Carotenoids modulate kernel texture in maize by influencing amyloplast envelope integrity

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The mechanism that creates vitreous endosperm in the mature maize kernel is poorly understood. We identified *Vitreous endosperm 1* (*Ven1*) as a major QTL influencing this process. *Ven1* encodes β-carotene hydroxylase 3, an enzyme that modulates carotenoid composition in the amyloplast envelope. The A619 inbred contains a nonfunctional *Ven1* allele, leading to a decrease in polar and an increase in non-polar carotenoids in the amyloplast. Coincidently, the stability of amyloplast membranes is increased during kernel desiccation. The lipid composition in endosperm cells in A619 is altered, giving rise to a persistent amyloplast envelope. These changes impede the gathering of protein bodies and prevent them from interacting with starch grains, creating air spaces that cause an opaque kernel phenotype. Genetic modifiers were identified that alter the effect of *Ven1A619*, while maintaining a high β-carotene level. These studies provide insight for breeding vitreous kernel varieties and high vitamin A content in maize.

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Maize kernel texture is a critical agronomic trait. It is influenced by the ratio of vitreous endosperm in the outer part of the kernel to the starchy endosperm in the center of the kernel\(^1\). Vitreous endosperm strengthens the kernel and protects the grains from mechanical damages during harvesting and shipping, whereas starchy endosperm is breakable and fragile, and is susceptible to pests and diseases\(^2\). Even though teosinte (wild maize) has a hard glume encasing the seed, it contains a large portion of vitreous endosperm. In this respect, there may have been evolutionary selection for vitreous endosperm in teosinte to enhance kernel integrity and passage through the gut of birds and other animals to aid seed dispersal. Maize grains containing a greater amount of vitreous endosperm have a higher kernel density and test weight and lower flotation indices\(^3,4\). The vitreous region is also important for food processing, including making of grits and cornflakes, and other food products\(^5\), and it is essential for making popcorn pop.

What creates vitreous endosperm is a longstanding question with many hypotheses, most of which are associated with synthesis of starch and storage proteins, and the factors affecting them. The major storage metabolites in the endosperm are starch and proteins, which account for 70 and 10%, respectively, of the endosperm dry mass. Starch is synthesized and accumulated as starch grains (SGs) in amyloplasts. The major proteins are storage proteins, prolamin, collectively called zeins. Zeins contain a signal peptide that directs their synthesis into the lumen of the rough endoplasmic reticulum (RER), where they become organized into accretions called protein bodies (PBs)\(^6,7\). During kernel maturation, SGs become surrounded by a proteinaceous matrix composed of PBs and amorphous cytoplasmic proteins. As the seed undergoes desiccation, the integrity of organellar membranes is lost and the peripheral endosperm cells, filled predominantly with PBs and SGs, form vitreous endosperm, while central endosperm cells, containing primarily SGs and few PBs, form the starchy endosperm\(^1\). Although a large number of mutations affecting zein proteins and starch synthesis have been identified and provided insight into the mechanisms that create vitreous endosperm\(^8,9\), little is known about quantitative trait loci and their metabolic products that explain the genetic variation in kernel texture found in natural populations.

Maize inbred lines have great natural variation in kernel texture, ranging from nearly completely vitreous to completely opaque. W64A and A619 are typical yellow dent inbred lines that illustrate the natural variation for a normal amount and a small amount of vitreous endosperm. In this work, through backcrossing, we are able to identify and clone the gene responsible for a major QTL, \textit{Ven1}, which affects kernel texture by regulating the \(\beta\)-carotene content. High levels of \(\beta\)-carotene appear to prevent the breakdown of amyloplast membranes and affect the quantity and composition of lipids in desiccating endosperm cells. This disrupts interactions between SGs and PBs, leading to air spaces and, consequently, opacity of the mature endosperm.

### Results

**\textit{Ven1} is a principal QTL controlling maize kernel texture.**

Maize inbred lines W64A and A619 display differences in endosperm vitreousness (Fig. 1a–d). When measured, W64A kernels contain 80% vitreous endosperm, whereas A619 kernels have only 20% (Fig. 1c, d). On a light box, light transmission was dramatically reduced compared with wildtype kernels, with more backcrossing, an equal segregation of vitreous and opaque kernels clearly occurred in F\(_1\)BC\(_3\) (Supplementary Fig. 2c). These results indicate the penetrance of the \(\text{Ven1\(^{W64A}\)}\) allele is strongly influenced by the genetic background. F\(_1\)BC\(_3\) seeds with vitreous and opaque phenotypes were propagated by self-pollination for two generations, yielding the homozgyous nearly isogenic lines (NILs), NILW64A and NILA619. The NILs were nearly identical in other plant phenotypes but exhibited an apparent difference in kernel texture (Fig. 1a–d). Although NILW64A contained a slightly smaller amount of vitreous endosperm than W64A, it had significantly more than A619 and NILA619 (Fig. 1c, d), thereby creating stronger light transmission on a light box (Fig. 1b).

We pooled the vitreous and opaque seeds segregating in the F\(_1\)BC\(_2\) population of the W64A by A619 cross (Supplementary Fig. 2f) and performed Bulked Segregant Analysis (BSA) sequencing. This revealed only one candidate region located on chromosome 10 (Fig. 1e). Using 2000 F\(_1\)BC\(_2\) individuals, the QTL was narrowed down to a 0.2 Mb region (Fig. 1f), where five genes were annotated based on the B73 genome. RNA sequencing (RNA-seq) analysis of 18-DAP (day after pollination) endosperms of W64A, A619, NILW64A and NILA619 revealed only Zm00001d026056 was differentially expressed in W64A and NILW64A, versus A619 and NILA619; the other four genes were expressed at a similar level or expressed at low levels in the endosperm of these lines (Fig. 1g). We amplified and sequenced the gene body and regions within 1.2 kb up- and downstream of the coding region and found the \(\text{Ven1\(^{A619}\)}\) allele contains a 1250-bp deletion in the 3’ end, resulting in truncation of the \(\text{Ven1\(^{A619}\)}\) protein (Supplementary Fig. 3). Using a specific primer pair flanking this deletion\(^10\), we verified that \(\text{Ven1\(^{A619}\)}\) was also associated with the opaque phenotype in the F\(_1\)BC\(_2\) populations created by A619 crosses with D1051 and P25 (Supplementary Fig. 4). Consistent with the transcript levels, the \text{VEN1} protein was barely detectable in A619 and NILA619, compared with W64A and NILW64A (Fig. 1h), indicating \(\text{Ven1\(^{A619}\)}\) is a nonfunctional allele.

To confirm that this mutation is responsible for the opaque endosperm phenotype of A619, we used RNA interference (RNAi) technology to knock down \textit{Ven1} expression in kn5585 (Supplementary Fig. 5a), a vitreous inbred line wildly used in China for maize transformation. Twenty independent \(T_0\) \text{Ven1RNAi} plants were recovered and all segregated vitreous and opaque seeds when self-pollinated. The transcript and protein levels of \textit{Ven1} in progeny bearing the RNAi transgene were dramatically reduced compared with wildtype kernels, as exemplified in photographs of two events (Fig. 1i, j). A representative ear is shown in Fig. 1k and Supplementary 5b, where three quarters of the seeds inherited the RNAi construct and exhibited the opaque endosperm phenotype (Supplementary Fig. 5b). Homozygous \text{Ven1RNAi} ears showing a fully opaque...
seed set were obtained after propagating the transgenic seeds (Supplementary Fig. 5c). When NILW64A ears were fertilized by the pollen from heterozygous VenIRNAi/+ plants, vitreous and opaque F1 seeds segregated at a 1:1 ratio (Fig. 1l, Supplementary Fig. 5d). These genetic data showed that Zm00001d026056 is the gene responsible for the opaque endosperm phenotype in A619.

**Ven1A619 causes over-accumulation of nonpolar carotenoids.** Quantitative RT-PCR revealed Ven1 is highly expressed in W64A endosperm, where RNA transcript levels gradually increased after 12 DAP, reaching a peak at 35 DAP (Fig. 2a). RNA in-situ hybridization showed Ven1 was predominantly expressed in starchy endosperm cells from the crown to the kernel center (Fig. 2b). To determine its subcellular localization, the VEN1 protein was fused to GFP and the resulting construct transiently expressed in Nicotiana benthamiana leaf cells. The signal from VEN1-GFP was observed to be localized to chloroplasts (Fig. 2c). Immunohistochemical assays clearly showed the distribution of VEN1 is restricted inside the amyloplast envelope in endosperm cells (Fig. 2d–f).

**Ven1 encodes β-carotene hydroxylase 3 (HYD3),** which hydroxylates the β-ionone ring of α/-β-carotene, producing xanthophylls (e.g. lutein and zeaxanthin). We measured the
total carotenoid content in 24- and 35-DAP endosperms and found it was not apparently different between NILW64A and NILA619 (Table 1). However, we found that the nonpolar carotenoids such as phytoene, lycopene, α-carotene and β-carotene, molecules produced upstream of HYD3, accumulated at a significantly higher level in NILA619 than NILW64A, whereas the levels of the polar carotenoids, like lutein and zeaxanthin, downstream of HYD3, were the inverse (Table 1).

**Ven1A619 influences SG-PB interactions.** To investigate the mechanisms by which Ven1A619 affects vitreous endosperm
Table 1 | Altered carotenoid composition in endosperm tissues of NILW64A and NILA619.

| Nonpolar carotenoids | Polar carotenoids | Total |
|----------------------|------------------|-------|
| α-(E/Z)-Phytoene     | Lycopene         | β-Carotene |
| NILW64A-24 DAP       | 2.38 ± 0.19      | 0.012 ± 0.004 |
| NILA619-24 DAP       | 6.80 ± 0.47**    | 0.96 ± 0.20** |
| NILW64A-35 DAP       | 4.03 ± 1.10      | 0.01 ± 0.002 |
| NILA619-35 DAP       | 3.13 ± 0.41      | 0.96 ± 0.04   |

** and *** significant differences at < 0.05 and < 0.01 in two-sided Student’s t-tests; mean and SD were calculated. Source data are provided as a Source Data file.

Ven1A619 affects the integrity of amyloplast membranes and lipid composition. To investigate the process of SG and PB interaction during desiccation of NIL-Ven1A619, we examined the formation, we first examined zein protein synthesis. Ven1A619 affects PB and SG development, and subsequently their interaction (Supplementary Fig. 8). At 24 DAP, TEM revealed that SGs and PBs developed normally and showed no apparent difference in size and number between NILW64A and NILA619 (Supplementary Fig. 8a and c). This is consistent with the observation that stalk and zein proteins were similarly accumulated in the NILs. When the endosperm cells were examined by SEM, we found that SGs in NILW64A were tightly surrounded by PBs, appearing to be embedded in a proteinaceous cytoskeletal matrix. In NILA619, the majority of SGs were contacted by a small number of PBs and they maintained their spherical morphology (Supplementary Fig. 8a, c). When observed by TEM, the SGs in NILW64A became embedded in a dense matrix of PBs and cytoplasmic contents. Probably due to tight compacting, the SGs began to form a polygonal shape. In NILA619, the majority of SGs were contacted by a small number of PBs and they maintained their spherical morphology (Supplementary Fig. 8g). Thirty five DAP is a critical time point for vitreous endosperm formation. By then most of the starchy endosperm cells have undergone programmed cell death and storage metabolite synthesis is slowing down, water content is rapidly decreasing, and the endosperm is entering the dough stage. In NILW64A, TEM showed that all the PBs converged together, creating a continuous network of matrix grids around the SGs; these grids, with SGs capsuled inside, appeared to function as the basic unit of vitreous endosperm formation. One could imagine that the later-formed vitreous endosperm is composed of numerous interconnecting units like this. In contrast, this grid structure was not organized in NILA619 (Supplementary Fig. 8i, k). In SEM, the SGs in NILW64A were observed to be tightly packed, with PBs developing distinctive features of vitreous endosperm formation, whereas the SGs in NILA619 were loosely packed with air space between them, a typical characteristic of opaque endosperm (Supplementary Fig. 8j, l). These results demonstrated that compaction of PBs and SGs is affected during desiccation of NILA619 endosperm.
organelles by TEM at higher resolution. Amyloplasts are a specialized plastid for starch synthesis and are bounded by two lipoprotein membranes. At 18 DAP, when the SGs in NILW64A were undergoing rapid development, we observed that the amyloplast membranes were intact, indicating they are functional for starch synthesis (Fig. 3a). At 24 DAP, when many SGs in NILW64A were reaching maturity, the amyloplast membranes began to break down, as evidenced by debris in their periphery (Fig. 3b). Because SGs at this stage differed in maturity, those with membranes in integrity or complete degradation (in the latter, the debris was absent in their periphery) were also observed (Supplementary Fig. 9a). At 35 DAP, the amyloplast membranes in NILW64A completely disappeared, enabling PBs to physically interact with SGs (Fig. 3c). It appeared that PBs were pulled together by undefined cytoplasmic components, perhaps by liposome-like structures, through hydrophobic interactions (Fig. 3d). In contrast, the amyloplast membranes in NILA619 behaved differently. At 18 DAP, the membranes appeared to be irregularly expanded, leading to formation of an amorphous structure (Fig. 3e). The membranes in NILA619 appeared to be more persistent; they were intact at 24 DAP (Fig. 3f) and remained intact at 35 DAP (Fig. 3g). It appears the stability of the membranes creates a barrier that prevents PBs and the cytoplasmic matrix from approaching the SGs. There was little of the
dense cytoplasmic contents around PBs compared to NILW64A (Fig. 3h). These results suggested Ven1A619 somehow influences the breakdown of amyloplast membranes, which is critical for allowing for SG-PB interaction.

Isolated amyloplast membranes contain about 70% galactolipid and 25% phospholipid (PL), but no neutral lipids like triacylglycerides (TAG) and fatty acids (FA)\textsuperscript{13}. Monogalactosyldiacylglycerol (MGDG) and digalactosyldiacylglycerol (DGDG) provide both structural and functional organization of amyloplast. We measured the quantity and composition of lipids in the endosperm at 24 and 35 DAP and found they were markedly different between NILA619 and NILW64A (Fig. 3i–k). In the endosperm, there are mainly neutral lipids, e.g., FA and TAG. In each NIL, total lipid content was not apparently altered at the two time points, but it was always significantly higher in NILA619 than NILW64A (Fig. 3i), suggesting an enhanced capacity for lipid synthesis. At 24 DAP, when amyloplast membranes of many SGs began to break down in NILW64A, only a few polar lipids (PA, LPC and LPE) accumulated at a mildly higher level in NILA619 than NILW64A (Fig. 3j, k). This suggested that although the breakdown of amyloplast membranes occurred in NILW64A, their main lipids, i.e. galactolipids were still present in endosperm cells. The apparent difference between the two NILs were the levels of FA and TAG (Supplementary Fig. 9c). By 35 DAP, when amyloplast membranes of most SGs completely degraded in NILW64A, the level of DGDG, which is a predominant galactolipid in amyloplast membranes, was significantly higher in NILA619 than NILW64A, as well as PL (PC, PE, PG) and lysOPL (LPC, LPE, LPG) (Fig. 3k, Supplementary Fig. 9b). It was evident that the contents of amyloplast lipids, MGDG, DGDG and other membrane phospholipids (PA, PC, PE and PG), were reduced at 35 DAP compared to those at 24 DAP in the two NILs. At this stage, FA and diacylglycerides (DAG, the minor neutral lipid constituent) were significantly higher in NILA619 than NILW64A, although TAG was not apparently different (Supplementary Fig. 9c–e). The difference in lipid composition between NILW64A and NILA619 is consistent with the differential status of amyloplast envelope integrity (Fig. 3b, c, f and g; Supplementary Fig. 9a).

Epistatic effect of carotenoid synthesis on the opaque phenotype in A619. To further investigate whether over-accumulation of non-polar carotenoids in A619 is associated with the opaque endosperm phenotype, we treated A619 with ethyl methane-sulfonate (EMS) and screened for mutants with defects in carotenoid synthesis by observing endosperm color. We identified four single-gene recessive mutants displaying varying differences in endosperm color, suggesting these mutations occurred in different steps of carotenoid synthesis (Fig. 4a–e). A common feature of these mutants was the conversion of the opaque to a vitreous endosperm phenotype. Because these mutations function as suppressors of the Ven1A619 phenotype, we designated them vitreous endosperm suppressor 1, 2, 3-1 and 3-2 (ves1, ves2, ves3-1 and ves3-2, the last two were subsequently shown to be allelic mutations) (Fig. 4f–g). We measured the carotenoid composition and found the synthesis of lycopene, α-carotene and β-carotene was suppressed in all these mutants (Fig. 4h), confirming a close relationship of the excessive accumulation of non-polar carotenoids and the opaque phenotype in A619 endosperm. We then cloned these genes by BSA sequencing (Supplementary Fig. 10a–d). The carotenoid biosynthesis pathway is conserved in maize and other plants, and the candidate genes, quantitative trait loci, and phenotypic loci are well characterized\textsuperscript{11,14–19}. According to these reports, we summarized the maize carotenoid biosynthesis pathway and included the suppressors in Supplementary Fig. 10e. Ves1 encodes 4-hydroxyphenylpyruvate dioxygenase 1 (HPPD), which is the first committed step in synthesis of both plastoquinone and tocopherols in plants and functionally reverses the pds1 mutant phenotype in Arabidopsis\textsuperscript{20} (Supplementary Fig. 10e). The enzyme directly downstream of HPPD is homogenitase solanesyl transferase (HST), and its mutation in maize, w3, showed blocked carotenoid synthesis by repressing the synthesis of PQ9, the oxidant required for phytene desaturase (PDS). The w3 mutant accumulates phytene and exhibits a white kernel and seeding phenotype. Ves2 (also known as seed carotenoid deficient, Scd) encodes a hydroxymethylbutenyl diposphosphate synthase (HDS) converting 2C-methyl-D-erythritol 2,4-cyclodiphosphate to 1-hydroxy-2-methyl-2-(E)-butenyl 4-diphosphate. Mutation of Scd inhibits the biosynthesis of MEP-derived isoprenoids (carotenoids, chlorophylls, and tocopherols) and results in a white kernel, albino seeding and eventually lethal phenotype\textsuperscript{21}, similar to the ves2 phenotypes. Ves3 encodes the 1-deoxy-D-xylulose-5-phosphate synthase (DXS), which is in upstream of UES2, a limiting enzyme for 2-C-methyl-D-erythritol 4-phosphate (MEP) for plant plastidic isoprenoid biosynthesis (Supplementary Fig. 10e).\textsuperscript{22,23} Sequencing revealed that ves1, 2 and 3-1 resulted from a premature stop codon in the second exon (Supplementary Fig. 10a–c) and ves3-2 was a missense mutation of DXS, in which Arg-210 was changed to Trp (Supplementary Fig. 10d). In contrast with the persistent amyloplast membranes in A619, TEM showed that the amyloplast membranes completely degraded in endosperm cells of ves1, 2, 3-1 and 3-2 at 30 DAP, which allowed the PBs to converge around SGs and surround them (Fig. 4i). SEM revealed that the SGs in all suppressor mutant endosperms were tightly compacted with PBs and cytoplasmic contents (Fig. 4j). These results genetically supported that the upstream carotenoid biosynthetic pathway is epistatic to the Ven1A619 phenotype.

Screening Ven1A619 endosperm modifiers. During recurrent introgression of VenW64A into A619, we did not observe segregation of the opaque phenotype until F3BC\textsubscript{3} (Supplementary Fig. 2a–c). This was different from the cross of NILW64A and Ven1RNAI/+ , where the F\textsubscript{1} hybrids of W64A and Ven1RNAI/+ were vitreous (Supplementary Fig. 11), suggesting the presence of genetic modifiers in W64A that influence the opaque endosperm phenotype (caused by Ven1A619 or Ven1RNAI). We analyzed the Ven1 genotype in the 262 inbred lines (Supplementary Data 1) and found that 14 bear the Ven1A619 allele, of which 12 are vitreous and only two have an opaque endosperm in the mature kernel (Supplementary Fig. 12a). To identify Ven1A619 modifiers (Vem) in the natural population, we screened the 262 lines by crossing them with Ven1RNAI/+ , no matter whether they are of the Ven1A619 genotype or not. If the inbred lacks modifiers, the F\textsubscript{1} progeny should segregate vitreous and opaque kernels at a 1:1 ratio; if does, all the progeny are expected to be vitreous. This screen resulted in identification of varying degrees of phenotypic modification in the natural population, wherein 182 were fully modified, 35 partially modified (showing a mosaic phenotype) and 45 totally unmodified (Supplementary Fig. 12a, b; Supplementary Data 1). Examples for each kind of modification are shown in Fig. 5a–c. These data demonstrated that a great proportion of germplasm (30%) in the natural population are not suitable for provitamin A biofortification due to incomplete modification, while Ven1A619 is a rare allele that has potential to increase β-carotene in maize grain. A genome-wide association study (GWAS) for modifiers identified six obvious peaks (\(P = 5 \times 10^{-6}\)) distributed on chromosome 1, 3, 6 and 8 (Fig. 5d). Genetic variation and
RNA-seq transcriptome analyses of W64A and A619 endosperms at 18 DAP revealed that nine genes in these regions were highly expressed in W64A, but lowly expressed in A619 (Supplementary Table 1). Among them, glutathione S-transferase 1 catalyzes conjugation of glutathione (GSH; γ-Glu–Cys–Gly) with electrophilic compounds, such as ROS and GSH, becoming an oxidized form, glutathione disulphide (GSSG)24. Therefore, higher expression of \textit{Gst1} could alter the redox state in endosperm cells, which may in turn affect the stability of amyloplast membranes and SG-PB interaction during vitreous endosperm formation.

**Discussion**

One of the first descriptions of cells in the vitreous endosperm was by Donald Duvick, who likened them to a box of white marbles (SGs) in which buckshot (PBs) was packed between the marbles and viscous cytoplasm functioned as a transparent glue between them, forming a structure like rigid concrete25. In this model, mutations affecting synthesis of SGs and zein PBs would affect the formation of vitreous endosperm9. Indeed, the starch mutants, \textit{waxy1} and \textit{amylose extender1}, with a low- and high-amylose content in SGs, develop an opaque and nearly fully vitreous endosperm phenotype, respectively. Mutations either reducing zein protein synthesis or altering its deposition in PBs also create an opaque phenotype in the mature endosperm. Besides these mutations that directly affect starch and PB synthesis, there are many others that indirectly affect their synthesis and cause an opaque endosperm phenotype26–28. Evidence from these studies support the hypothesis that the number of PBs, the organization of zein proteins within them, and their interaction with SGs are essential factors for vitreous endosperm formation9. However, our understanding of the process by which vitreous endosperm forms...
is incomplete, as other factors can influence it. For example, some mutations, like *opaque1*, have no apparent effect on zein or starch synthesis\(^{29}\). Indeed, the chemistry that creates the matrix between PBs and SGs during kernel desiccation is not understood. In this study, we demonstrated that *Ven1*, a QTL affecting the nature of carotenoids in the amyloplast and the stability of the amyloplast membrane, impacts interactions between SGs and PBs. An influence of the amyloplast membrane on vitreous endosperm formation was reported for the *opaque5* mutation\(^{30}\). Although the reason is unclear, this mutation affects mono-galactosyldiacylglycerol synthase, the enzyme responsible for synthesis of amyloplast membrane galactolipids. Whether or not this influences amyloplast membrane stability during kernel desiccation was not investigated in detail.

Vitreous endosperm begins to form at the periphery of the kernel by the late dough stage (35–40 DAP). This process could simply be a consequence of condensation of PBs and cellular contents onto SGs. At seed maturity, endomembrane systems would be expected to degrade as a result of PCD and water loss. On the one hand, during vitreous endosperm formation the amyloplast envelope appears to break down before PB membranes (Fig. 3b, c). This would be critical to facilitate PBs and cytoplasmic contents condensing onto the SG surface and creating the matrix grid (Supplementary Fig. 8i, j; Fig. 3c). Our results support the hypothesis that the SG-PB interaction requires degradation of amyloplast membranes, which otherwise would form a barrier to this process (Supplementary Fig. 8k, l; Fig. 3f, g).

On the other hand, PBs appear to stay in positions surrounding SGs around the time amyloplast membranes are about to degrade. Timing could be critical to achieve an optimal spatial arrangement of SGs and PBs, otherwise once the starchy endosperm tissue is dehydrated, PBs are unable to approach SGs and condense around them.

While the number and size of PBs matters for vitreous endosperm formation, the assembly of PBs is also a determining factor, as suggested by the *opaque1* and *floury1* mutations, which are associated with defects in synthesis of myosin or myosin-related proteins, part of the cytoskeleton surrounding the RER\(^{32}\).

**Fig. 5 Screening *Ven1* endosperm modifiers by GWAS and proposed model for vitreous endosperm formation.** a–c Representative ear phenotypes showing full modification (a), partial modification (b) and totally unmodified (c) pollinated by *Ven1RNAi/+*. d Manhattan plot of whether the endosperm contains *Ven1* modifiers. The dashed line indicates the significance threshold of *P*-value $5 \times 10^{-8}$. 27 unique SNPs are labelled with red dots, and the candidate genes are highlighted. The numbers on the horizontal axis indicate the maize chromosomes. e Proposed model for vitreous endosperm formation.
Although the two NILs had identical zein protein abundance (Supplementary Fig. 6b), and thus a similar number and size of PBs (Supplementary Fig. 8a, c), it is evident that PBs don’t gather around SGs (Supplementary Fig. 8c, g, k; Fig. 3e–g). A tentative explanation for the dispersal of PBs in NILA619 is the altered lipid content (Fig. 3i–k and Supplementary Fig. 9). Because zeins are a group of hydrophobic proteins, the association of PBs would involve hydrophobic interactions with lipids in the cytoplasm. In wheat, dark lipid inclusions (containing glycolipids, phospholipids, free fatty acids and monoglycerides) were found to organize in a liquid crystalline phase with protein matrix and SGs during vitreous endosperm formation. The higher level of galactolipids (DGDG and MGDG) in NILA619 endosperms is associated with persistent amyloplast membranes at 35 DAP. In NILW64A, coincident with amyloplast membrane degradation, the membrane lipids significantly decreased (Fig. 3). As a result, the residual membrane lipids might organize the FA and DAG into liposome-like structures. Condensation of liposome-like structures between neighboring PBs is evident in NILW64A, but not NILA619 (Fig. 3c, d, g, h, indicated by asterisks), supporting this hypothesis. Vitreous endosperm formation might be related to the rearrangement of PB membranes and the lipid-mediated aggregation of PBs could be a prerequisite for the adhesion of PBs. As observed in durum wheat, there might be a fusion between PB and amyloplast-derived membranes, especially during the dehydration phase, which will orientate the glass transition of the endosperm matrix towards a vitreous or an opaque solid state. Although maize endosperm contains a low level of lipids, many opaque and floury mutants exhibit defects in the structure of subcellular organelles, e.g., ER, PB, and amyloplasts, which are bounded by lipoprotein membranes. Alteration in lipid content might affect the PB-PB and SG-PB interactions (Fig. 3i, j), although further molecular and biochemical analyses are required to address this question.

Our data indicate that the composition of carotenoids in amyloplasts influence the stability of amyloplast membranes during kernel desiccation. Although there have been many studies of carotenoid interactions with membranes, little is known about the nature of carotenoids in amyloplast membranes. Ven1 is a non-functional allele of Hyd3, and results in an increased level of non-polar carotenoids in the kernel (Table 1). Since the Ven1 protein localizes to amyloplast membranes (Fig. 2d–f), there are two possibilities that could explain the abnormalities and persistence of these membranes in mature endosperm of A619. First, Ven1 could function as a structural protein that negatively regulates the membrane stability. This is unlikely, however, because mutations in upstream genes affecting carotenoid synthesis also suppress the opaque kernel phenotype in the presence of Ven1 (Fig. 4). Although these mutations affect many other secondary metabolites such as tocopherol biosynthesis, apocarotenoid derivatives and other isoprenoids, they all have the common feature that the levels of lycopene, α-carotene and β-carotene are concurrently reduced, indicating that upstream carotenoid biosynthetic genes are epistatic to the Ven1 phenotype and the altered carotenoid composition may trigger an unknown physicochemical or metabolic mechanism to protect amyloplast membranes from breaking down. Second, in contrast to polar carotenoids, e.g., lutein, zeaxanthin, and violaxanthin, that are primarily synthesized in amyloplast membranes in most inbred lines, non-polar carotenoids are in preponderance in NILA619. These carotenoids appear dispensable for the biological functions of amyloplasts, at least in terms of starch synthesis, which is supported by many high yielding commercial white maize varieties containing small amounts of carotenoids. Since the ves1 endosperm accumulated only one kind of carotenoid, i.e., phytene, and the level is obviously higher than that in the NILs and other suppressor lines (Fig. 4h), phytoene could be excluded and other nonpolar carotenoids (lycopene, α-carotene or β-carotene) considered most likely responsible for the irregular and persistent amyloplast membranes. Moreover, carotenoids are precursors to hormones and many other apocarotenoids (including many new apocarotenoids recently discovered, such as β-cyclocitrin derived from β-carotene, strigolactones, and others)18,34. The contents of these apocarotenoids may be changed in the endosperm of NILs and suppressors, which might also influence the stability of amyloplast membranes.

Since carotenoids are located in the membranes of the plastid envelop (here the non-photosynthetic amyloplast), it was logical to hypothesize a connection between the role of carotenoid composition and the physicochemical properties of amyloplast membranes. This hypothesis was strengthened by numerous works showing that, in vitro, carotenoids modulate the physical properties of model lipid membranes. For example, β-carotene tends to be randomly distributed within the hydrophobic interior of the bilayer envelope and increase the membrane fluidity, whereas polar carotenoids span lipid bilayer and have their polar groups anchored in the opposite polar zones of membranes. As a result, they increase the viscosity of the membrane35–37. In addition, zeaxanthin, but not β-carotene, increases the rigidity of model phospholipid membranes35,36. This effect of polar carotenes is similar to the effect of cholesterol, i.e., limits the molecular motion of the lipid alkyl chains and favors the extended conformation of the alkyl chains of bilayer membranes. Probably, the elevated nonpolar carotenoids, particularly β-carotene, and decreased polar carotenoids, particularly zeaxanthin, created by Ven1 could increase the fluidity and hydrophobicity of amyloplast membranes and result in the irregularly expanded amyloplast membranes (Fig. 3e, f). The altered membrane physical properties could delay breakdown of amyloplast membranes in NILA619 endosperm, although the mechanism is unclear. As a result, NILA619 endosperm cells contain more amyloplast-specific lipids, DGDG, and phospholipids with a higher level of unsaturation than NILW64A (Fig. 3j–k and Supplementary Fig. 9b). Scavenging singlet oxygen (1O2) in membranes by β-carotene and its polar derivative, zeaxanthin, is effective in membranes containing unsaturated lipids (such as DGDG), but not in membranes containing saturated lipids. One could envision that the higher level of unsaturated DGDG in NILA619 endosperm cells enhances the β-carotene-mediated antioxidant activity to protect the amyloplast membranes from oxidative damage and degradation during kernel dehydration. Ven1 previously identified as crtRB1 or Hyd3, has potential to be applied for high β-carotene breeding in corn10,11. Since knock-down of Ven1 expression causes an undesirable opaque kernel phenotype in many genetic backgrounds, use of the Ven1 allele or Ven1RNAi could be limited to germplasms with appropriate genetic modifiers (Supplementary Fig. 12). We created an effective screen for genetic modifiers in the natural population that can modify the opaque phenotype caused by Ven1RNAi (Fig. 5a). Indeed, several loci were identified, of which Gs1 encoding Glutathione S-transferase 1 was a major QTL (Fig. 5d), although their mechanisms remain to be understood.

Vitamin A deficiency (VAD) is the leading cause of preventable childhood blindness and is estimated to affect about one-third of the world’s children under the age of five (information from the World Health Organization, https://www.who.int/nutrition/topics/vad/en/). VAD usually results from dietary deficiency, as it mostly occurs in developing countries where the staple food is devoid of β-carotene. One economical approach to prevent VAD is to increase β-carotene levels in cereal grains. Maize, which is one of the most globally grown crops that provides nutritious food for humans, could serve as an ideal crop for provitamin A
biofortification. Although the level and composition of total carotenoids in natural maize populations varies dramatically, alleles that increase \( \beta \)-carotene in the grain are rare, suggesting that, for an unknown reason, most selected varieties tend to contain a low \( \beta \)-carotene level. Because carotenoids are more concentrated in the vitreous endosperm than in the starchy endosperm, a better understanding of vitreous endosperm formation is potentially valuable for increasing \( \beta \)-carotene in maize grain.

In any case, maize kernel vitreousness is a complex phenotype and we presented new data that offers insight into how vitreous endosperm forms. Figure 5e shows a model illustrating one possible mechanism for interfering with the vitreous phenotype through carotenoids acting on amyloplast membranes. In white corn varieties or lines with a small amount of non-polar carotenoids, degradation of amyloplast membranes occurs coordinately with interaction between SGs and PBs as endosperm is entering the dough stage, creating the foundation for vitreous endosperm formation. These interactions can be disrupted by increased accumulation of non-polar carotenoids in amyloplast membranes, which stabilizes the membranes during endosperm desiccation by changing the membrane physical properties. In combination, these biochemical changes impair the condensation of PBs and cytoplasm contents onto SGs, leading to the formation of an opaque endosperm phenotype. This phenotype could hinder utilization of superior alleles that increase \( \beta \)-carotene content. Although the nature and mechanisms of genetic modifiers need to be further investigated, their presence in natural populations broadens the germplasm available for breeding high \( \beta \)-carotene varieties of corn that would benefit children suffering from vitamin A deficiencies.

**Methods**

**Genetic materials.** Yellow dent inbred lines W64A and A619 exhibiting vitreous and opaque endosperm phenotypes, respectively, were used for Ven1 mapping. By recurrent backcrossing, the Ven1W64A allele from W64A was introduced into the A619 background for genetic analysis. At F2, the vitreous and opaque progeny seeds, segregating at a 1:1 ratio, were separated and divided into two pools for BSA sequencing. The vitreous and opaque seeds from F3, were self-pollinated for two generations, yielding vitreous and opaque NILW64A and NILA619, respectively. Similar experiments were also conducted for two other vitreous inbred lines D1051 and P25 with A619.

Two hundred sixty two temperate maize inbred lines (Supplementary Data 1) provided by Dr. Linsheng Lai at China Agricultural University, were used for the population genetic analysis.

To make the Ven1RNAi construct, reverse and forward Ven1 cDNA fragments (400 bp in length) were amplified using two primer pairs: Ven1-SP2XbaI and Ven1-SR2SacI, and Ven1-ABFpE1 and Ven1-ARBamH1. Expression of the Ven1RNAi cassette was driven by the 254D y-zsn promoter. The construct was transformed into K5585S, a vitreous inbred line. Transgenic plants were confirmed by PCR using primers described elsewhere.

**Mapping Ven1**. More than 100 vitreous and opaque kernels were selected for germination from the F2 F2BC1 ears described above. Then high-quality genomic DNA was extracted individually at the seedling stage for subsequent sequencing. Libraries were constructed using the standard protocol of the Illumina TruSeq library DNA PCR-free prep kit. Next-generation sequencing (NGS) technology was used to perform paired-end (PE) sequencing of the libraries based on the Illumina sequencing Nova Seq platform. The sequencing depth was 20x for the parents, and 50x for each mixed pool. Reads of the two bulks and two parents were filtered, quality evaluated, and sequence aligned to the 9T27 reference genome using bwa software. SNP calling was performed by GATK software and then calculated SNP index. Delta-SNP, the difference between the two bulks, was then determined and the statistical confidence intervals of index were calculated. Sequencing service was provided by Personal Biotechnology Co., Ltd. Shanghai, China.

**RNA-seq and RT-qPCR analysis**. Developing endosperm were ground in liquid nitrogen. Total RNA was extracted by TRizol reagent (Invitrogen, catalog number 1596,018) and then purified with an RNeasy Mini Kit (Qiagen, catalog number 74,106) after DNase I digestion (Qiagen, catalog number 79,234) following the manufacturer’s protocol.

RNA-seq libraries of W64A, A619, NILW64A and NILA619 were prepared with an Illumina Standard library preparation kit and sequenced by HiSeqTM 2500 at OE Biotech Co, Ltd (Shanghai, China). Quality control checks on raw sequencing data were performed by FastQC combined with the NGS QC TOOLKIT v2.3.3. Clean reads were aligned to the B73 reference genome (RefGen_v3) and the reference gene model dataset (FGS 5b) using TopHat/ Bowtie2 (cc.jhu.edu/software/tophat/). The gene expression value was calculated as fragments per kilobase of transcript per million mapped reads (FPKM). The FPKM value should be greater than zero for an expressed gene in the three biological replicates. Candidate genes and genes involved in synthesis of zeins and starch were selected and analyzed, respectively. The related web links are as follow: reference gene [ftp://ftpensemblgenomes.org/pub/plants/current/fasta/zae_mays/], reference transcript [ftp://ftpensemblgenomes.org/pub/plants/current/fasta/zae_mays/cdna/], and annotations [ftp://ftpensemblgenomes.org/pub/plants/current/fasta/zae_mays/cdna/].

For RT-qPCR analysis of each gene during seed development and filling, the digested RNA was used for reverse transcription with Super Script III First Strand Kit (Invitrogen). The resulting cDNA was diluted to 50 ng/\( \mu \)l for RT-qPCR with SYBR Green (TAKARA) on a CFX Connect Real-Time System (Bio Rad). The relative gene expression level was calculated using the comparative CT method (\( \Delta \Delta CT \) method), and the maize Actin gene was used as a control. The expression level in seeds at 8 DAP was set to 1. In addition, expression of genes involved in zein and starch synthesis were also analyzed using 24 DAP and 30 DAP endosperms. Primers were listed in Supplementary Table 2.

**Amplification of Ven1 in NILW64A and NILA619.** Genomic DNA of NILW64A and NILA619 were extracted from seedling leaves using the CTAB method and used to amplify of the full-length genomic sequence of Ven1 using primers Ven1-GF and Ven1-GR. The cDNAs reverse-transcribed from mRNAs of NILW64A and NILA619 endosperms were used for amplification of the Ven1 coding sequence (CDS) with the primers Ven1-CE+Ven1W64A-GR, and Ven1-CE+Ven1A619-G, respectively. The PCR products were then cloned into the pEASY-Blunt Zero Cloning Kit (TransGen, China), and six clones for each fragment were sequenced (TSINGKE Co., Ltd., China).
DNA sequencing. The BSA sequencing and analyses were provided by OE biotech Co., Ltd. (Shanghai, China).

Screening genetic modifiers of VenRNAI in the natural population. Two hundred sixty two inbred lines were pollinated by VenRNAI-t+. The F2 endosperm phenotypes were divided into 3 levels: vitreous (completely modified), mosaic (partially modified) and opaque (completely unmodified). According to the phenotype, GAMS analysis was performed using the vitreous and opaque groups to screen the modifiers in the natural population following the standard protocol.

Statistics and reproducibility. Three independent replications were performed to observe the vitreous endosperm formation in W64A, A616, NIL-64W4A and NIL-A619 by SEM shown in Fig. 1c and at least six kernels were analyzed in each experiment. Western blotting experiments in Fig. 1h and j were repeated three times. Confocal analyses of the HYD3 localization in leaf and endosperm cells were performed three independent times in Fig. 2c–f. TEM and SEM analyses in Figs. 3, 4 and Supplementary Fig. 8 were performed with 20 kernels from three cobs in each experiment. At least 100 endosperm cells were observed to investigate the changes in starch grain membranes shown in Figs. 3a–h, 4i and Supplementary Fig. 9a. 182 and 200 individual plants were used to perform genetic linkage analysis by PCR in two BC4FI populations, respectively, shown in Supplementary Fig. 4b and d. Three independent replications were performed to analyze the zein protein content shown in Supplementary Fig. 6a, b, g.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. Data supporting the findings of this work are available within the paper and its Source Data. A reporting summary for this Article is available as a Supplementary Information file. The genetic materials generated and analyzed during the current study are available from the corresponding author upon request. Source data are provided with this paper.

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