Costa et al., 2003).

We surveyed published primary data for B-A translocation phenotypes and found the vast majority of dek mutants show developmental autonomy (i.e., developmental patterning of the tissue is in accordance to the genotype) (see Supplemental Table 1 online). In these studies, two types of nonautonomous developmental interactions were observed. First, dek2, dek12, dek13, or dek32 mutant embryos are rescued by wild-type endosperm (Chang and Neuffer, 1994). Second, globby1 (glo1), discolored1 (dsc1), or dek26 mutant endosperm inhibits wild-type embryo development (Chang and Neuffer, 1994; Scanlon and Myers, 1998; Costa et al., 2003). Both glo1 and dsc1 cause defects in endosperm cellularization, suggesting a cellular endosperm is needed for embryo development in maize.

Here, we report a maize mutant, rough endosperm3 (rgh3), with a nonautonomous endosperm function that promotes embryo development. We show Rgh3 is required in the endosperm to switch from cellular proliferation to differentiation. These data suggest a unique mechanism by which endosperm cell differentiation influences embryo patterning and development. Molecular cloning of Rgh3 implicates alternative RNA splicing as an essential process to endosperm cell differentiation as well as embryo and plant development.

RESULTS

rgh Mutants Show Endosperm–Embryo Developmental Interactions

To identify seed mutants affecting endosperm–embryo developmental interactions, we crossed B-A translocation stocks for chromosome arms 5L, 6L, and 9L onto 140 rgh mutants. These mutants have a characteristic rough surface to the endosperm that is also termed etched or pitted (Scanlon et al., 1994; Neuffer et al., 1997). We also completed crosses for a small number of rgh mutants using 19 translocations that uncover all chromosome arms except 8S. These experiments uncovered 27 isolates distributed over five chromosome arms (see Supplemental Table 2 online). The uncovered rgh kernels were analyzed with hand sections to identify isolates with endosperm–embryo interactions (Figure 1). Mutants were considered not to have an interaction when they showed autonomous defective endosperm or embryo phenotypes in the two classes of nonconcordant kernels (Figure 1B). An endosperm–embryo interaction was inferred when nonconcordant kernels with both defective endosperm and defective embryo tissues were present. Among the uncovered rgh isolates, 40% (11/27) showed nonautonomous development (see Supplemental Table 2 online). By contrast, we estimate <20% (7/42) of defective kernels described in the literature have nonautonomous endosperm and embryo phenotypes (see Supplemental Table 1 online). These data suggest rgh mutants are enriched for loci required in endosperm–embryo developmental interactions.

We also inferred the direction of the endosperm–embryo interaction from the phenotypes of nonconcordant kernel classes in uncovering crosses. For example, uncovering crosses with the rgh3 mutant produced an embryo defective class with a normal endosperm as well as a defective kernel class with both endosperm and embryo defects. The embryo-defective class suggests wild-type endosperm can develop in conjunction with a mutant embryo. Thus, the defective endosperm and embryo phenotype suggests rgh3 endosperm negatively impacts wild-type embryo development. To confirm this inference, we developed marked nonconcordant kernels by crossing rgh3 with the

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Figure 1. B-A Translocation Crosses Reveal a Nonautonomous Role for rgh3 in Seed Development.

(A) to (C) Sagittal hand sections comparing recessive mutant and TB-5La translocation-uncovering kernel phenotypes. Arrowheads indicate normal embryo shoots and roots. The translocation is indicated by purple aleurone or embryo tissues in the uncovering panels. Bars = 1 mm. (A) pr1 anthocyanin marker. (B) rgh*-01F12226 in a pr1/pr1 background. (C) rgh3 in a pr1/pr1 background. (D) Schematic of expected anthocyanin and developmental phenotypes of autonomous endosperm and embryo defective mutants. Δ represents deletion of the long arm of chromosome 5.
linked pr1 marker. The B-A translocation carries the dominant Pr1 allele, which confers purple color to the aleurone or embryo (Figure 1A). Figure 1B shows marked uncovering crosses for an autonomous isolate, rgh-01F12226, in which mutant, uncovered tissues are red. In rgh3 uncovering crosses, the embryo-defective kernel class has a purple aleurone, indicating the endosperm has the translocation and the embryo was uncovered (Figure 1C). In the nonautonomous kernels, the aleurone is red, indicating the endosperm was uncovered. Rarely (8/82 kernels sectioned), the nonautonomous kernels would develop sufficient embryo tissue to score anthocyanin color, and these embryos are purple, confirming a wild-type embryo with two copies of the normal Rgh3 allele fails to develop in conjunction with rgh3 mutant endosperm.

Inheritance of rgh3

Imprinted loci have multiple roles in seed development, and some imprinted genes have biparental expression at later stages of seed development (Danilevskaya et al., 2003; Gutiérrez-Marcos et al., 2004; Raissig et al., 2011). It is formally possible to explain the rgh3 seed uncovering phenotypes if Rgh3 is an imprinted locus required specifically for early embryo development. Based on this hypothesis, we would expect rgh3 to show transmission bias due to the parent of origin. In self-pollinations of rgh3/+ plants, we observe a 3:1 ratio of normal to rgh3 endosperm phenotypes, suggesting that rgh3 is a recessive allele for endosperm defects (see Supplemental Table 3 online). We did not observe a kernel class with a normal endosperm and a defective embryo in these self-pollinations. To test male and female transmission, we made reciprocal crosses with the reference UniformMu inbred W22 and self-pollinated the F1 progeny to score ears segregating for rgh3. We found ratios close to 1:1 of normal to rgh3 segregating ears regardless of the direction of the cross (see Supplemental Table 3 online). These results indicate rgh3 transmits equally through both gametes and is unlikely to have a parent-of-origin effect on embryo development. Combined with the marked B-A translocation phenotypes, we conclude that rgh3 mutant endosperm has a negative effect on wild-type embryo development. To understand the developmental basis of this interaction, we characterized rgh3 in more detail with a focus on the mutant endosperm.

Seed and Seedling Phenotype of rgh3

Mature rgh3 seeds show a rough, etched, or pitted endosperm surface as well as a reduced seed size compared with normal siblings (Figure 2A). Within individual segregating ears, rgh3 seeds show variable severity, ranging from almost normal grain fill to a nearly empty pericarp phenotype (Figure 2B). rgh3 kernels are reduced in weight but have an overlapping distribution with normal siblings (Figure 2C). The severity of rgh3 embryo defects is correlated to total seed weight with the least severe mutants developing near normal embryos. The most severe class of rgh3 mutants have similar endosperm and embryo phenotypes to rgh3 uncoverings (cf. Figures 1C to 2B), suggesting that a hypoploid endosperm or embryo has a more severe phenotype than the recessive mutant.
determine if rgh3 impacts seed storage protein accumulation, zein and non-zein protein fractions were analyzed from 17-DAP kernels (Figure 2D). In comparison to normal siblings, rgh3 has reduced zein proteins relative to cellular proteins, suggesting rgh3 delays zein accumulation. In mature rgh3 kernels, zein accumulation is correlated to total seed weight with less severe kernels accumulating near normal levels (Figure 2E). Mature rgh3 kernel composition was determined from average weight mutants with standard analytical methods (see Supplemental Table 4 online). Only a slight reduction in kernel fat content was observed, which would be consistent with reduced embryo development of typical rgh3 kernels. These data suggest that rgh3 does not specifically limit protein, starch, or oil biosynthesis.

The range of rgh3 mutant embryo phenotypes led us to test whether rgh3 plants are able to develop. Less than 50% of rgh3 seeds germinate in soil, and the seedlings are smaller and stunted when compared with normal siblings 10 d after planting (Figure 3). All rgh3 seedlings exhibit aberrant development with adherent, narrow leaves resulting in hooked seedlings that died 15 to 18 d after planting. Lethal seedling phenotypes were also observed in F2 progeny after rgh3 was crossed to the B73, Mo17, and A636 inbreds (see Supplemental Figure 1 online). These lethal phenotypes suggest rgh3 impacts multiple developmental processes or Rgh3 is required for cell viability.

**Rgh3 Regulates Endosperm Cell Proliferation in Culture**

To differentiate between these hypotheses, we used cell culture to test rgh3 mutants for a general role in cellular viability. Shannon and Batley (1973) developed an endosperm culture system that produces immortalized, callus-like endosperm cells.

Endosperm culture growth is best with the A636 inbred. We crossed rgh3 into A636 and initiated cultures from self-pollinations of the BC1 and BC2 generations. Mutant and normal seeds could be scored readily at 12 DAP due to a pale color in rgh3 seeds (Figure 4A). Growth in culture was scored with individual endosperm tissues. More than 87% of rgh3 endosperms grew in vitro compared with only 22% of normal siblings (Figure 4B). After 35 d in culture, rgh3 shows massive growth compared with normal siblings (Figures 4C and 4D). Calcofluor white staining of mutant and normal calli was used to visualize cell walls and estimate cell size (Figures 4E and 4F). The mutant cells were smaller than normal siblings, suggesting the excess growth is due to greater cell division in rgh3. Flow cytometry of nuclei extracted from normal endosperm cultures had 3C, 6C, and 12C DNA content, indicating the wild-type Rgh3 genotype undergoes endoreduplication in culture (Figure 4G). By contrast, rgh3 nuclei were mostly 3C with some 6C DNA content, consistent with mitotic cycling (Figure 4H). We then tested the endosperm culture system using 6-, 13-, and 18-DAP ears (see Supplemental Figure 2 online). In these assays, wild-type endosperm tissue readily initiates culture at 6 DAP, while rgh3 homozygous tissue grows readily even if harvested at 18 DAP. These data show Rgh3 inhibits the mitotic cell cycle, although this may be an indirect effect of the mutant. We conclude Rgh3 has roles in endosperm, embryo, and seedling development.

**rgh3 Shows Multiple Endosperm Differentiation Defects**

The proliferative state of rgh3 endosperm cells in tissue culture is not simple to reconcile with the reduced endosperm of rgh3 kernels. Developing maize kernels transport nutrients from the maternal plant through the basal endosperm transfer cell layer (BETL), and prior reports of reduced grain fill mutations suggest that defects in transfer cells are associated with reduced seed size (Lowe and Nelson, 1946; Brink and Cooper, 1947; Maltz et al., 2000; Costa et al., 2003; Gutiérrez-Marcos et al., 2007). We analyzed sections stained with toluidine blue for BETL morphological defects. A developmental time course of wax-embedded sections from 12 to 16 DAP compares mutant and normal differentiation of the BETL region (see Supplemental Figure 3 online). Figures 5A and 5B show higher resolution semithin sections of BETL cells are distributed throughout the cytosol (Figure 5E, white arrowheads) as opposed to normal, polar distribution along on the basal cell wall of differentiating BETL cells (Kang et al., 2009). Moreover, rgh3 cells in the BETL region have a cuboidal cell shape and contain electron-dense bodies similar to protein storage bodies in aleurone cells (Figure 5E). These observations suggest rgh3 BETL differentiation is impaired or delayed.
To test this hypothesis, we analyzed expression of BETL specific markers by quantitative RT-PCR (Figure 5F) and gel analysis of PCR products (see Supplemental Figure 4 online). Aberrant expression was observed for three BETL markers tested. Figure 5F shows that the BETL marker \textit{Tcrr1} is expressed at a basal level in \textit{rgh3} kernels at 15 and 18 DAP, which is the normal peak of expression. The same pattern of reduced expression was observed using an embryo surrounding region (ESR) marker, \textit{ESR1} (Figure 5F). ESR is an endosperm cell type that is physically associated with the developing embryo and expresses secreted peptides (Opsahl-Ferstad et al., 1997; Bonello et al., 2002; Balandín et al., 2005). These cell-type marker data indicate that \textit{rgh3} has defects in both the BETL and ESR.

We extended cellular marker studies to immunolocalization of BETL and aleurone marker proteins at 10 and 20 DAP (Figure 6). At 10 DAP, normal and \textit{rgh3} kernels cannot be distinguished prior to sectioning, but microscopy readily identified kernels with altered endosperm differentiation (Figures 6A and 6B). The \textit{aBETL2} antibody (Serna et al., 2001) labels transfer cells, and BETL2 accumulation reaches a maximum at early developmental stages in normal kernels. Normal sibling kernels had elongated transfer cells accumulating sufficient BETL2 to exclude the blue, azure B, counterstain for secondary cell walls (Figure 6A, BETL signal in brown to black/gray). By contrast, the presumably \textit{rgh3} kernels had flat BETL cells with few cell wall ingrowths and showed greatly reduced BETL2 signal and enhanced azure B counterstaining (Figure 6B).

To investigate the possible causes of \textit{rgh3} variable phenotypic severity, we sorted 20-DAP \textit{rgh3} into classes of phenotypic severity and compared with normal sibling kernels. At 20 DAP, \textit{aBETL2} signal is very weak in normal, fully developed transfer
The most severe in mild rgh3 (Figure 6F). These results suggest BETL development is delayed no cell wall ingrowths, and only a slightly thickened cell wall were decorated with masses outside the BETL region, in the position of ESR cells, significant amounts of BETL2 (Figures 6D and 6E). Occasionally cell densely covered by cell wall ingrowths and accumulating significant of total RNA extracted from the lower half of normal kernels. En, inner endosperm; Pd, pedicel; TCL, transfer cell layer. Bars = 10 μm.

(C) to (E) TEM of rgh3 (C) and (E) and normal (D) BETL region at 15 DAP. Black arrows mark cell wall ingrowths. White arrowheads indicate mitochondria, and black arrowheads point to protein bodies. Bars = 10 μm.

(F) Quantitative RT-PCR of total RNA extracted from the lower half of normal (wild-type [WT]) and rgh3 developing kernels. Error bars show SE from two biological replicates.

cells (Figure 6C, gray/black color), which appear densely covered by cell wall ingrowths (arrowheads, blue stain). Mild rgh3 kernels at 20 DAP had apparently younger transfer cells, less densely covered by cell wall ingrowths and accumulating significant amounts of BETL2 (Figures 6D and 6E). Occasionally cell masses outside the BETL region, in the position of ESR cells, were decorated with αBETL2 (Figure 6E, yellow arrowheads). The most severe rgh3 kernels showed no αBETL2 signal, almost no cell wall ingrowths, and only a slightly thickened cell wall (Figure 6F). These results suggest BETL development is delayed in mild rgh3 mutants and is not specified correctly when the kernel phenotype is severe.

The rgh3 mutants were also examined with an antibody directed against a protease expressed in immature aleurone (G. Hueros, unpublished data). In normal 10 DAP kernels, the aleurone-specific antibody decorated, in addition to the aleurone layer, a few cells at the border between the transfer cell layer and aleurone (Figures 6G and 6H). In mutant 10-DAP kernels, the aleurone marker was found throughout most of the transfer cell layer (Figures 6I and 6J). Similar to 10-DAP kernels, normal 20-DAP kernels had weak signal at the abgerminal edge of the BETL that extends for several cells into the transfer cell area (Figures 6K and 6L). No other signal is detected in normal transfer cells by this antibody at any developmental stage. By contrast, the aleurone marker showed significant signal in the BETL region of severe rgh3 mutants (Figures 6M and 6N). Figures 6F, 6M, and 6N are sections from the same mutant kernel. Thus, endosperm regions that lack expression of BETL2 express an immature aleurone marker. Consistent with reduced expression of the Esr1 marker (Figure 5F), internal endosperm cell masses, in the position of ESR cells, expressed the aleurone marker in severe rgh3 mutants (Figure 6M, red arrowhead, and Figure 6P). In mild rgh3 kernels, the aleurone marker labeled only the abgerminal edge of the BETL in a pattern similar to normal kernels except that the labeling was more intense (Figure 6O). These results suggest that aleurone cells are developmentally delayed in mild rgh3 mutants and that the BETL, ESR, and other endosperm cell types are incorrectly specified when the kernel phenotype is severe.

**Rgh3 Encodes a Predicted RNA Splicing Factor**

We mapped rgh3 to a 0.7-centimorgan interval between mmc0081 and AC207400-2 using an F2 population from crosses of rgh3 with the B73 inbred as well as two-point linkage data with pr1 (Figure 7A). Based on the Genetic 2005 map at MaizeGDB (Sen et al., 2009), two seed mutant loci map near pr1: dek33 and pitted rough germless1 (prg1). Three other seed mutant loci, dek9, dek26, and dek27, were mapped to 5L with B-A translocations. All five mutants are seed lethal and were tested for genetic complementation with rgh3 by crossing normal progeny from segregating ears. Results from these tests suggest all five loci are not alleles of rgh3 (see Supplemental Table 5 online).

DNA gel blots identified a Mu1 insertion linked to the rgh3 phenotype (see Supplemental Figure 5 online). Robertson’s Mutator insertion site thermal asymmetric interlaced PCR was used to amplify and clone the linked Mu1 insertion site (Settles et al., 2004), and linkage analysis was expanded to 1020 meiotic products without any recombinants between the Mu1 element and rgh3 observed. This Mutator-induced rgh3 reference allele was then named rgh3-umu1. We identified a second allele of rgh3 using chemical mutagenesis tagging crosses. Ethyl methanesulfonate (EMS)–mutagenized W22 pollen was crossed onto rgh3-umu1/+ plants. Among the F1 progeny, we identified four rgh3/rg3 rgh3 seedlings that included three homozygous rgh3-umu1 self-contaminants (Figure 7B). The remaining rgh3-umu1/rg3* seedling was analyzed by PCR amplification of the genomic locus and Cell digest to identify two point mutations in the Rgh3 gene (see Supplemental Figure 6 online). One mutation within intron 6 is predicted to be silent. The second mutation was a G-to-A transition resulting in a premature stop codon (Figure 7B, right side). Combined, these data indicate that we identified the Rgh3 locus.
Figure 6. Immunolocalization of Transfer and Aleurone Cell Markers.

(A) to (F) Kernel sections decorated with antibodies against BETL2.

(G) to (P) Kernel sections decorated with antibodies against an aleurone protease.

Normal Rgh3 kernels are grouped in the left panels and are outlined with a black border ([A], [C], [G], [H], [K], and [L]). Mutant rgh3/rgh3 kernels are shown in the right panels ([B], [D] to [F], [I], [J], and [M] to [P]). Kernel sections are organized chronologically with (A), (B), and (G) to (J) from 10-DAP kernels. (C) to (F) and (K) to (P) are from 20-DAP kernels. Black arrowheads point to cell wall ingrowths in (C), (D), and (O). Yellow arrowheads in (E) point to a mass of cells in the ESR decorated with αBETL2. Red arrowhead in (M) points to a mass of cells, at the position of the embryo suspensor, decorated with the aleurone marker. Red dotted lines mark the border between aleurone and transfer cells ([H], [J], and [L]). Red boxes labeled L, N, and P frame areas shown at higher magnification in adjacent panels ([L], [N], and [P]). Bar = 1 mm in (G), (I), and (K) and 100 μm in all other panels. Positive immunostaining appears gray, black, or dark brown against blue counterstaining.

(A) Normal Rgh3 kernels, 10 DAP, anti-BETL2 antibodies.
(B) Mutant rgh3/rgh3 kernels, 10 DAP, anti-BETL2 antibodies.
(C) Normal Rgh3 kernels, 20 DAP, anti-BETL2 antibodies.
(D) Mutant rgh3/rgh3 kernels, 20 DAP, anti-BETL2 antibodies.
(E) Mutant rgh3/rgh3 kernels, 20 DAP, anti-BETL2 antibodies.
(F) Mutant rgh3/rgh3 kernels, 20 DAP, anti-BETL2 antibodies.
(G) Normal Rgh3 kernels, 10 DAP, antialeurone protease antibodies.
(H) Normal Rgh3 kernels, 10 DAP, antialeurone protease antibodies.
(I) Mutant rgh3/rgh3 kernels, 10 DAP, antialeurone protease antibodies.
(J) Mutant rgh3/rgh3 kernels, 10 DAP, antialeurone protease antibodies.
(K) Normal Rgh3 kernels, 20 DAP, antialeurone protease antibodies.
(L) Normal Rgh3 kernels, 20 DAP, antialeurone protease antibodies.
(M) Mutant rgh3/rgh3 kernels, 20 DAP, antialeurone protease antibodies.
(O) Mutant rgh3/rgh3 kernels, 20 DAP, antialeurone protease antibodies.
(P) Mutant rgh3/rgh3 kernels, 20 DAP, antialeurone protease antibodies.

Al, aleurone; Em, embryo; En, inner endosperm; Pd, pedicel; TCL, transfer cell layer.
overlapping fragments from the maize BAC ZMMBBc497J22. AGPv2 reference, and we cloned and sequenced the locus in two RT-PCR products. These clones identified 19 Rgh3 and U12 introns in mammals (Tronchère et al., 1997; Shen et al., 2000). URP is involved in RNA splicing of both U2 and U12 introns in mammals (Tronchère et al., 1997; Shen et al., 2000). URP is involved in RNA splicing of both U2 and U12 introns in mammals (Tronchère et al., 1997; Shen et al., 2000). URP is involved in RNA splicing of both U2 and U12 introns in mammals (Tronchère et al., 1997; Shen et al., 2000).

The Rgh3 locus is not correctly assembled in the current B73 AGPv2 reference, and we cloned and sequenced the locus in two overlapping fragments from the maize BAC ZMMBBc497J22. The predicted RGH3 protein is homologous to human U2AF35 Related Protein (URP). URP is involved in RNA splicing of both U2 and U12 introns in mammals (Tronchère et al., 1997; Shen et al., 2010). We amplified, cloned, and sequenced 45 independent Rgh3 RT-PCR products. These clones identified 19 Rgh3 splice variants with the potential to encode seven protein variants. The Rgh3 transcript variants result from variable intron retention, exon skipping, and alternative 5’ and 3’ splice sites. Figure 7B shows schematics for seven examples of Rgh3 isoforms that code for the different protein sequences. The Rgh3α isoform encodes a predicted 86.1-kD protein that is orthologous to URP. Splicing of intron 1 produces a likely noncoding message that deletes the start codon but still contains an uninterrupted URP reading frame that could initiate translation within the UHM domain. The other 17 isoforms are predicted to encode partial proteins ranging from 20 to 41.1 kD (see Supplemental Figure 7 online). There were four Rgh3 transcript variants that are predicted to code for the RGH3β protein and nine transcript variants predicted to code for the RGH3γ protein. Only a single clone was recovered for each of the four other predicted RGH3 protein variants.

To estimate the relative levels of transcripts coding for the RGH3α protein, we designed a quantitative RT-PCR assay for the exon skipping RNA splicing event required to code for the full-length protein (skipping both exons 6 and 7). We compared the relative abundance of the exon 6/7 skip to three splicing events that cause truncation of the open reading frame (ORF) (retaining intron 6, skipping exon 7, and splicing intron 5 to include exon 6). The levels of these Rgh3 splicing events are regulated in a tissue- and developmental-specific manner (see Supplemental Figure 8 online). The exon 6/7 skip event accumulates at its highest levels in 3-cm immature ears and late in seed development, post 30 DAP. The exon 6/7 skip is also found at high levels in early seed development from 6 to 16 DAP as well as in the ligule and midvein of adult leaves. The usage of Rgh3 alternative splice sites is regulated with the exon 6/7 skip, representing 8.1 to 26.7% of the total Rgh3 transcripts assayed (see Supplemental Figure 8C online). Rgh3α-like transcripts were most efficiently spliced at 20-, 34-, and 45-DAP in seeds.

Surprisingly, Rgh3α-specific primers that flank the Mu1 insertion were able to amplify a cDNA fragment from rgh3-umu1 mutant tissues with only a small size shift relative to wild-type products (Figure 7C). Sequencing of eight rgh3-umu1 cDNA clones identified 141 bp from the Mu1 terminal inverted repeat, indicating Mu1 is partially spliced from mutant transcripts. The spliced Mu1 sequence deletes 12 amino acids of the wild-type protein and inserts 47 new amino acids, suggesting the mutation is likely to be a hypomorph (see Supplemental Figure 7 online).

The full-length RGH3α protein encodes a 755–amino acid peptide with a domain structure similar to URP (Figure 8A). URP has an N-terminal acidic domain, a central U2AF Homology Motif (UHM) flanked by zinc-finger motifs, and a C-terminal Ser/Arg-rich region (RS domain). The UHM is related to the RNA Recognition Motif but is responsible for protein–protein interactions rather than binding RNA (Kielkopf et al., 2004). The zinc-finger and UHM region of the protein is highly conserved between maize, Arabidopsis, and humans, whereas the N- and C-terminal sequences are variable with only sequence composition being conserved (see Supplemental Figure 9 online). The U2AF35 protein lacks the acidic N-terminal domain. Phylogenetic analysis of the zinc-finger/UHM region indicates that URP is a distinct clade from U2AF35 proteins and is a single-copy gene in maize, Arabidopsis, and rice (Oryza sativa; Figure 8B). The Rgh3 transcript variants coding for RGH3ζ and RGH3η truncate the ORF.
such that the conserved zinc fingers, UHM, and RS domains are out of frame with the N-terminal acidic domain (see Supplemental Figure 7 online). RGH3β, RGH3γ, and RGH3δ include the N-terminal zinc finger in the ORF. RGH3ζ is predicted to code for protein that includes the acidic, one zinc finger, and the UHM domain, while the EMS allele only truncates the RS domain.

**RGH3 Is Localized to Subnuclear Compartments**

Transient expression of an RGH3α-green fluorescent protein (GFP) fusion in Arabidopsis mesophyll protoplasts localized to compartments of varying size within the nucleus (Figure 9). Analysis of 200 transformed protoplasts gave an equal number of cells with RGH3α-GFP localized to the nucleolus or with a speckled pattern within the nucleoplasm plus localization to the nucleolus (Figure 9B). A similar pattern of localization to the nucleolus or nuclear speckles was observed when URP, the Arabidopsis ortholog of RGH3, was fused to GFP and expressed in Arabidopsis protoplasts (see Supplemental Figure 10A online). These data suggest the maize localization pattern is not an artifact of heterologous expression. Coexpression of RGH3α-GFP with an established nucleolus marker, Arabidopsis PRH75 (Lorkovic et al., 2004), confirmed RGH3α localization to the nucleolus (see Supplemental Figure 10B online). RNA processing proteins, such as RNA splicing factors, are typically localized to nuclear speckles and can be localized to the nucleolus in response to cellular stress (Tillemans et al., 2005; Koroleva et al., 2009). We then expressed a GFP fusion with the EMS-truncated reading frame, RGH3EMS-GFP, which resulted in cytoplasmic localization (Figure 9A). Using an N-terminal GFP fusion expressed in Nicotiana benthamiana, we found GFP-RGH3α localized to the nucleolus and nuclear speckles as in Arabidopsis protoplasts (Figure 9C). We then used this expression system to localize RGH3umut-GFP as well as three additional wild-type RGH3 isoforms: β, γ, and ζ. These fusions expressed at low levels and localized to the nucleolus as well as diffusely throughout the nucleoplasm. These data suggest RGH3 localization, and potentially protein stability, result from signals or interacting domains within the acidic, UHM, and RS domains. Both the Mu element insertion and EMS alleles alter RGH3α protein localization, suggesting the alleles alter protein function.

To confirm that the localization patterns of GFP fusions with RGH3α represent RNA splicing complexes, we coexpressed RGH3α-GFP with a core splicing factor. We fused the ORF of the maize U2 Auxiliary Factor 65-kD subunit locus a (U2AF65a) to red fluorescent protein (RFP), Figure 9D shows that RGH3α-GFP expression overlaps with U2AF65a-RFP expression in both the nucleolus and in speckles. These results suggest the speckled pattern in RGH3α fusion proteins is likely to represent spliceosomes. Interestingly, expression of GFP-U2AF65a resulted in a speckled pattern, with U2AF65a being excluded from the nucleolus. Thus, high levels of RGH3α expression influence U2AF65a subnuclear localization. These data are consistent with the known protein–protein interaction of human URP and U2AF65 (Tronchère et al., 1997).

**Rgh3 Is Required for a Subset of RNA Splicing Events**

In human cells, URP participates in both U2 and U12 splicing and is essential for cell culture survival (Tronchère et al., 1997; Shen et al., 2010). The viability of rgh3 endosperm cells in culture suggests the rgh3-umu1 hypomorphic allele does not disrupt core splicing functions. To assay rgh3 mutants for altered splicing, we selected 21 maize genes with evidence for alternative splicing in maize, Arabidopsis, and rice (Wang and Brendel, 2006; see Supplemental Table 6 online). Only three genes had differences in isoform usage in rgh3 and wild-type tissues. GRMZM2G165901 encodes a Gly-rich RNA binding protein with two major splice variants (Figure 10A). Sequencing of these isoforms found a noncanonical intron retention with GU-CG dinucleotides at the 5’ and 3’ splice sites instead of GU-AQ. The rgh3 mutation reduces the level of intron retention in seed and seedling tissue, suggesting noncanonical splicing is more efficient in the mutant. The smaller variant contained different splice junctions in rgh3 and the wild type with rgh3 shifting the splice acceptor site by three bases.
GRMZM2G051276 encodes a putative inositol monophosphatase with three splice variants (Figure 10B). Sequencing of these variants found complex alternative splicing within annotated intron 10. First, a four-base exon is found that results from splicing of two noncanonical introns with UU-AA and AU-UU dinucleotides. Second, an alternative variant contains only a single intron between the annotated exons 10 and 11. Assuming the same donor site is used in the alternative variant, the dinucleotides for the predicted intron are UU-UU, indicating this intron is also noncanonical. Relative to the wild type, the alternative, retained intron variant accumulates to higher levels in rgh3 seed tissue but to lower levels in endosperm culture.

GRMZM2G081642 encodes a protein of unknown function, which is conserved within plants and algal species. Sequencing of four variants identified an alternative donor site for intron 2 in which the shorter variant is a GC-AG intron. The longer variant splices with a canonical GU-AG intron, and the rgh3 tissues have higher levels of this canonical donor site. Intron 3 has three...
alternative acceptor sites that all have canonical dinucleotides. Combined, these data suggest that RGH3\textsuperscript{umu1} modulates splice site selection for a subset of noncanonical introns.

**DISCUSSION**

A Role for Endosperm Cell Differentiation in Endosperm–Embryo Developmental Interactions

The endosperm is thought to have a relatively minor role in embryo development but important roles in embryo and seed growth by providing nutrients and acting as a regulator of seed size (De Smet et al., 2010). Maize \textit{dek} B-A translocation uncoverings generally support this theory with most types of reduced or defective endosperm tissues being tolerated by genetically wild-type embryos (Neuffer and Sheridan, 1980; Neuffer et al., 1986; Chang and Neuffer, 1994). For a few cases, mutant embryo growth benefits from the presence of a wild-type endosperm, supporting the model in which the endosperm nurtures the embryo (see Supplemental Table 1 online). By contrast, we found the \textit{rgh} mutant class to be enriched for nonautonomous endosperm–embryo developmental interactions (see Supplemental Table 2 online). Most of these interactions are consistent with mutant endosperms causing wild-type embryos to abort or develop abnormally. For \textit{rgh3}, we confirmed the endosperm to embryo interaction by marking the nonconcordant genotypes using the \textit{pr1} anthocyanin locus. These data suggest the \textit{rgh} mutant class is likely to reveal mechanisms by which the endosperm promotes embryo development. Our characterization of the \textit{rgh3} phenotype illustrates a unique developmental mechanism for endosperm–embryo interactions.

As far as we are aware, \textit{glo1} is the only other \textit{dek} mutant with a similar endosperm–embryo developmental interaction for which the mutant endosperm defects have been characterized at a cellular level (Costa et al., 2003). Cellularization and cytokinesis are affected in \textit{glo1}, and \textit{Glo1} is likely to be essential for cell viability. The \textit{dsc1} mutation has a similar endosperm to embryo developmental interaction. At a macroscopic level, \textit{dsc1} endosperm fails to solidify by 16 DAP, suggesting defects in endosperm cellularization (Scanlon and Myers, 1998). The \textit{dsc1} and \textit{glo1} defects imply endosperm cellularization is critical for embryo development. By contrast, \textit{Arabidopsis} mutants in the \textit{SP\textATZLE} and \textit{ENDOSPERM DEFECTIVE1} (\textit{EDE1}) loci can produce homozygous viable embryos even though the endosperm never cellularizes, suggesting endosperm cellularization is not an absolute requirement for embryo development (Sørensen et al., 2002; Pignocchi et al., 2009).

The \textit{rgh3} mutant further argues that cellularization is not the key endosperm event required for embryo development. The \textit{rgh3-umu1} allele does not impact endosperm cell viability or cellularization. Bulk kernel composition analysis suggests there are no major differences in the ability of \textit{rgh3} cells to accumulate seed storage molecules (see Supplemental Table 4 online).
Instead, the cellular phenotypes of rgh3 indicate the gene is necessary for cells to switch from a proliferative to differentiating state. Multiple lines of evidence argue for this interpretation. The in vitro rgh3 cell cultures have smaller cell size and do not undergo endoreduplication (Figures 4E to 4H), while developing endosperm cells in rgh3 kernels are reduced in size (Figures 5A and 5B). Morphological, molecular, and protein markers indicate that rgh3 cells have endosperm cell differentiation defects. The starchy endosperm has reduced storage protein levels (Figures 2D and 2E); the BETL and ESR do not express cell type markers at wild-type levels (Figure 5F); an aleurone marker is delayed in expression and expressed in inappropriate cell positions (Figures 6G to 6P); and rgh3 endosperm retains the ability to grow in vitro 12 d later in development than the wild type (see Supplemental Figure 2 online). These data argue that one or more aspects of endosperm cell differentiation are needed for the embryo to develop. We suggest mutants affecting endosperm cellularization are likely upstream of the cell differentiation pathways affected in rgh3.

**Is There a Key Endosperm Cell Type Required for Embryo Development?**

A common endosperm phenotype in both glo1 and rgh3 is a defective BETL. It is tempting to conclude that loss of BETL function would limit the sink strength of a developing cell and lead to defective embryo development. However, multiple dek mutants disrupt BETL differentiation without inhibiting embryo development. Mutants in defective endosperm17, minute1, reduced grain-filling1, and baseless1 (bs1) produce viable embryos but have morphological BETL defects correlated with reduced grain fill (Lowe and Nelson, 1946; Brink and Cooper, 1947; Matiz et al., 2000; Gutiérrez-Marcos et al., 2006; Kang et al., 2009). Similar to rgh3, bs1 shows a range of seed phenotypes, including an embryo-lethal phenotype in seeds at wild-type levels (Figure 5F); an aleurone marker is delayed in expression and expressed in inappropriate cell positions (Figures 6G to 6P); and rgh3 endosperm retains the ability to grow in vitro 12 d later in development than the wild type (see Supplemental Figure 2 online). These data argue that one or more aspects of endosperm cell differentiation are needed for the embryo to develop. We suggest mutants affecting endosperm cellularization are likely upstream of the cell differentiation pathways affected in rgh3.

**Weak Alleles of rgh3 Reveal Developmental Roles for Maize URP**

The Rgh3 gene is a predicted core RNA splicing factor. In plants, a variety of RNA splicing factors have been shown to have specific roles in abiotic stress tolerance and flowering time (reviewed in Lorkovic, 2009). Mutants and misexpression transgenics of both U2 and U12 splicing factors can have pleiotropic developmental defects (Lopato et al., 1999; Kalyna et al., 2003; Ali et al., 2007; Chung et al., 2007; Coury et al., 2007; Kim et al., 2010). Most plant genes contain introns that must be removed for protein expression; therefore, RNA splicing should be an essential function for cell viability. Consistent with this reasoning, loss-of-function alleles of core splicing factors are gametophyte or seed lethal (Liu et al., 2009; Wang and Okamoto, 2009; Kim et al., 2010).

The RGH3/ZmURP homologs in animals have only been studied to a limited extent. Human URP is a required core factor for both U2 and U12 splicing in vitro (Tronchère et al., 1997; Shen et al., 2010). URP proteins have a similar domain structure to U2AF35 and can interact with U2AF65 (Tronchère et al., 1997). We found RGH3x to colocalize with maize U2AF65a and redirect U2AF65a into the nucleolus supporting a similar model for URP protein–protein interactions in plants (Figure 9D). URP contacts the 3' splice site in both U2 and U12 splicing reactions but has different roles for each type of intron with U2AF65 absent from U12 spliceosomes (Shen et al., 2010). Small interfering RNA knockdowns of URP and U2AF35 show that the human genes are essential for cell viability (Shen et al., 2010). Although the rgh3-umu1 mutation is seed and seedling lethal, RNA splicing is not broadly affected in rgh3 mutants, and rgh3 endosperm is completely viable in cell culture. These data are consistent with the recessive inheritance and molecular nature of rgh3-umu1, suggesting the allele is hypomorphic. The Mu insertion disrupts the coding sequence of the locus, but the transposon contains a cryptic splice site that creates an insertion-deletion mutation in the acidic domain of the RGH3x protein (Figure 7; see Supplemental Figure 7 online). The primary amino acid sequence of the
acidic and RS domains of URH homologs is not highly conserved. Instead, these domains have conserved amino acid composition. The acidic domain of the RGH3umu1α protein has a pI of 5.64, which is identical to the wild-type protein. B-A translocation uncovering of rgh3 further supports the hypothesis that rgh3-umu1 encodes a weak allele. Uncovered embryo and endosperm phenotypes are more severe than recessive rgh3umu1 mutant seeds (cf. Figures 1 and 2). In uncovered kernels, the entire 5L chromosome arm is deleted. It is possible there are rgh3 modifier loci on 5L that are haploinsufficient. However, we think it more likely that a single dose of RGH3umu1α is insufficient to promote embryo or endosperm development. The EMS-induced rgh3* allele may also be hypomorphic. The EMS allele supported embryo development and retains the conserved domains predicted to be involved in protein–protein interactions with U2AF65.

GFP fusions with RGH3 isoforms and mutant alleles provide insight on the function of RGH3 protein domains. For RGH3α, we observed exclusive localization to the nucleolus as well as a distribution between nucleolar and speckled localization. The speckled localization is likely to represent splicing complexes because RGH3α colocalizes with U2AF65α in the nucleoplasm. The distribution of nucleolar and speckle localization of RGH3α is not typical for plant pre-mRNA splicing factors (Docquier et al., 2004; Fang et al., 2004; Lorkovic et al., 2008; Tanabe et al., 2009). A few splicing factors show dynamic localization between the nucleolus and the nucleoplasm (Tillemans et al., 2005; Koroleva et al., 2009). The targeting of these splicing factors to the nucleolus is influenced by cellular stress, such as hypoxia, and levels of protein phosphorylation. RGH3α may show similar dynamic regulation of its localization. It is also possible that overexpression of RGH3α fusion proteins may stress the cell sufficiently to cause nucleolar localization.

Nuclear speckle localization is lost in RGH3umu1α and in RGH3 splice variants lacking the UHM, one zinc-finger, and the RS domains. The RGH3umu1α protein suggests that the acidic domain is important for the protein to be incorporated into nuclear speckles, while the RGH3β, γ, and ζ proteins implicate the UHM, C-terminal zinc-finger, or RS domain of the protein for nuclear speckle localization. In Arabidopsis, deletion of the RS domain from RGSzp22 results in a similar loss of speckle localization and directs the splicing factor to the nucleolus (Tillemans et al., 2005). However, the rgh3* EMS allele truncates only the RS domain and shows cytoplasmic localization. The RS domain may be required to unmask the nuclear localization signal in the N-terminal acidic domain. Alternatively, a nuclear export signal may reside in the C-terminal region of the UHM domain or the C-terminal zinc finger. Combined, the GFP fusion data suggest RGH3α contains multiple targeting signals that can direct nuclear versus cytoplasmic localization as well as subnuclear localization to the nucleolus or to speckles.

Alternative RNA Splicing in Endosperm Cell Differentiation

In animals, alternative splicing is associated with cell-type and tissue-specific expression patterns (Blencowe, 2006). Multiple genetic diseases have been identified that show alternative RNA splicing is essential for different cell types to function correctly (Ferreira et al., 2010; Twine et al., 2011). However, only a few proteins with KH-type RNA binding domains have been shown to act as cell differentiation regulators (Ule et al., 2005; Volk et al., 2008). Endosperm cell differentiation is clearly defective in rgh3 mutants. We observe delayed or misexpression of all endosperm cell types, including zein storage proteins in the starchy endosperm, a protease in the aleurone, multiple BETL markers, and Esr1. Moreover, rgh3 endosperm retains the ability to grow in culture up to the latest time point tested, 18 DAP, while wild-type endosperm only grows at similar efficiency at 6 DAP. These data indicate a role for RNA splicing in endosperm cell differentiation. Our survey of splicing defects in rgh3 suggests that the RGH3umu1α protein alters splicing in a subset of noncanonical introns. Genome-wide-level analysis of rgh3 splicing defects would help identify the specific genes required to promote normal endosperm cell differentiation. Similar differentiation defects could be the cause of the rgh3 lethal seedling phenotype, but we did not investigate the rgh3 seedling phenotype in detail in this study.

In conclusion, we have shown that endosperm cell differentiation is required for wild-type embryo development. By comparing rgh3 cellular defects with other maize seed mutants, we speculate that the ESR is a critical cell type to promote embryo development. The cloning of rgh3 identifies alternative RNA splicing as a key process required for endosperm cell differentiation. The hypomorphic rgh3-umu1 allele suggests the acidic domain of URP has a role in alternative splicing in maize.

METHODS

B-A Translocation Genetics

All rgh mutants were heritable isolates from the UniformMu population (McCarty et al., 2005). Plants segregating for rgh/+ genotypes were crossed as females with hyperploid or heterozygous B-A translocations (Birchler and Alfenito, 1993; Beckett, 1994a, 1994b). For each translocation, 10 to 15 crosses were attempted with multiple B-A plants. A mutant was considered uncovered when multiple F1 crosses showed rgh phenotype specific to a single chromosome arm. To identify endosperm–embryo interactions, sagittal sections of mature kernels were cut with a utility knife. To generate marked uncovering crosses, rgh3/+ plants carrying dominant alleles for anthocyanin genes were crossed onto a red aleurone (pr1) stock in the W22 inbred. F2 pr1/pr1 kernels were self-pollinated to identify rgh3 recombinants. Normal kernels from segregating F3 ears were used as females for B-A crosses.

Molecular and Genetic Mapping

F2 progeny from B73 × rgh3/+ crosses were scored for rgh3/rgh3 seedling phenotypes. DNA was extracted from seedling leaves as described by Settles et al. (2004). Simple sequence repeat markers from chromosome 5 were amplified and scored for recombination, and Haldane map distances were calculated. PCR conditions were optimized for each marker based on initial conditions from Martin et al. (2010). Primer sequences for the markers are in Supplemental Table 7 online.

Genetic complementation tests used reciprocal crosses between plants segregating for rgh3, dek9, dek26, dek27, dek33, and pr1 heterozygotes. The first ears of male parents in the crosses were self-pollinated to determine plant genotypes. The second ears of female parents in each cross were self-pollinated when possible. Crosses were
screened visually, and seed phenotypes were expected to approximate segregation ratios of self-pollinated parents.

The rgh3 Mu1 insertion site was cloned from a PstI digest of genomic DNA from a rgh3/rgh3 mutant seedling at the BC2_S1 generation of a Mu-intrinsically introgression into W22. The DNA fraction from 3.5 to 4.2 kb was gel purified and used as template for Robertson’s Mutator insertion site thermal asymmetric interlaced PCR (Settles et al., 2004). PCR products were cloned and sequenced as described (Settles et al., 2004). The majority of the Rgh3 locus was subcloned from a BamHI digest of the B73 BAC, ZMMBBc49722. A 6-bp overlapping 3’ region was amplified from the BAC using TaKaRa LA Taq DNA Polymerase and cloned using a pTOPO vector (Invitrogen). Full-length Rgh3 cDNA was amplified using 5’ and 3’ rapid amplification of cDNA ends PCR with the GeneRacer kit (Invitrogen), and splice variants were amplified using Phusion high fidelity polymerase (Finnzymes). cDNA products were cloned with the Zero Blunt TOPO PCR cloning kit (Invitrogen).

The rgh3* allele was generated by pollinating 150 rgh3-umu1/+ plants with EMS-mutagenized W22 pollen as described (Neuffer, 1993; Douglas et al., 2010). Approximately 4000 progeny were screened for the rgh3 seedling phenotype. Point mutations in the Rgh3 locus were identified using the Surveyor mutation detection kit (Transgenomics).

**Quantitative Kernel Phenotypes**

Normal and rgh kernels from 10 rgh3 segregating ears were separated. Seed weights were collected from a automatically microbalance as described by Spielbauer et al. (2009). Kernels were ground to a fine meal using 15 g of meal from normal and average weight kernels (140 to 200 mg) with Association of Analytical Communities methods. Bulk tissues were used to initiate cultures on Murashige and Skoog media, pH 8.0 with 3% Suc, 4 ppm thiamine, 0.2% Asn, and 2% phytagel. Cultures were described by Gibbon et al. (2003). Zein and non-zein protein fractions were analyzed on a fresh weight basis with 12.5% SDS-PAGE. Bulk cultures were gravity filtered through four layers of cheesecloth and then 100-mL of extraction buffer (10% glycerol, 2% SDS, 100 mM DTT, 50 mM Tris-HCl, pH 6.8, and 1× Complete protease inhibitor cocktail [Roche Applied Science]) at 37°C with agitation for 2 h. Extracts were centrifuged at 12,000g for 10 min and supernatants processed for zein purification as described by Gibbon et al. (2003). Zein and non-zein protein fractions were analyzed on a fresh weight basis with 12.5% SDS-PAGE. Bulk composition analysis was completed by Waters Agricultural Laboratories using 15 g of meal from normal and average weight rgh3 kernels (140 to 200 mg) with Association of Analytical Communities methods.

**Endosperm Culture**

Self-pollinations from A636 × rgh3/+ BC1, and BC2 generations were used for endosperm cultures as described by Shannon (1994). Briefly, 6-, 12-, 13-, or 18-DAP ears were surface sterilized, and individual endosperm tissues were used to initiate cultures on Murashige and Skoog media, pH 5.7, with 3% Suc, 4 ppm thiamine, 0.2% Asn, and 2% phytagel. Cultures were incubated at 30°C in the dark for 35 d. Growth was assessed for over 250 cultured endosperms for both rgh3 and normal siblings. Fragments of calli were stained with calcofluor-white (0.01% in 1× PBS), mounted on glass slides, and observed using a Nikon Diaphot inverted microscope with epifluorescence illumination and a UV filter. Representative pictures were taken with a Leica DC300 digital camera.

**Flow Cytometry**

Approximately 150 mg of endosperm culture was chopped in 1.5 mL of ice-cold chopping buffer (20 mM MOPS, pH 7.2, 45 mM MgCl2, 30 mM Na3Citrate, 0.1% Triton X-100, and 0.01% RNase A) using a razor blade. Homogenates were gravity filtered through four layers of cheesecloth and then 100-μm nylon mesh. Samples were centrifuged for 2 min at 600g. Pelleted nuclei were resuspended in chopping buffer with 0.05% propidium iodide, BD FACSort (BD Biosciences) with “no gate” for the setting parameter was used for flow cytometry. Flow cytometry plots are representative of 10 replicates of 5000 events counted.

**Microscopy**

Purple kernels from crosses of Rgh3 pr1/Rgh3 pr1 × rgh3 pr1/Rgh3 pr1 were selected to enrich for ~93% rgh3/+ plants. Developing ears were harvested from 10 to 20 DAP. Kernels were processed for semithin (500 nm) and thin (80 nm) sections as described by Kang et al. (2009). Representative pictures of semithin, toluidine blue (0.05%) stained sections were taken using an Olympus BH2 microscope and Retiga 2000R digital camera (QImaging). Transmission electron micrographs of thin sections were captured with a Hitachi H-7000 microscope operated at 80 kV. For the developmental time course and immunolocalizations, kernels were fixed at 4°C overnight in FAA solution (3.7% formaldehyde, 5% glacial acetic acid, and 45% ethanol). Kernels were dehydrated in an ethanol series, embedded in paraffin or Paraplast plus embedding medium (Sigma-Aldrich), and cut into 8- to 10-μm sections. The sections in the developmental time course were stained with toluidine blue. Immunolocalization used each preserum-serum pair at equivalent dilutions. Slides were rehydrated in an ethanol series, and endogenous peroxidase activity was deactivated in 0.3% hydrogen peroxide. The sections were blocked with 2% normal donkey serum and incubated with antisera or presera. The slides were washed twice in 140 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4, and 0.1% Tween-20, pH 7.3, and incubated in biotin-conjugated α-rabbit goat antibody at 1/750 (Sigma-Aldrich). After two washes, an Extravidin-peroxidase (Sigma-Aldrich) solution at 1/800 was incubated with each slide. Following two washes, the slides were blocked with 2% normal donkey serum and incubated with antisera or presera. The slides were washed twice in 140 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4, and 0.1% Tween-20, pH 7.3, and incubated in biotin-conjugated α-rabbit goat antibody at 1/750 (Sigma-Aldrich). After two washes, an Extravidin-peroxidase (Sigma-Aldrich) solution at 1/800 was incubated with each slide. Following two washes, the slides were blocked with 2% normal donkey serum and incubated with antisera or presera. The slides were washed twice in 140 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4, and 0.1% Tween-20, pH 7.3, and incubated in biotin-conjugated α-rabbit goat antibody at 1/750 (Sigma-Aldrich). After two washes, an Extravidin-peroxidase (Sigma-Aldrich) solution at 1/800 was incubated with each slide. Following two washes, the slides were blocked with 2% normal donkey serum and incubated with antisera or presera. The slides were washed twice in 140 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4, and 0.1% Tween-20, pH 7.3, and incubated in biotin-conjugated α-rabbit goat antibody at 1/750 (Sigma-Aldrich). After two washes, an Extravidin-peroxidase (Sigma-Aldrich) solution at 1/800 was incubated with each slide. Following two washes, the slides were blocked with 2% normal donkey serum and incubated with antisera or presera. The slides were washed twice in 140 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4, 1.8 mM KH2PO4, and 0.1% Tween-20, pH 7.3, and incubated in biotin-conjugated α-rabbit goat antibody at 1/750 (Sigma-Aldrich).

**Subcellular Localization**

N- and C-terminal GFP fusion proteins were constructed by cloning cDNAs for RGH3 isoforms and the coding Arabidopsis thaliana URP isoform into pDONR221 or pENTR (Invitrogen) according to the manufacturer’s instructions. The ORFs were subcloned into p2GW7 for C-terminal GFP fusions or pB7-WGF2 for N-terminal GFP fusions (Karimi et al., 2002, 2007) by LR recombination (Invitrogen). Transient expression experiments in Arabidopsis were performed with protoplasts isolated from leaves or cell suspension culture (Yoo et al., 2007). Transient expression experiments in Nicotiana benthamiana were originally described by Sofia (1999) with the following modifications. Binary vectors were transformed into Agrobacterium tumefaciens strain ABI by a freeze-thaw method (Wise et al., 2006). MES was not included in the Agrobacterium growing media, and N. benthamiana infiltration was completed with a 10-mL needleless syringe on 4- to 5-week-old plants grown in growth chamber at 22 to 24°C with 16h/8h day/night. For coclonalization experiments, Agrobacterium strains carrying individual plasmids were mixed in a 1:1 ratio prior to infiltration. Fusion protein expression was visualized 24 h after transient transformation, and representative pictures of subcellular localization were obtained using a Zeiss Pascal LSM5 confocal laser scanning microscope as previously described (Pribat et al., 2010). GFP/RFP colocalization images were obtained with a Leica TCS SP2 confocal laser scanning microscope (Leica Microsystems). GFP was excited at 488 nm and detected with an emission band of 500 to 545 nm. Monomeric RFP was excited at 594 nm and detected with an emission band of 605 to 645 nm.

**Expression Analysis**

Total RNA was extracted from maize (Zea mays) tissues as described by Reid et al. (2006) using 10 mL of extraction buffer/g of fresh weight. For molecular markers, maize kernels were cut in half with a transverse...
section as described by Gómez et al. (2009), and RNA was extracted from the basal sections. RNA samples were treated with RNase-free DNase I (New England Biolabs), extracted with phenol-chloroform, and ethanol precipitated. RT-PCR was performed on first-strand cDNA and synthesized by Superscript III (Invitrogen) using gene-specific primer sets from Supplemental Tables 6 and 7 online. PCR cycles were optimized for each gene to identify the linear amplification range. Representative results from two biological replicates are shown. Quantitative RT-PCR used first-strand oligo(dT)-primed cDNA, 500 nM of gene-specific primer sets from published cell-type markers or from Rgh3 isofrom specific primers, and 1 × SYBR Green Master Mix (Applied Biosystems). PCR was conducted on a StepOnePlus real-time PCR machine (Applied Biosystems). All quantifications were normalized against the Ubq control gene (Bommert et al., 2005), and data were expressed as 2–Ct (cycle threshold) (Livak and Schmittgen, 2001). Primer specificity was confirmed by agarose gel electrophoresis and melting curve analysis.

**Phylogenetic Analysis**

Predicted protein sequences of URP and U2AF35 were identified through BLASTP and TBLASTN searches in the National Center for Biotechnology Information databases (http://www.ncbi.nlm.nih.gov/), the Joint Genome Institute (www.jgi.doe.gov) servers, and the plaza plant comparative genomics resource (Proost et al., 2009) using default settings at each server. Conserved domains were identified using Prosite scans at the European Bioinformatics Institute server. Multiple sequence alignments and phylogenetic tree of the zinc-finger-UHM-zinc finger region of the proteins were generated with MEGA4 (Tamura et al., 2007). The multiple sequence alignment (see Supplemental Data Set 1 online) was completed with the ClustalW module within MEGA4 using default parameters: gap opening penalty = 10, gap extension penalty = 0.2, protein weight matrix = Gonnet with residue-specific and hydrophobic penalties, gap separation distance = 4, and a 30% delay divergent cutoff. A rooted phylogenetic tree was constructed using the neighbor-joining algorithm within MEGA4 according to a Poisson model with uniform rates among sites and complete deletion of gaps and missing data. Bootstrap values were calculated from 1000 iterations. Protein sequence alignments in Supplemental Figures 7 and 9 online were completed with ClustalW2 at the European Bioinformatics Institute server using default parameters: Gonnet protein weight matrix, gap open = 10, gap extension = 0.2, gap distances = 5, no end gaps = no, iteration = none, numiter = 1, clustering = NJ.

**Accession Numbers**

Sequence data for the maize Rgh3 genomic locus and cDNA clones can be found in the GenBank/EMBL data libraries under accession numbers JN692485, JF681144, and JN791417 to JN791437. The accession number for Arabidopsis URP is At1g10320.

**Supplemental Data**

The following materials are available in the online version of this article.

**Supplemental Figure 1.** Inbred Background Effects on the Phenotype of rgh3 Seedlings 10 d after Planting.

**Supplemental Figure 2.** Endosperm Tissue Culture Response at Multiple Developmental Time Points.

**Supplemental Figure 3.** Developmental Series of the BETL Region in rgh3 and Normal Siblings.

**Supplemental Figure 4.** RT-PCR of Endosperm Cell-Type Markers.

**Supplemental Figure 5.** Mu1 Element Linked to rgh3-umu1.

**Supplemental Figure 6.** Point Mutations in the EMS-Tagged rgh3* Allele.

**Supplemental Figure 7.** Multiple Sequence Alignment of RGH3 Protein Isoforms and Mutant Alleles.

**Supplemental Figure 8.** Relative Expression of Rgh3 Alternative Splicing Events.

**Supplemental Figure 9.** Multiple Sequence Alignment of the Conserved Zinc-Finger-UHM-Zinc-Finger Region of URP and U2AF35 Proteins.

**Supplemental Figure 10.** Transient Expression of AtURP-GFP in Arabidopsis Mesophyll Protoplasts and Coexpression of RGH3α-GFP with AtPRH75-RFP.

**Supplemental Table 1.** Summary of Inferred Seed Developmental Phenotypes from Published B-A Translocation Uncovering Crosses.

**Supplemental Table 2.** Inferred Developmental Phenotypes from B-A Translocation Uncovering Crosses of Uniform Mu rgh Mutants.

**Supplemental Table 3.** Transmission of rgh3.

**Supplemental Table 4.** Bulk Seed Composition Analysis.

**Supplemental Table 5.** Complementation Tests for rgh3 with Seed Loci on Chromosome Arm 5L.

**Supplemental Table 6.** Maize Genes Surveyed for Splicing Defects in rgh3 Mutants.

**Supplemental Table 7.** Primers Used in This Study.

**Supplemental Data Set 1.** Text File of the Sequences and Alignment Used for the Phylogenetic Analysis Shown in Figure 8B.

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Maize *Rough Endosperm3* Encodes an RNA Splicing Factor Required for Endosperm Cell Differentiation and Has a Nonautonomous Effect on Embryo Development

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