Nonredundant Function of Zeins and Their Correct Stoichiometric Ratio Drive Protein Body Formation in Maize Endosperm

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Zeins, the maize (Zea mays) prolamin storage proteins, accumulate at very high levels in developing endosperm in endoplasmic reticulum membrane-bound protein bodies. Products of the multigene α-zein families and the single-gene γ-zein family are arranged in the central hydrophobic core and the cross-linked protein body periphery, respectively, but little is known of the specific roles of family members in protein body formation. Here, we used RNA interference suppression of different zein subclasses to abolish vitreous endosperm formation through a variety of effects on protein body density, size, and morphology. We showed that the 27-kilodalton (kD) γ-zein controls protein body initiation but is not involved in protein body filling. Conversely, other γ-zein family members function more in protein body expansion and not in protein body initiation. Reduction in both 19- and 22-kD α-zein subfamilies severely restricted protein body expansion but did not induce morphological abnormalities, which result from reduction of only the 22-kD α-zein class. Concomitant reduction of all zein classes resulted in severe reduction in protein body number but normal protein body size and morphology.

The zein storage proteins account for 70% of maize (Zea mays) endosperm protein but are devoid of essential amino acids Lys and Trp. As a result, the overall protein quality of maize grain is poor. Paradoxically, many studies have shown that high-level zein accumulation in endoplasmic reticulum (ER)-derived protein bodies normally plays a central role in vitreous endosperm formation, an essential characteristic underlying most maize functional grain properties. This relationship has been summarized in recent reviews (Holding and Larkins, 2006, 2009; Holding and Messing, 2013). Studies to understand this process more completely have been aimed at improving the protein quality of maize and were largely inspired by the opaque2 (o2) mutant. o2 has double the wild-type levels of Lys and Trp (Mertz et al., 1964) because it accumulates low levels of zein proteins and high levels of nonzein proteins that impart a better balance of essential amino acids. The poor agronomic properties associated with kernel opacity prevented implementation of o2, but subsequent breeding efforts led to the development of Quality Protein Maize (QPM) (Vasal et al., 1980; Geetha et al., 1991; Lopes and Larkins, 1991; Holding et al., 2011); the degree of endosperm vitreousness closely correlates with the level of 27-kD γ-zein protein (Lopes and Larkins, 1991), and this increase causes the observed elevation in protein body number in QPM (Lopes and Larkins, 1995; Moro et al., 1995) consistent with the proposed role of 27-kD γ-zein protein body initiation. Some other opaque mutants are the result of aberrations in processes unrelated to protein bodies, such as amino acid biosynthesis, plastid development, and cytoskeletal function (Holding et al., 2010; Myers et al., 2011; Wang et al., 2012). This indicates that further work is needed to generate a more complete understanding of the factors that control endosperm maturation.

Although the highly duplicated nature of the α-zein gene families prevents the identification of loss-of-function mutants, several dominant opaque mutants have been characterized. These result from the accumulation of a defective zein that interferes with normal zein deposition and results in ER stress responses (Coleman et al., 1997; Kim et al., 2004, 2006). The very high-level accumulation of the hydrophobic α-zeins is made possible by their efficient packaging within a shell of cross-linked γ-zeins within ER membranes. The
proteins and their retention in the ER as properly formed protein bodies (Woo et al., 2001; Kim et al., 2002). It is thought that direct interactions between zeins are essential for their retention in the ER as properly formed protein bodies (Kim et al., 2002). Experiments using transgenic expression of both α- and γ-zeins in heterologous species shed some light on their interaction in protein body formation (Williamson et al., 1988; Bagga et al., 1995, 1997; Coleman et al., 1996). However, interpretation of such studies was limited because they did not take into account endogenous nonzein accessory factors such as Flouiry1 (Fig. 1), which are thought to be important determinants of protein body formation (Holding et al., 2007).

We now have the ability to routinely transform maize as well as an ability to selectively eliminate specific zein proteins, multiple products of gene subfamilies, or entire zein classes using RNA interference (RNAi). Previous use of RNAi to selectively eliminate specific γ-zein transcripts and α-zein subfamily transcripts have advanced our understanding of zein function and their inverse relationship to kernel protein quality (Segal et al., 2003; Huang et al., 2004, 2006; Wu et al., 2010; Wu and Messing, 2010, 2011), but questions concerning their distinct roles remain unanswered. This study took advantage of RNAi technology to address the extent of functional nonredundancy within the γ- and α-zein subclasses.

**RESULTS**

**Zein RNAi Lines Abolish Vitreous Endosperm Formation through Discrete Reduction of Zein Transcripts and Proteins**

We used three RNAi transgenes to target all α-zeins, a 27-kD γ-zein, and other γ-zein classes (16-kD γα, 50-kD γβ, and 15-kD β-zeins). We also crossed these RNAi events to derive the pairwise and triplicate combinations that resulted in kernel opacity and the additive effects on zein depletion at the transcript and protein levels. The effects of each RNAi event on transcript level, protein level, and opacity are described in turn. For expression analysis of Z1A, Z1B, Z1C, and Z1D α-zein subfamilies, we used subfamily-specific primers previously described (Feng et al., 2009).

**α-Zein RNAi**

Our strategy for complete knockdown of 19- and 22-kD α-zeins involved a synthetic gene composed of approximately 220-bp conserved regions of a highly expressed member of each of the 19-kD α-zein subfamilies (Z1A, Z1B, and Z1D) and the 22-kD subfamily (Z1C; Fig. 2; Supplemental Fig. S2). Endosperm-specific expression of this RNAi cassette was driven by the 27-kD γ-zein promoter. To measure the effect of the RNAi transgene on endogenous α-zein genes, we used quantitative real-time reverse transcription (qRT)-PCR.
of endosperm RNA from kernels PCR genotyped using DNA extracted from their dissected embryos. The $\alpha$-zein RNAi transgene reduced the level of detectable 19-kD subclass transcripts (Z1A and Z1B) to less than 1% of the level of the wild-type control (Fig. 3, A and B) and Z1D to approximately 2% to 5% of the wild-type control (Fig. 3D). The 22-kD subclass transcripts (Z1C) were reduced to approximately 10% of the wild-type level (Fig. 3C). These reductions were observed both in the single $\alpha$-zein RNAi line and in combination with

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**Figure 3.** qRT-PCR expression analysis of $\alpha$- and $\gamma$-zein genes in RNAi lines relative to wild-type nontransgenic (HiII). Each graph represents the measurement of a different zein gene in all the RNAi lines, which are shown on x axes. Values and SD (shown in insert) are the average of three biological replicate kernels and also incorporated three technical replicates of each measurement.
opaque to vitreous. These results indicate the presence of a single transgene locus and dominant action of the transgene. All opaque kernels showed dramatic reduction in total \( \alpha \)-zeins in mature kernels (Supplemental Fig. S1). To isolate the specific effects of a transgene from nonspecific zein degradation as a result of improper protein body formation, we routinely analyzed zeins at 18 d after pollination (DAP) during midkernel filling (Fig. 4). This corresponds to the latest stage that developing kernels can be reliably fixed and infiltrated for TEM. At 18 DAP, endosperm from kernels containing the transgene accumulated only a small fraction of \( \alpha \)-zein compared with the wild type (Fig. 4E). Curiously, \( \alpha \)-zein RNAi resulted in a noticeable increase in the apparent abundance of 50-kD \( \gamma \)-zeins (Fig. 4E, lanes 2 and 6; Supplemental Fig. S1; data not shown). This 50-kD \( \gamma \)-zein increase was also observed at the transcript level (Fig. 3G).

16-kD \( \gamma \)-Zein, 50-kD \( \gamma \)-Zein, and 15-kD \( \beta \)-Zein RNAi

\( \gamma \)-Zein family members 27-kD, 50-kD, and 16-kD \( \gamma \)-zeins and the 15-kD \( \beta \)-zein share a common location in the protein body periphery and have been assumed to have redundant function. To address the question of functional redundancy, we made transgenic lines containing two different RNAi constructs. The first of these used a synthetic gene composed of 212-bp segments of the 50-kD and 16-kD \( \gamma \)-zeins and the 15-kD \( \beta \)-zein (Fig. 2, Supplemental Fig. S2). This construct, whether alone or in combination with other RNAi constructs, reduced the abundance of all of the three targeted \( \gamma \)-zein proteins to very low levels, usually less than 1% of the wild-type level (Fig. 3, F–H). This construct also reduced to level of the 27-kD \( \gamma \)-zein to approximately 20% to 30% of the wild-type level (Fig. 3E), probably due to the similarity of the 16-kD and 27-kD \( \gamma \)-zeins, and such cross repression was previously described (Wu and Messing, 2010).

We recovered self-pollinated T1 ears, often from several plants per event, from seven events. Six events had ears that apparently segregated three to one, semiopaque to vitreous. However, because the semiopaque phenotype on the light box was subtle and kernel numbers were often low, it was not possible to obtain rigorous segregation ratios. Hill or B73 outcross ears from these events, where recovered, segregated approximately one to one, semiopaque or opaque (Fig. 4B). Although scarce in normal kernels, the 50-kD \( \gamma \)-zein was rendered undetectable in other RNAi constructs. Slight nonspecific reduction in expression of Z1A, Z1B, Z1C, and Z1D class transcripts also resulted from expression of both \( \gamma \)-zein RNAi constructs described below (Fig. 3, A–D). This may indicate that high-level expression of any transgene in the endosperm can reduce endogenous expression of \( \alpha \)-zeins through competition for transcriptional machinery. Alternatively, improper protein body formation caused by \( \gamma \)-zein RNAi-mediated defects may cause negative feedback on expression of \( \alpha \)-zeins.

The \( \alpha \)-zein RNAi construct, we recovered multiple self-pollinated T1 ears from each of 11 events. Ten events had ears with a majority of completely opaque kernels and fewer normal, vitreous kernels. Seed fill was variable and often low in these ears, but we were able to make kernel counts and establish three-to-one opaque-to-vitreous segregation ratios in adequately filled ears from four events (Supplemental Table S1). Hill or B73 outcross ears from these events, where recovered, segregated approximately one to one.
developing and mature kernels of this line. The 16-kD γ-zein ranged from reduced (Fig. 4E, lane 5) to undetectable (Fig. 4E, lane 3) in kernels containing this transgene. Though still detectable, the 15-kD β-zein was substantially reduced in this line. Through sequence similarity to the 16-kD γ-zein and consistent with the reduced transcript level, the 27-kD γ-zein was usually slightly reduced by this transgene, consistent with its reduced transcript level.

27-kD γ-Zein RNAi

We sought to specifically target 27-kD γ-zein knockdown using an RNAi hairpin composed of the entire 27-kD γ-zein open reading frame (ORF; Fig. 2). At the transcript level, this transgene resulted in 1% or less of the wild-type transcript level of both 27-kD and 16-kD γ-zeins (Fig. 3, E and F) and partial suppression (46% of the wild type) of the 50-kD γ-zein (Fig. 3G) and 15-kD β-zein (12% of the wild type; Fig. 3H). At the protein level, this transgene resulted in total suppression of the 27-kD γ-zein as well as substantial suppression of 16-kD γ-zeins (Fig. 4E, lane 4). When combined with the α-zein RNAi, the suppression of the 27-kD γ-zein and the 16-kD γ-zein were less pronounced both at the RNA level (Fig. 3, E and F) and the protein level (Fig. 4E, lane 6). The reduced 50-kD γ-zein transcript correlated with reduced 50-kD γ-zein protein below wild-type levels (Fig. 4E, lane 4), but the protein was present at normal levels in mature kernels (not shown). Interestingly, when combined with the α-zein RNAi, as described above, the suppressive effect of this transgene on the 50-kD γ-zein was outweighed because both the transcript (Fig. 3G) and the protein (Fig. 4E, lane 6) are substantially increased. Despite the fact that this transgene substantially reduced 15-kD β-zein transcript, it did not markedly reduce the level of 15-kD β-zein protein (Fig. 4E, lanes 4 and 6).

We recovered self-pollinated T1 ears, often several ears per event, from four events. Two of these events had ears that apparently segregated three to one, opaque to vitreous. Although this transgene resulted in more complete opacity than 50-kD, 16-kD, and 15-kD RNAi transgenes, light box opacity was not as clear as the α-zein RNAi. This ambiguity, as well as low kernel numbers, again prevented us from obtaining segregation ratios. Hill or B73 outcross ears from these events, where recovered, segregated approximately one to one, opaque to vitreous. The other two events produced self or outcross ears, with all opaque kernels indicating the presence of more than one transgenic locus.

RNAi Effects on Protein Body Density, Size, and Morphology as Well as Zein Distribution Suggest Nonredundant Roles of Zeins in Protein Body Formation

Figures 5, 6, and 7 show low-magnification TEM analysis of protein body density (where density refers to number of protein bodies per unit area, not buoyant density), high-magnification TEM analysis of protein body size and morphology, and high-magnification TEM immunogold analysis of zein distribution, respectively. To account for natural variation in protein body size and density, these analyses were standardized by sampling the fourth subaleurone starchy endosperm cell layer. In our growth conditions, at 18 DAP, the fourth subaleurone cell layer had densely clustered protein bodies in wild-type endosperm, whereas the number and size of protein bodies increased in a gradient in the first through third cell layers. In immunogold TEM, control labeling without primary antibody showed no nonspecific secondary antibody labeling. α-Zein antiserum showed very low labeling outside protein bodies, and the γ-zein antibody showed no labeling outside protein bodies.

α-Zein RNAi

Dramatic reduction of total α-zeins resulted in a similar effect to the δ2 mutant. α-Zein filling was
reduced, which resulted in protein bodies approximately one-quarter to one-third of their normal diameter (Figs. 5B, 6B, and 7, D–F). Unlike α2 however, no reduction in protein body density was observed, indicating that α-zéins are not required for protein body initiation (Fig. 5B; Table I). Normally, α-zéins are located in the center of the protein body and are largely encapsulated by and excluded from an outer peripheral ring of γ-zéins (Fig. 7, B and C). α-Zéin and γ-zéin distribution appeared normal in α-zéin RNAi protein bodies (Fig. 7, E and F), and residual α-zéins were densely packaged within the smaller protein bodies. Notably, unlike selective repression of the 22-kD α-zéin, where the normal ratio of 22- to 19-kD α-zéins is disturbed (Segal et al., 2003), we did not observe any abnormal protein body shape, such as lobing, aggregation, or reticulation.

27-kD γ-Zéin RNAi

Elimination of the 27-kD γ-zéin and the severe reduction of the 16-kD γ-zéin caused by this RNAi transgene resulted in substantial reduction in protein body number per unit area (Fig. 5C; Table I), consistent with the role of the 27-kD γ-zéin in protein body initiation, although this does not rule out the 16-kD γ-zéin as also playing a role in initiation. Interestingly, protein body size was normal in this line (Figs. 5C, 6C, and 7, G–I), suggesting that the 27-kD γ-zéin is not required for the bulk of protein body filling. Minor distortions in protein body shape were usually observed (Figs. 6C and 7, G and H) that may result from α-zéin not being properly encapsulated by γ-zéin and is supported by an increased amount of α-zéin deposited in the protein body periphery and membrane (Fig. 7H). The dark-staining protein body periphery was visibly thinner (Figs. 6C and 7, G–I), and γ-zéin was less abundant, as suggested by a noticeable reduction in the number of gold particles (Fig. 7I).

16-kD γ-Zéin, 50-kD γ-Zéin, and 15-kD β-Zéin RNAi

Severe reduction of these three γ-zéin family proteins, while leaving the 27-kD γ-zéin only slightly reduced, had very different effects. First, protein body number was normal (Fig. 5D; Table I), suggesting the other γ-zéin family members are not required for protein body initiation. However, protein body size was reduced by approximately 50%, suggesting that some or all of these zeins have roles in protein body expansion. Although these smaller protein bodies showed minor shape distortions (Fig. 7, J–L), the dark-staining protein body periphery was of normal appearance (Fig. 7, J–L). Furthermore, α-zéin did not appear ectopically in the protein body periphery, suggesting the 27-kD γ-zéin plays the dominant role in packaging α-zéins and consistent with it being by far the most abundant of the γ-zéin family members.

Total γ-Zéin Family RNAi

Combining the 27-kD γ-zéin RNAi to the 16-kD γ-zéin, 50-kD γ-zéin, and 15-kD β-zéin RNAi resulted in very low levels of all the γ-zéin family proteins (Fig. 3E). Effects of this were additive in terms of severely reduced protein body density (Fig. 5E), reduced protein body size, and moderate to severe protein body distortion or reticulation (Figs. 6E and 7, M–O).

α-Zéin RNAi Combined with Individual γ-Zéin RNAi Events

Combination of the 27-kD γ-zéin RNAi or the 16-kD γ-zéin, 50-kD γ-zéin, and 15-kD β-zéin RNAi to the α-zéin RNAi resulted in protein body size reductions...
similar to the α-zein RNAi alone (Fig. 5, F and G). Although protein bodies in 16-kD γ-zein, 50-kD γ-zein, and 15-kD β-zein RNAi × α-zein RNAi had a slightly reduced density (Table I; Fig. 5G), the 27-kD γ-zein × α-zein RNAi combination resulted in a similar severe loss of protein body density to that of 27-kD γ-zein RNAi alone (Table I; Fig. 5F). This supports the suggestion that the 27-kD γ-zein has the major role in protein body initiation.

**α-Zein RNAi Combined with Total γ-Zein RNAi Events**

To achieve low levels of all the major α- and γ-zeins, we combined all three RNAi constructs. This resulted in the lowest density of protein bodies (Table I; Fig. 5H). However, protein body size ranged from only slightly reduced to normal, compared with the wild type, when all classes of zeins were reduced (Figs. 5H, 6H, and 7, P–R). α-Zein and γ-zein distribution and density in these sparse protein bodies were relatively normal (Fig. 7, Q–R).

**DISCUSSION**

The ability to use RNAi to dominantly reduce or eliminate either single gene or whole gene family products, coupled with routine transformation of maize, has enhanced our ability to decipher the structural functions of zein storage protein classes. Because of the high level of α-zein gene duplication, recessive knockout
mutations for functional analysis are not recoverable, but we now have the technology to selectively or collectively eliminate α-zein subfamilies. This study sought to address unresolved questions about the differential roles of zeins in protein body formation. First, are the colocalized γ- and β-zeins functionally redundant or do they have specific roles? Second, what is the effect of total α-zein knockdown and does the 22-kD α-zein function to constrain 19-kD α-zeins in the protein body center? The data also provided new insight into role of transcript levels in determining zein protein abundance.

**Targeted Zein Transcript Reduction Causes Major Zein Protein Reduction, while Minor Nonspecific Transcript Reductions Do Not Substantially Affect Protein Level**

The chimeric α-zein RNAi construct achieved the targeted reduction of Z1A, Z1B, Z1C, and Z1D transcripts. Z1C (22-kD) class transcripts were reduced to approximately 10% of wild-type levels, and the protein was not reduced to quite the same extent as the 19-kD class (Fig. 4E, lane 2). In the case of the 19-kD α-zein classes, we detected less than 1% of the relative wild-type level of transcript. Although dramatically reduced, reduction of the 19-kD α-zein protein does not appear to be as profound as the transcript. Moreover, we observed a nonspecific transcript reduction of all four classes of α-zein caused by the γ-zein RNAi construct. Despite transcript reductions of around 50%, these nonspecific effects were not seen at the protein level, suggesting that transcript level is in excess and zein abundance is also likely governed by available amino acids and translational machinery. Another nonspecific transcript reduction was that of the 15-kD β-zein, which was caused by both the α-zein RNAi and the 27-kD γ-zein RNAi constructs. Although the basis for these substantial transcript reductions is unknown, they do not result in noticeably lower-than-wild-type levels of 15-kD β-zein protein, again suggesting that this protein is not primarily limited by transcript level.

It was observed that the 27-kD γ-zein RNAi also suppressed the 16-kD γ-zein at the RNA and protein level. However, when combined with the α-zein RNAi, while the 27-kD γ-zein remains suppressed, the 16-kD γ-zein suppression is largely lifted at both the RNA and protein level (Figs. 3F and 4E, lane 6). A similar effect was observed with the 50-kD γ-zein, whose transcript and protein was increased in both the γ-zein RNAi construct lines when combined with the α-zein RNAi. Together, these may suggest that with reduced α-zein synthesis, nonspecific effects of other RNAi events may be less pronounced because of increased availability of translational resources.

**Separable Roles of γ-Zeins in Protein Body Initiation and Filling**

Because γ-zein family member genes are generally not duplicated, it is perhaps surprising that recessive mutants have not been recovered for these, although a double null mutant for the 10- and 18-kD δ-zein was recovered (Wu et al., 2009), which had no observable effects on protein body morphology (Wu and Messing, 2010). With careful design, it is possible to knock down all γ-zein family members individually, but sequence conservation, such as between the 27- and 16-kD γ-zeins, as well as the high cost and time input for working with maize transgenic plants place restrictions on the experimental design. For example, Wu and Messing (2010) chose to simultaneously target the 27- and 16-kD γ-zeins with a single construct, suppress the 15-kD β-zein with another construct, and not target the 50-kD γ-zein because of its low abundance on SDS-PAGE gels. However, using our protocol, we find the 50-kD γ-zein abundance approaching that of the 16-kD γ- and 15-kD β-zeins, and by maturity, it is usually highly abundant (Supplemental Fig. S1). During zein purification, nonzeins are separated by precipitation with 70% (v/v) ethanol (Wallace et al., 1990). When a lower ethanol percentage is used at this step (45%), we have found that, although there is more contamination

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**Table 1. Protein body number per 100 μm² of fourth subaleurone starchy endosperm cell layer**

Protein body counts were made by sampling 100-μm² regions, which excluded nuclei and cell walls, within 18-DAP fourth subaleurone endosperm cells. Average is derived from counts of four TEM sections (two independent kernels and two independent embedments of each kernel).

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Protein Body Number per 100 μm² ± st</th>
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<tbody>
<tr>
<td>Wild type (HiI)</td>
<td>50.0 ± 3.2</td>
</tr>
<tr>
<td>α-Zein RNAi</td>
<td>20.4 ± 3.4</td>
</tr>
<tr>
<td>27- and 50-kD γ-zein RNAi</td>
<td>46.0 ± 3.8</td>
</tr>
<tr>
<td>16- and 50-kD γ-zein and 15-kD β-zein RNAi</td>
<td>12.0 ± 2.1</td>
</tr>
<tr>
<td>α-Zein and 27-kD γ-zein RNAi</td>
<td>25.5 ± 3.4</td>
</tr>
<tr>
<td>α-Zein, 16- and 50-kD γ-zein, and 15-kD β-zein RNAi</td>
<td>36.3 ± 6.6</td>
</tr>
<tr>
<td>α-Zein, 27- and 50-kD γ-zein, and 15-kD β-zein RNAi</td>
<td>15.0 ± 1.6</td>
</tr>
</tbody>
</table>

*Significantly different from the wild type at the 0.01 level.  †Significantly different from the wild type at the 0.001 level using a two-tailed Student’s t test.
with nonzeins, the 50-kD γ-zein band is even more intense (not shown). We therefore targeted the 50-kD γ-zein with RNAi.

The 27-kD/16-kD γ-zein and 15-kD β-zein RNAi events were previously reported to cause only very minor morphological changes to protein bodies (Wu and Messing, 2010). Although an incidence of very small protein bodies was shown in both cases, the majority of protein bodies were of normal size, although standardization of which subaleurone endosperm cell layer was used for TEM imaging was not indicated (Wu and Messing, 2010). Our 27-kD γ-zein RNAi construct was similar because it resulted in complete elimination of 27-kD γ-zein and almost complete elimination of the 16-kD γ-zein. While we also showed that protein body size is normal in developing kernels of this RNAi mutant, later-stage protein body filling is compromised because most protein bodies had a distorted, undulating surface. Our study also incorporated quantitative counts of protein body density. Because protein body density in the developing endosperm was reduced by more than 50%, this suggests that the 27-kD and/or 16-kD γ-zeins are required for protein body initiation. Our 16-kD γ-, 50-kD γ-, 15-kD β-zein constructs completely eliminated the 16-kD γ-zein as well as the 50-kD γ-zein and strongly reduced the 15-kD β-zein. Significantly, this construct did not reduce protein body density, thus indicating that the 16-kD γ-zein does not function in protein body initiation. This construct did result in a significant reduction in protein body size. Because this work and the previous study (Wu and Messing, 2010) showed that RNAi of 27-kD γ-, 16-kD γ-, or 15-kD β-zeins (but not the 50-kD γ-zein) do not markedly reduce protein body size, a significant role for the 50-kD γ-zein in α-zein-driven protein body expansion is supported. Interestingly, we consistently observed increased 50-kD γ-zein (transcript and protein) in α-zein RNAi lines, suggesting that the 50-kD γ-zein has a closer association to the central α-zein core than other γ-zeins and can partially compensate the loss of α-zeins. Combination of the γ-zein family RNAi constructs resulted in addition of the distortion and the low-density phenotypes. The combined RNAi in the previous study (Wu and Messing, 2010; 27-kD γ-, 16-kD γ-, and 15-kD β-zeins) did not target the 50-kD γ-zein and again did not reduce protein body size, which further supports a major role for the 50-kD γ-zein in protein body filling. Taken together, our results suggest nonredundant initiation and filling roles of γ-zeins.

Reduction of Both α-Zein Classes Affects Protein Body Size but Not Number or Morphology

At the outset of this work, previous efforts to use RNAi to eliminate α-zeins had been aimed solely at creating a dominant, nonpleiotropic low-zein line for Lys improvement. The first targeted only the 22-kD α-zein (Segal et al., 2003), and the second was private-sector work that targeted both the 19- and 22-kD α-zein but did not address the morphological effects on protein bodies (Huang et al., 2006). We required such a total α-zein RNAi line both as a nonpleiotropic, dominant opaque line for testing the activity of overexpressed o2 (QPM) modifier genes as well as to try to dissect possible nonredundant function between the 19- and 22-kD α-zeins. A total α-zein RNAi line that incorporated a dominant GFP marker was subsequently published, but morphological protein body data were not reported (Wu and Messing, 2011).

Protein bodies in o2 endosperm are small (Geetha et al., 1991), presumably because of reduced accumulation of both 22-kD and 19-kD α-zeins, the former resulting from reduced transcriptional activation and the latter as an indirect effect (Schmidt et al., 1990). Protein body density in cells is also reduced (Geetha et al., 1991), consistent with the fact that the mutation also indirectly reduces 27-kD γ-zein accumulation. However, o2 exhibits no misshapen protein body morphologies, as is seen in dominant negative α-zein mutants (Coleman et al., 1997; Gillikin et al., 1997; Kim et al., 2004). The 22-kD α-zein-specific RNAi event, which has substantial residual 19-kD α-zein, also has abnormally lobed and distorted protein bodies whose size is not significantly reduced over the wild type (Wu and Messing, 2010). This leads to the suggestion that the bulk of protein body expansion is driven by the 19-kD α-zein, which is consistent with its location throughout the protein body core within a ring of the 22-kD α-zein (Holding et al., 2007). This also suggests that the 22-kD α-zein may have a structural role in packaging the 19-kD α-zeins and may explain why protein body lobing and distortion result from specific reduction of the 22-kD α-zein. In support of this hypothesis, when we reduced both α-zein classes equivalently with RNAi, protein body morphology and zein distribution were normal and only a reduced size resulted.

Our results show that reduced protein body size results from reduced total α-zein as well as reduction in some but not all γ-zein classes. Reduced protein body initiation results specifically from the reduced 27-kD γ-zein and not from reductions in other γ-zein family members. However, when all the RNAi events were combined, all zeins were reduced roughly in proportion. Surprisingly, this resulted in protein bodies of relatively normal size and shape but at very low density. This indicates that correct stoichiometric ratio of the different zeins is essential for vitreous endosperm formation through correctly formed, concentrically organized protein bodies (Fig. 1).

MATERIALS AND METHODS

Constructs

The α-zein RNAi cassette harbored a synthesized gene (GenScript) containing 220-bp conserved regions derived from the GenBank accession numbers EU116446, EU116472, EU116497, and EU116477 for the 19- and 22-kD subfamilies. The synthetic gene was assembled into the hairpin element by the vector pUC18-RNAi (gift from Heriberto Cerutti [University of Nebraska-Lincoln]), a plasmid that carries the second intron of the Arabidopsis (Arabi-
dopsis thaliana) small nuclear ribonucleic protein D1 (locus AT4G2840). The intron is delineated by a set of compatible cohesive end restrictions sites that facilitate assembly of inverted repeats. The hairpin structure was subsequently
subcloned between the 27-kD γ-zein promoter and T3SS terminator of transcription within the binary vector pPZP212 (Hajdukiewicz et al., 1994).

For genotyping plants, a primer in the Z1A, Z1B, Z1D, and Z1C gene (5'-GCCAGAAGGGTCTCAGAC) and T3SS poly A sequence (5'-ATGCTCAACATGACCGAA) were used to give a 311-bp band.

For the 50-kD γ-, 16-kD γ-, and 15-kD β-zein RNAi, a synthetic gene was made and inserted with the 27-kD γ-zein promoter in pPZP212 as described above. For the synthetic gene, the regions were isolated from GenBank accession AF371263 (50-kD), DQ400403 (16-kD), and NM_01112534 (15-kD). For genotyping plants, a 27-kD γ-zein promoter primer (5'-GATGAACCATGACCGTACA) was used with a primer within the synthetic RNAi gene (5'-CCATGGTGAGCCTCAGCC) for insertion into the RNAi hairpin structure. For genotyping primers in the 27-kD γ-zein, ORF (5'-ACTGACCGCAGAATCTCG) and T3SS poly A sequence (5'-ATGCTCAACATGACCGAA) were used to give a 539-bp band.

For the pZ1C:27γ construct, the 27-kD ORF was inserted with the α-zein ZIC promoter in pPZP212. For genotyping, a primer with the ZIC promoter (5'-CGAATGCGATGATCCGATA) was used with a 27-kD γ-zein ORF primer (5'-GGGC CGTGGTAGGACTGAT) to give a 486-bp band. Constructs were transformed into Agrobacterium tumefaciens and integrated using an established procedure (Sattarzadeh et al., 2010).

Genotyping, Zein Protein Analysis, and Fixation for Microscopy

Given the highly readily discernible effects of reducing individual zein species at the protein level, we used protein level as the criterion for selecting lines for TEM analysis. For RNAi lines with dominant phenotypes, opaque T2 kernels were planted. RNAi and the pZ1C:27γ plants were selected for the presence of the transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008). For the 18-DAP stage, kernels were removed, transgene using leaf genomic PCR using DNA extracted according to the urea method (Holding et al., 2008).

Zein Gene Expression Analysis

We extracted endosperm RNA, DNase treated, made complementary DNA, and measured the relative levels of all α- and γ-zein subclasses using qRT-PCR as described previously (Holding et al., 2011) with the following modifications. The control gene used in this case was tubulin. RNA was extracted from individual halves of developing endosperm from frozen kernels derived from plants already genotyped as described in the previous section. The other half was used for zein extractions to confirm identity (not shown), and each kernel was genotyped using the dissected embryo as described in the previous section (not shown but illustrated in kernels used for TEM and protein analyses, Fig. 4). In these analyses, we compared RNA from three kernels positive for the transgene(s) from the same ear and, thus, the same event to exemplify the effect of each transgene. While expression levels could vary between different events of each RNAi construct, the purpose here was to exemplify the effects of each transgene on the expression of all zeins. However, for the wild-type control RNA, each of the three kernels was taken from separate ears to be sure to exclude ear-to-ear variation that might otherwise bias the relative expression levels of all RNAi construct lines. We averaged the three technical replicates first, and then the final average is between the biological replicates zeins. Zein primers are listed in Supplemental Table S2.

TEM

Selected kernel segments were further dissected following fixation to remove the apical and basal kernel portions, leaving two 1-mm midkernel pieces with pericarp still attached for orientation. Samples were postfixed by 1% osmium tetroxide. Samples were then dehydrated in a graded series of ethanol (30%, 50%, 70%, 95%, and 100%), followed by 100% acetone. Dehydrated samples were infiltrated in Epon 812 resin (Electron Microscopy Sciences) using a resin:acetic (1.2 and 2:1) and 100% resin series before polymerization at 60°C overnight. Ninety-nanometer sections were cut using a Diatome diamond knife and a LKB Ultrotome III microtome and attached to carbon- and Formvar-coated grids (Electron Microscopy Sciences). Sections were stained with 4% lead citrate and 3% uranyl acetate and air dried before being viewed on a Hitachi 7500 TEM.

Immunogold TEM

Selected kernel segments were further dissected and dehydrated as for regular TEM, except that the osmium tetroxide post fixation and acetone infiltration were omitted, and the samples were infiltrated in LR White resin (Electron Microscopy Sciences) before sectioning as for TEM. For immunogold labeling, grids were hybridized in 50 µL drops on Parafilm strips as follows. The grids were incubated for 30 min in blocking solution (pH 8.2) containing 0.2% bovine serum albumin and 0.06% Tween 20 in 20 mM Tris-HCl and 500 mM NaCl. Rabbit primary antisera were added (α-zein at 1:1,000 and γ-zein at 1:100) and incubated overnight at 4°C before three 10-min washes in 100 µL blocking solution. Secondary antibody (goat anti-rabbit, 15 nm, gold-conjugated; Electron Microscopy Sciences) was added at 1:20 dilution for 1 h at room temperature. The grids were washed three times and rinsed three times in deionized water and air dried. Grids were poststained in 4% lead citrate and 3% uranyl acetate and air dried again before being viewed on a Hitachi 7500 TEM.

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers EU116446, EU116472, EU116497, and EU16477 for the 19- and 22-kD alpha zein genes represented in the chimeric alpha zein RNAi gene. For the gamma zein genes represented in the chimeric RNAi gene, accession numbers are AF371263 (50-kD), DQ400403 (16-kD), and NM_01112534 (15-kD).

Supplemental Data

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

Supplemental Figure S1. Purified zeins from transgenic mature kernels.

Supplemental Figure S2. Design of synthetic zein RNAi transgenes.

Supplemental Table S1. Segregation ratios of opaque versus vitreous kernels from T1 ears of selected α-zein RNAi events.

Supplemental Table S2. Primers for qRT-PCR of zein expression.

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LITERATURE CITED


