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Aberrant seed development in *Litchi chinensis* is associated with the impaired expression of cell wall invertase genes

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Abstract

Cell wall invertase (CWIN) are known to play important roles in seed development. However, most reports to date have focused on a single gene family member, and have mainly investigated CWIN functions during the filling stage of seed development. In this study, we found significant lower levels of CWIN protein and activity associated with seed abortion in the *Litchi chinensis* cultivar “Nuomici.” We identified five litchi CWIN genes and observed that the expression of *LcCWIN5* was limited to the flower tissues and decreased sharply with fruit development. Silencing of *LcCWIN5* expression before 28 DAA (cell division stage) resulted in perturbed liquid endosperm development, smaller seeds, and higher seed abortion rate, while silencing after 28 DAA (filling stage) had no effect on seed development. In contrast, *LcCWIN2* was mostly expressed in the funicle and seed coat, and increased with fruit development. Decreased *LcCWIN2* expression and CWIN activity during early seed filling coincided with smaller seeds in the cultivar “Feizixiao.” Silencing of *LcCWIN2* caused a reduction in the seed size without inducing seed abortion. We propose that CWIN activity in seed maternal tissues during cell division stage is likely due to *LcCWIN5* expression, which regulates early seed development. On the other hand, CWIN activity during the filling stage is due to the expression of *LcCWIN2*, which may promote carbon import by creating a sucrose gradient. Comparable *LcCWIN5* expression, but much lower CWIN activity, detected in the funicle of “Nuomici” is consistent with post-translational regulation.

Introduction

Angiosperm seeds are important sources of food and animal feed. They are composed of the two fertilization products, the embryo and the endosperm, surrounded by the maternally derived seed coat. Coordinated interaction between all three seed components is a requirement for seeds to complete their development¹.

Molecular genetic studies have shown that cell wall invertase (CWIN) proteins play a key role in determining the sequential development of the endosperm and the

embryo², and a positive correlation has been observed between CWIN activity and seed development in a range of plant species including bean (*Vicia faba*)^{3,4}, barley (*Hordeum vulgare*)⁵, maize (*Zea mays*)⁶, rice (*Oryza sativa*)⁷, tomato (*Solanum lycopersicum*)⁸, cotton (*Gossypium hirsutum*)⁹, and soybean (*Glycine max*)¹⁰. Mutants of maize and rice with reduced CWIN expression produce smaller seeds^{7,11}, whereas elevation of CWIN activity by silencing its inhibitor in tomato increases the seed size¹².

CWIN is central to phloem unloading and carbon partitioning, especially in cellular sites lacking symplastic connections such as the filial tissues of seeds⁵ and plays a critical role in photoassimilate import in the developing seeds of maize¹¹, barley⁵, bean, and tomato^{4,13}. High CWIN activities correlate with high glucose levels, generally favoring cell division and promoting cell differentiation during seed development¹⁴. Thus, CWIN has

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been associated with seed development by regulating carbon import and/or cell division of filial tissues through sugar signaling.

CWIN multigene families of varying sizes in a number of plants have been described^{15–18}. However, most such studies to date have focused on a single CWIN gene family member, and have investigated its role mainly in the filling stage of seed development. In contrast, there are fewer studies on CWIN during early seed development, when the endosperm enters a phase of substantial nuclear division immediately after fertilization, along with rapid growth of the seed integuments. Early seed development, or seed set, is a critical stage that determines the yield potential. At this stage, seed development is highly sensitive to biotic or abiotic stresses, which can lead to abnormal development or even abortion, and consequently irreversible yield loss⁹. In this regard, a growing body of literature suggests that invertases are a central component of the plant response to biotic and abiotic stimuli¹⁹. Determining how CWIN genes regulate seed development will not only advance the understanding of plant reproductive biology, but may also suggest strategies to manipulate crop yield and improve both fruit quality and tolerance to stress.

Litchi (*Litchi chinensis* Sonn.) is an evergreen fruit crop that is widely cultivated in warm regions of the world. The production of small sterile seeds is an economically desirable trait of litchi fruits, and cultivars have been bred with small seeds or stenosperry, resulting in fruits with greater commercial value. In a previous study, we demonstrated that seed abortion in a stenospermic litchi cultivar, “Nuomici,” might be due to low CWIN activity in seed pedicel and seed coat²⁰. Larger seed, longer developmental period, and easily separated seed pedicel (funicle) make litchi an excellent model for exploring the roles of CWIN in seed development. In addition, an investigation into the stenosperry mechanism in litchi has the potential to expand our knowledge of the formation of stenospermic fruit and to provide an important reference for genetic improvement programs, and help to manage fruit set.

In this current study, big-seeded, small-seeded, and seed-aborted litchi cultivars were used to investigate the role of different CWIN genes in seed development. To this end, we measured CWIN activities in the funicle, seed coat, and cotyledon of different cultivars and detected the presence of CWIN proteins using immunocolloidal gold labeling in the funicle. Five litchi CWIN genes (*LcCWIN1–5*) were identified and characterized, and the roles of two of them in seed development were evaluated by suppressing their expression through virus-induced gene silencing (VIGS). We sought to answer the following questions: (i) does CWIN regulate seed development of litchi? (ii) which CWIN gene family member is critical for

the normal development of litchi seeds? (iii) do CWIN genes affect the early or late stage of seed development?

Results

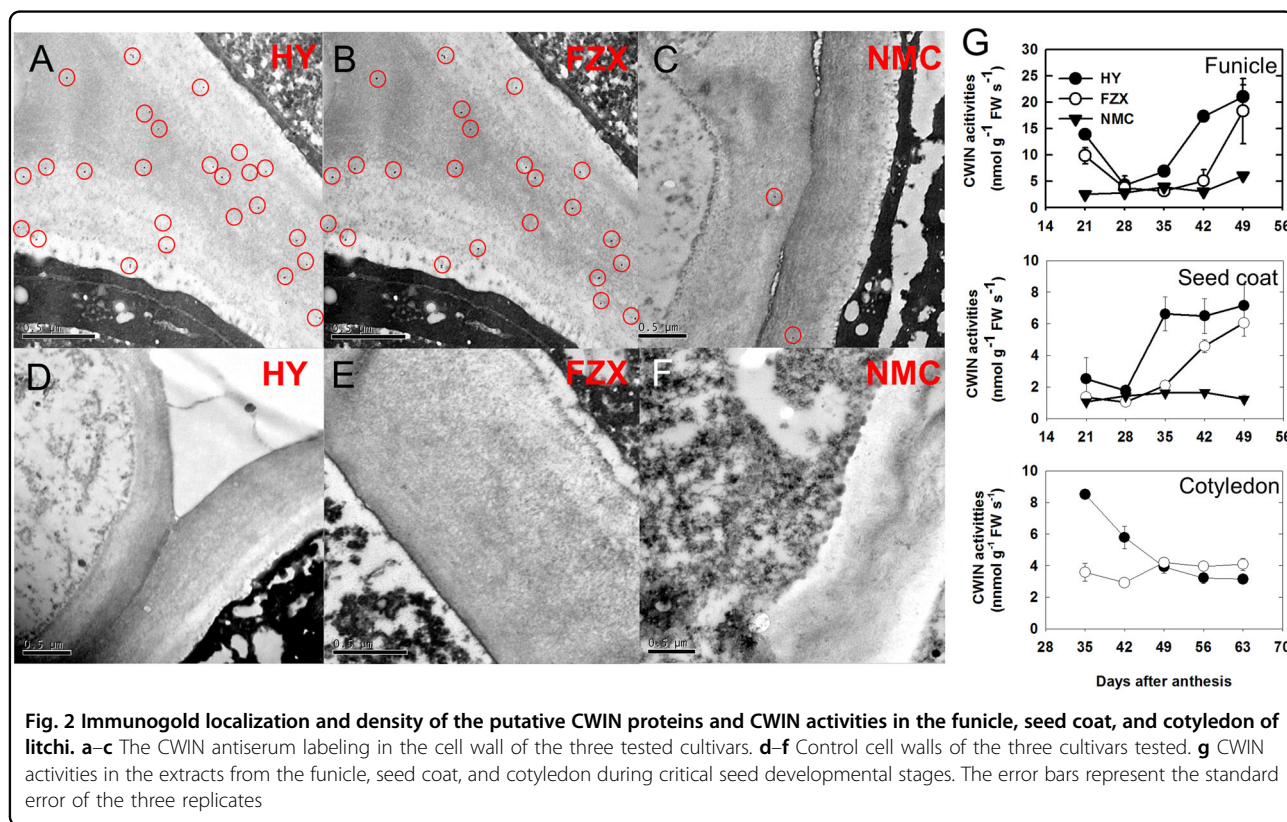
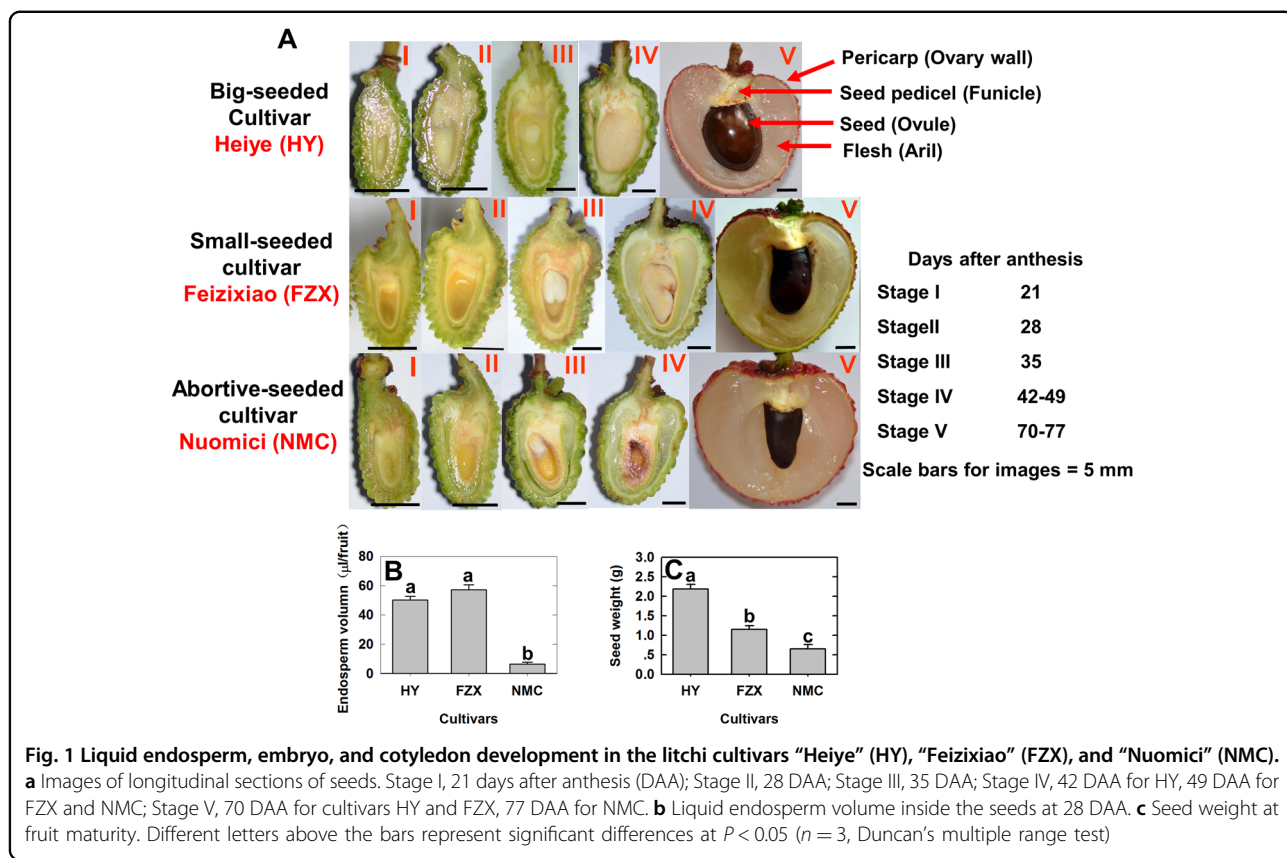
A distinct developmental transition between the cell division and filling stages, and the presence of less liquid endosperm in stenospermic “Nuomici”

According to Lü et al.²¹, double fertilization takes place in litchi 2–3 days after pollination, followed by division of the nuclei of the primary endosperm, which initially develops as a syncytium, leading to the formation of a large cell containing multiple nuclei surrounding the central vacuole (liquid endosperm). The zygote begins to divide after 3–7 days and reaches the heart stage around 30 days after anthesis, when the liquid endosperm is substantial²¹.

As shown in Fig. 1a, in both, a big-seeded cultivar, “Heiye” (HY), and a small-seeded cultivar, “Feizixiao” (FZX), visible liquid endosperm was observed around 21 days after anthesis (DAA, Stage I) and the volume increased rapidly over the next week. The liquid endosperm was most abundant at 28 DAA (Stage II) when the embryo reached the heart stage with a rudimentary cotyledon, and after 28 DAA, the liquid endosperm was absorbed by the quickly developing cotyledon. Seeds with a much larger embryo and cotyledons, but less liquid endosperm were observed at 35 DAA (Stage III). Thus, 28 DAA represents a transition point between the cell division stage and the filling stage during litchi seed development. In the seed-aborting cultivar, “Nuomici” (NMC), sequential liquid endosperm and embryo development was not observed (Fig. 1a), and the amount of liquid endosperm at 28 DAA was much lower in NMC than in HY or FZX (Fig. 1b). The three tested cultivars displayed significant differences in seed size at maturity (Fig. 1c). HY produced larger seeds with normally developed embryos and cotyledons, and FZX had smaller seeds with normally developed embryos, but smaller cotyledons, while NMC produced seeds without embryos or cotyledons.

Differences in the levels of CWIN protein and activity

We used a CWIN antiserum coupled with immunogold labeling to investigate the presence CWIN proteins in the funicle at the early stage of fruit development (7 DAA). In contrast to the scarcely found immunogold particles in a control without antiserum, the proteins reacting with anti-CWIN serum were clearly observed in the cell walls of all the three cultivars tested (Fig. 2a–f). The density of immunogold particles was significant lower in the cell walls of NMC than in those of FZX and HY. In addition, we measured the CWIN activities in protein extracts from the funicle, seed coat, and cotyledon, and detected significantly lower activity in NMC than in HY and FZX at



most sampled points (Fig. 2g). These results were consistent with the lower fructose and glucose contents observed in the funicles of NMC, compared to those of FZX and HY²². In HY and FZX, the funicle CWIN activities were high at 21 DAA, followed by a short decrease and an increase again at 28 DAA and 42 DAA, respectively. From 35 to 42 DAA, the CWIN activities in the funicle, seed coat, and cotyledon of FZX were much lower than those in HY (Fig. 2g).

Identification and characterization of the Litchi CWIN gene family

In order to identify litchi CWIN genes, sequences annotated as putative glycoside hydrolases were obtained by searching the *L. chinensis* genome database (<http://litchidb.genomics.cn/page/species/index.jsp>) and CWIN-specific regions including conserved functional motifs and a signal peptide were analyzed. Five putative CWIN genes, named *LcCWIN1-5*, were identified, and full-length cDNAs of each were cloned by RT-PCR using gene-specific primers. The cDNA and deduced amino acid sequences were deposited in the GenBank with the following accession numbers: *LcCWIN1* (KX981206), *LcCWIN2* (KX981207), *LcCWIN3* (KX981208), *LcCWIN4* (KX981209), and *LcCWIN5* (KX9812010).

The predicted open reading frames (ORFs) of the five genes were between 1677 and 1740 bp, corresponding to deduced amino acid sequences ranging from 558 to 579 (Table 1). The most homologous sequences in the GenBank, identified using BLAST (<https://blast.ncbi.nlm.nih.gov>), ranged from 69 to 79% identity (Table 1). The sequences of *LcCWIN2* and *LcCWIN5* ORFs were identical, while other *LcCWINs* displayed 2–7 SNPs among the three cultivars tested. With the exception of a three amino acid deletion in the NDPNG motif (β -fructosidase motif) in *LcCWIN1*, the three conserved sequence domains that represent the predicted active sites (i.e., NDPNG, RDP, and WEC[P]VD) of the five *LcCWIN* proteins were intact (Fig. 3). To investigate the phylogenetic relationship of the *LcCWIN* proteins with homologs from different plant species, an unrooted phylogenetic tree was constructed

using the neighbor-joining method (Fig. S1). The sequences were classified into five major groups: monocot-I, monocot-II, dicot-I, dicot-II, and CWINs with an acidic pI from both dicots and monocots. *LcCWIN1* belonged to the subgroup with an acidic pI, together with *Arabidopsis thaliana* AtCWIN6, and maize ZmCWIN4. CWIN proteins characterized by a high pI are thought to interact with the cell wall, while invertases with an acidic pI are thought to be soluble and localized intracellularly^{12,23,24}. Another phylogenetic tree was constructed without CWIN proteins with acidic pI (Fig. 4), and *LcCWIN2* was found to belong to the dicot-I group, along with *A. thaliana* AtCWIN2/4, as well as sequences from tomato (Lin5-8) and potato (StinvGE, StinvGF, StpCD111 and StpCD141). *LcCWIN3-5* and *A. thaliana* AtCWIN1/5 grouped with other dicot-II proteins, with *LcCWIN3/5* being most closely related to the CWIN from grapevine.

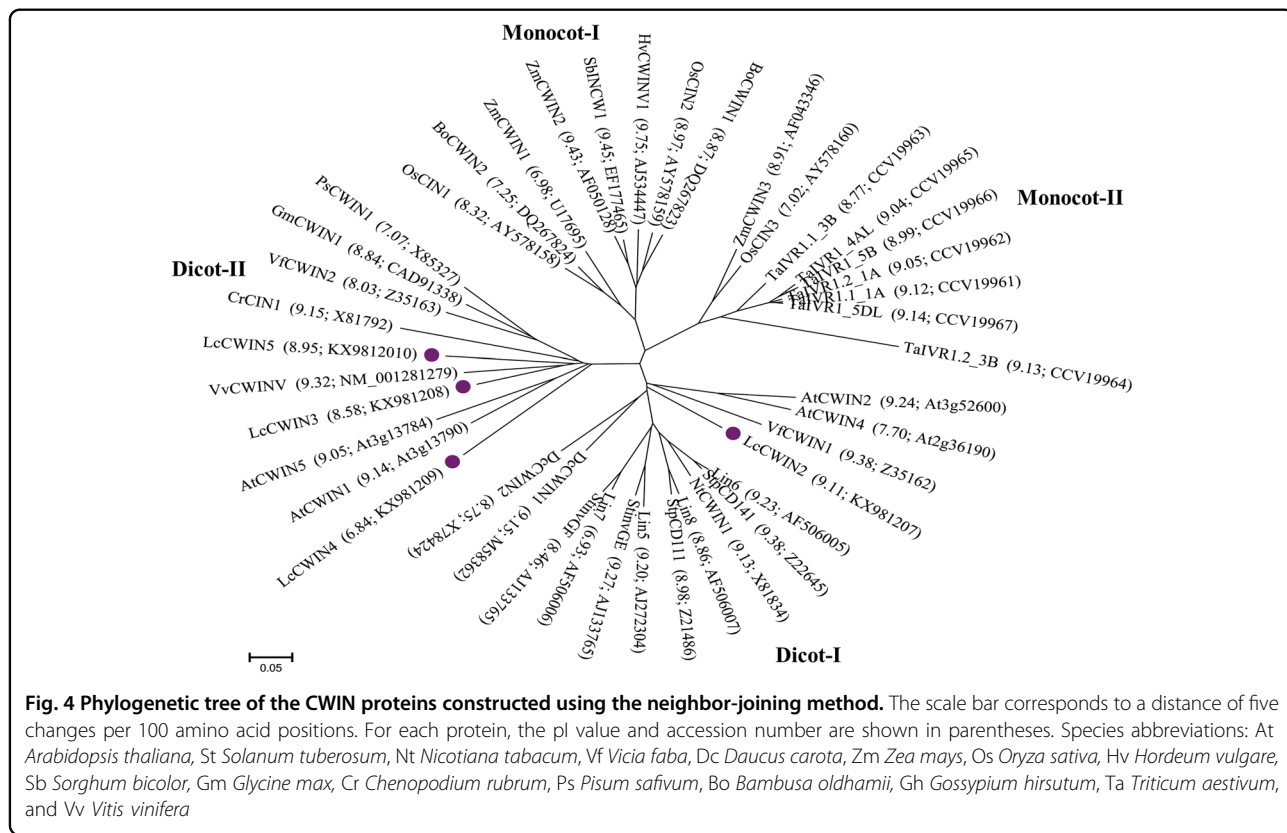
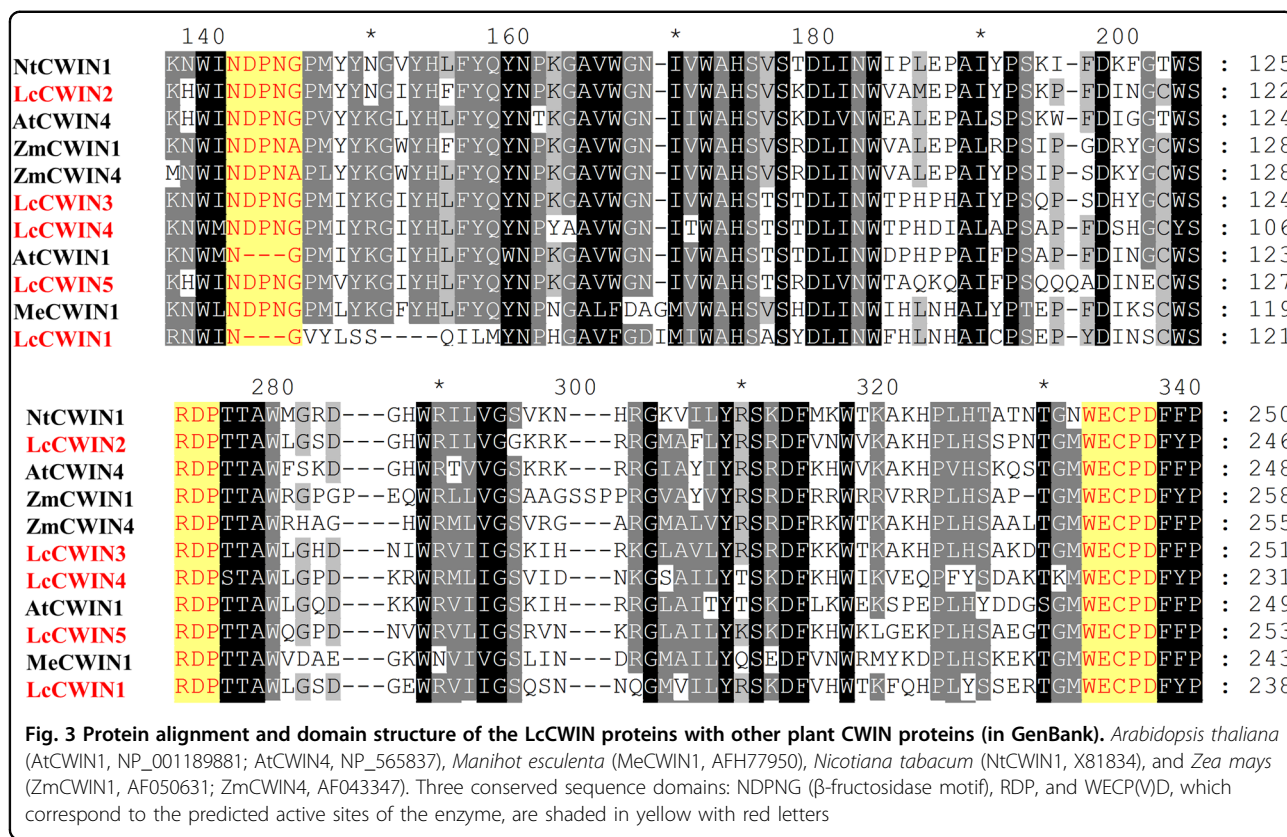
LcCWIN expression patterns and CWIN activities

The transcript levels of the *LcCWIN* genes and the CWIN activities in different litchi tissues were measured (Fig. 5), and we observed notable differences. *LcCWIN1-4* were expressed in all tissues, while *LcCWIN5* was specifically expressed in anthers and pistils. Among the four constitutively expressed *LcCWIN* genes, we observed that *LcCWIN2* was predominantly expressed in the funicle and the seed coat, and so we focused and targeted this gene in subsequent studies of the role of CWIN genes in seed development. *LcCWIN1* also showed relatively high expression levels in the funicle and seed coat, but the predicted protein had a three amino acid deletion in the β -fructosidase motif. The expression pattern of *LcCWIN5* was generally consistent with that of LIN5, a tomato CWIN gene that was shown to be required for seed development⁸. Thus, *LcCWIN2* and *LcCWIN5* were selected for further studies. Consistent with the predominant *LcCWIN5* expressions, anther and pistil displayed the highest CWIN activities, followed by the funicle and the seed coat (Fig. 5).

The developmental patterns of *LcCWIN1*, *LcCWIN2*, and *LcCWIN5* expression in the funicle of the cultivars tested are shown in Fig. 6a–c. *LcCWIN1* was preferentially

Table 1 Sequence homologies based on amino acid sequences of litchi CWIN isolated from litchi cv. “Feizixiao,” and single-nucleotide polymorphism (SNP) numbers detected in *LcCWINs*, among the three cultivars tested

Gene	GenBank number	ORF(bp)	Length(AA)	Top BLAST match (GenBank number)	Homology(%)	SNP numbers
<i>LcCWIN1</i>	KX981206	1731	576	<i>Theobroma cacao</i> (XM_018128260.1)	69	2
<i>LcCWIN2</i>	KX981207	1722	573	<i>Citrus sinensis</i> (XM_006481682.1)	75	0
<i>LcCWIN3</i>	KX981208	1734	577	<i>Citrus sinensis</i> (XM_006426486.1)	79	7
<i>LcCWIN4</i>	KX981209	1677	558	<i>Dimocarpus longan</i> (KP769771.1)	72	2
<i>LcCWIN5</i>	KX981210	1740	579	<i>Theobroma cacao</i> (XM_007024428.2)	69	0



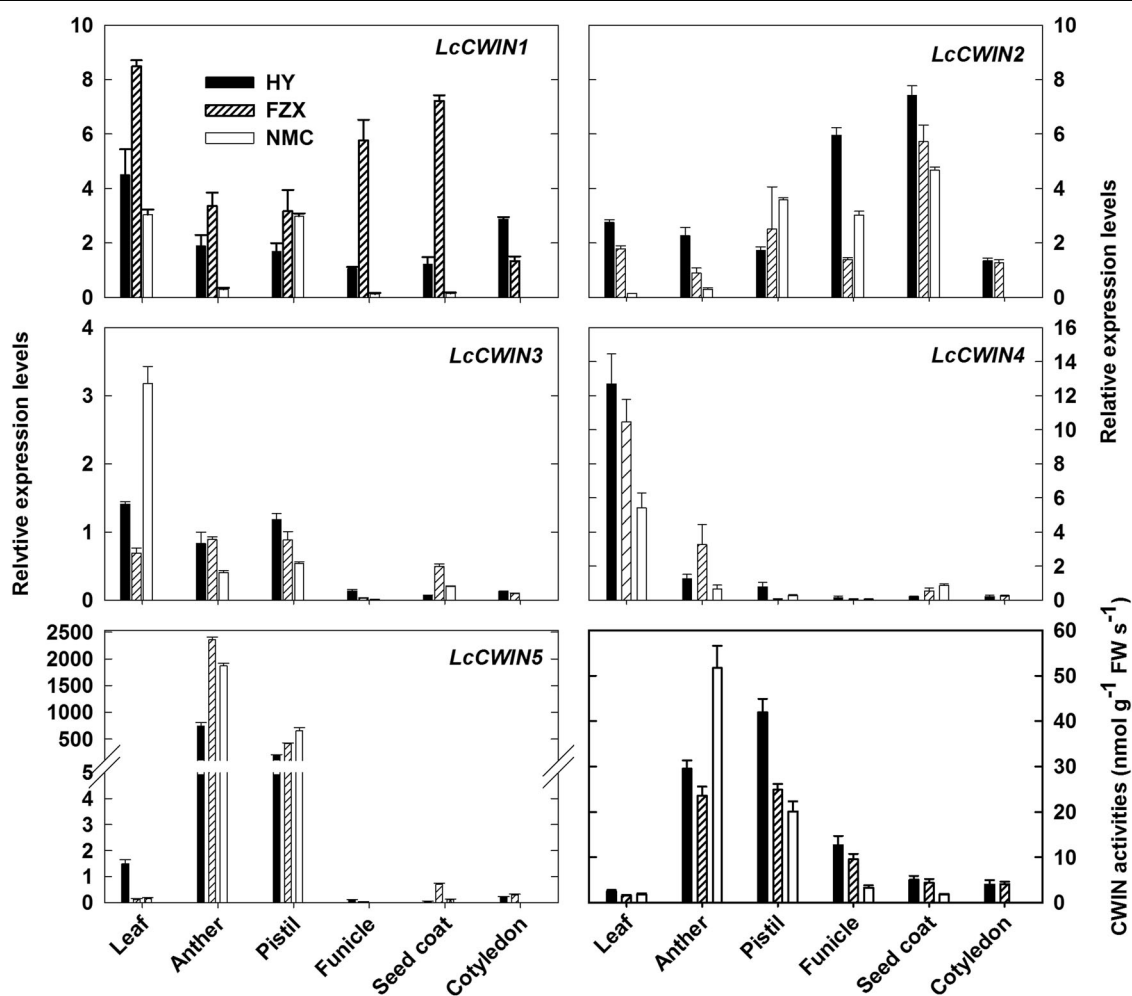


Fig. 5 The expression patterns of the five litchi CWIN genes and CWIN activities in different organs/tissues. New mature leaves were taken from the first autumn shoots. Anther and pistil were taken on the day of full bloom, while funicle, seed coat, and cotyledon were sampled from fruits at 49 days after anthesis. The error bars represent the standard error of three replicates

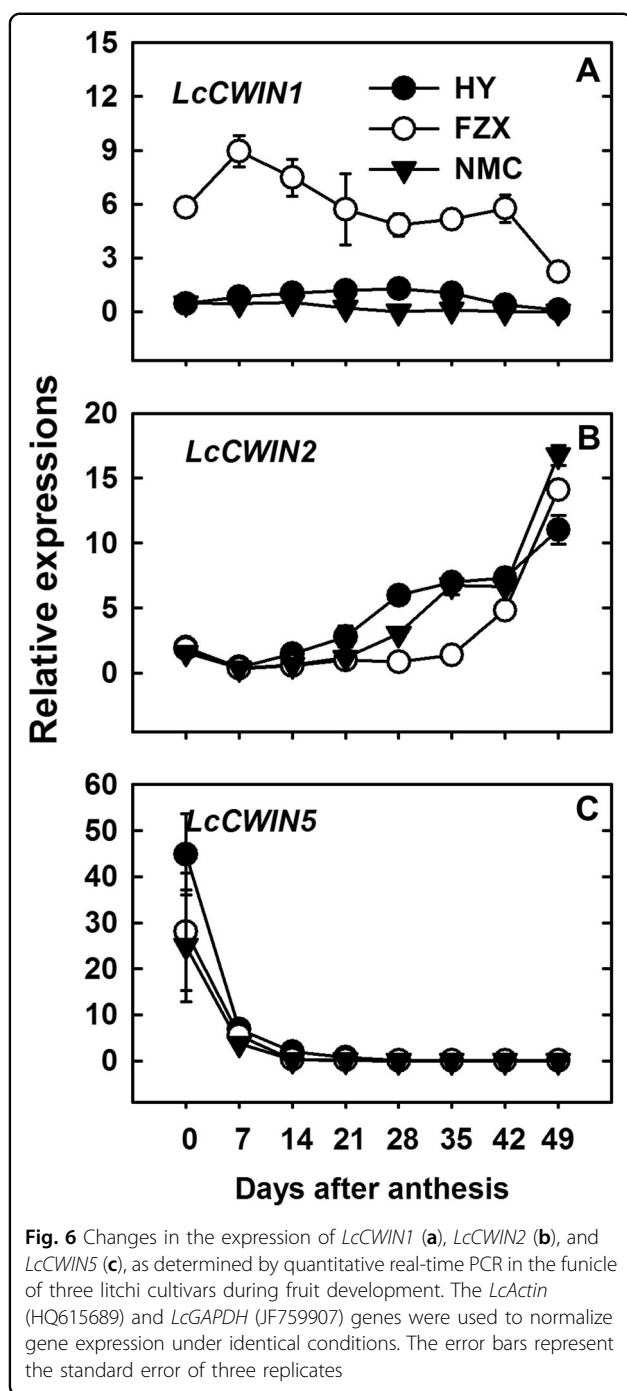
expressed in FZX throughout fruit development. The expression of *LcCWIN2* remained low before 14 DAA in all the three cultivars tested. Increased expression occurred later, but at different time points in different cultivars. A noticeable increase in *LcCWIN2* expression was evident at 21 DAA in HY, at 28 DAA in NMC, and at 42 DAA in FZX. In the funicle of all the cultivars, the expression of *LcCWIN2* increased, while that of *LcCWIN5* decreased with seed development. *LcCWIN5* expression peaked at the day of blooming, followed by a sharp decrease with seed development.

In vivo functional analysis of *LcCWIN2* and *LcCWIN5* using VIGS in normal seed cultivars

VIGS is a powerful and rapid tool that enables targeted gene loss-of-function studies, and has been successfully applied to induce the silencing of phytoene desaturase (PDS, EC 1.3.99.30) gene in woody litchi leaves and of

LcUFGT1 in the pericarp²⁵. In the present study, to test the function of *LcCWIN2* and *LcCWIN5* in litchi seed development, the vectors pTRV2-*LcCWIN2* and pTRV2-*LcCWIN5* were generated by inserting fragments of *LcCWIN2* or *LcCWIN5* into the pTRV2 vector. An *Agrobacterium tumefaciens* suspension containing the pTRV2-*LcCWIN2* or pTRV2-*LcCWIN5* constructs was injected into the fruit stalks still attached to the plant at 21–35 DAA, or wetted the panicles on the day of female blooming.

First, we explored the effects of fruit-stalk injection with pTRV2-*LcCWIN5* at 21 DAA on seed development of normal seed litchi cultivars, HY and 9911 (Fig. 7). Ten days after the treatment, the pTRV2-*LcCWIN5* infiltration resulted in a significantly lower fruit retention rate and liquid endosperm volume (Fig. 7a). At 3 weeks (cultivar HY) or 5 weeks (cultivar 9911) after the injection, significantly lower fruit and seed weight were observed in



fruit that had been infiltrated with pTRV2-*LcCWIN5*. Longitudinal sections confirmed a higher incidence of embryo and cotyledon development failure in fruits treated with pTRV2-*LcCWIN5* than those treated with pTRV2-Empty control vector (Fig. 7b, c).

We also silenced the expression of *LcCWIN2* and *LcCWIN5* through panicle dipping and examined the effects on seed development in the cultivars “Heiye” (HY) and “Baitangying.” To confirm the suppression effects, we

performed quantitative RT-PCR and measured CWIN enzyme activities in different fruit tissues of HY at 4 weeks after panicle dipping (Fig. S2). The primers that annealed to a region outside the *LcCWIN2* and *LcCWIN5* sequences were used for silencing. We observed that in the pTRV2-*LcCWIN5*-infiltrated fruits, the *LcCWIN5* transcript levels were significantly reduced in the funicle, when compared with the pTRV2-Empty control (Fig. S2A). The ineffective silencing in the other tissues might reflect the extremely low expression of the *LcCWIN5* gene in those tissues. In contrast, *LcCWIN2* expression was reduced in the funicle, seed coat, and cotyledon of the pTRV2-*LcCWIN2*-infiltrated fruit, compared with the fruit infiltrated with the control vector (Fig. S2B). Both the silencing of *LcCWIN5* and *LcCWIN2* resulted in decreased CWIN activities in the funicle, seed coat, and cotyledon, although the decrease was considerably greater in the case of *LcCWIN5* silencing (Fig. S2C–E). In both cultivars, the reduction in *LcCWIN2* transcript levels did not affect the fruit retention and seed abortion rate, but significantly reduced the seed weight, while the silencing of *LcCWIN5* led to significantly lower fruit retention, liquid endosperm volume, and seed weight, but a higher seed abortion rate (Table 2).

When *LcCWIN2* and *LcCWIN5* were silenced through fruit-stalk injection at different stages of seed development in two big-seeded cultivars, “Heiye” and “Shuilin,” we observed different effects on the seed weight depending on the silenced gene and the developmental stage (Fig. 8). The fruits infiltrated with a *LcCWIN5* antisense construct at 21 and 28 DAA produced significantly smaller seeds, while the silencing at 35 DAA did not significantly affect the seed weights in both cultivars. On the other hand, the silencing of *LcCWIN2* at 21, 28, or 35 DAA decreased the seed weight in both cultivars, although it was less apparent as comparing to the silencing of *LcCWIN5*.

Discussion

CWIN regulates early seed development

Seed development depends on the coordinated development of the seed coat, endosperm, and embryo. In the stenosperric litchi cultivar NMC, a normal seed coat develops, but visible liquid endosperm development does not occur and zygote development is retarded (Fig. 1). In parallel with the failure of sequential liquid endosperm and embryo development, we saw a significant lower density of CWIN-immunogold particles and reduced CWIN activities at the site of seed assimilate unloading, compared with the big-seeded cultivar, HY, and the small-seeded cultivar, FZX (Fig. 2). Furthermore, silencing of *LcCWIN5* resulted in decreased CWIN activity, a significantly lower liquid endosperm volume, and a higher seed abortion rate in the big-seeded cultivars (Table 2 and

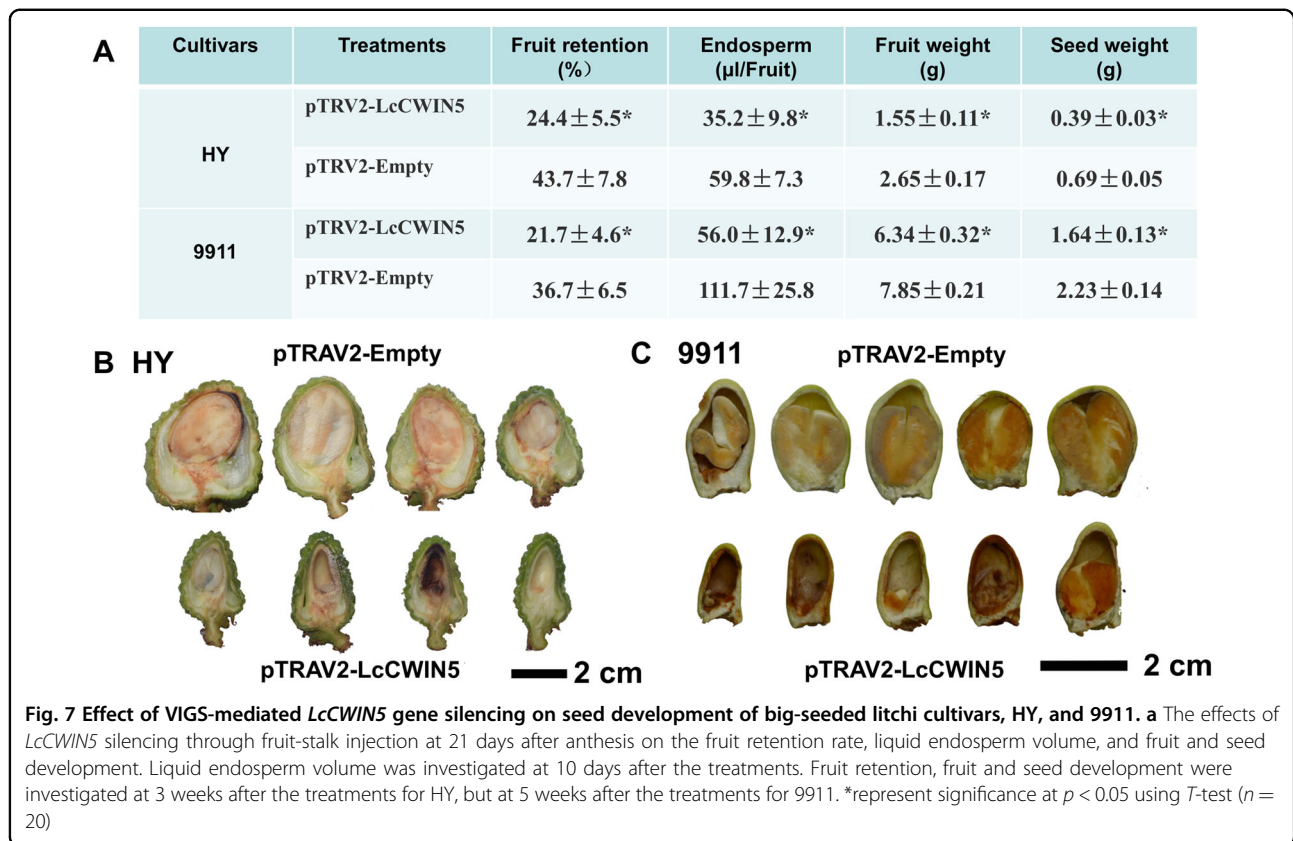


Table 2 The effects of *LcCWIN2* and *LcCWIN5* silencing, through panicle dipping in a bacterial suspension at female blooming, on the fruit retention rate and seed development of the litchi cultivars “Heiye” and “Baitangying”

Cultivar	Treatments	Fruit retention(%)	Liquid endosperm($\mu\text{l}/\text{seed}$)	Seed weight(g)	Abortion rate(%)
Heiye	pTRV2-LcCWIN2	23.7 \pm 4.5b	50.2 \pm 2.5a	1.37 \pm 0.11b	19.2 \pm 5.1b
	pTRV2-LcCWIN5	7.5 \pm 1.4a	38.6 \pm 5.5b	0.73 \pm 0.10c	50.3 \pm 11.2a
	pTRV2-Empty	25.6 \pm 6.6b	54.6 \pm 5.3a	1.87 \pm 0.14a	12.4 \pm 3.5b
Baitangying	pTRV2-LcCWIN2	38.2 \pm 3.8a	13.3 \pm 0.76a	0.75 \pm 0.02b	51.3 \pm 2.3b
	pTRV2-LcCWIN5	33.7 \pm 4.2a	9.0 \pm 0.61b	0.70 \pm 0.04b	69.4 \pm 4.8a
	pTRV2-Empty	38.8 \pm 4.4a	15.2 \pm 0.68a	0.86 \pm 0.08a	50.0 \pm 5.0b

Liquid endosperm volume was investigated at ten days after treatments. Fruit retention was investigated at three weeks after treatments. Seed weight and abortion rate were investigated at maturity. Different letters after the values in the same cultivars indicate significant differences at $P < 0.05$ among treatments, according to Duncan's Multiple New Range Test ($n = 20$).

Fig. 7). *LcCWIN5* expression in the funicle, but not in the cotyledon, was significantly reduced, paralleling with the perturbed seed development in response to *LcCWIN5* silencing (Fig. S2). These results suggest a key role for maternal CWIN in litchi seed development.

There is considerable evidence of CWIN playing a role in seed development^{3–10}. However, these studies mainly focused on the filling stage, and the underlying mechanism of CWIN in regulating seed development was unclear. In the present study, litchi seed development could be clearly divided into the cell division stage and the filling

stage at around 28 DAA, when the zygote reaches the heart stage (Fig. 1). There is compelling evidence that CWIN activity correlates with mitotic activity, cell number, and seed size. In large-seeded genotypes of fava bean (*Vicia faba*), CWIN is active for longer time in the enlarged seed coat and, consequently, more cells are produced in the embryo⁵. Additionally, in maize, a CWIN deficiency mutation (*mn1*) impairs endosperm development by reducing the cell number^{6,26}. It is also noteworthy that the effect of CWIN on the rate of cell division most likely results from higher hexose levels, rather than

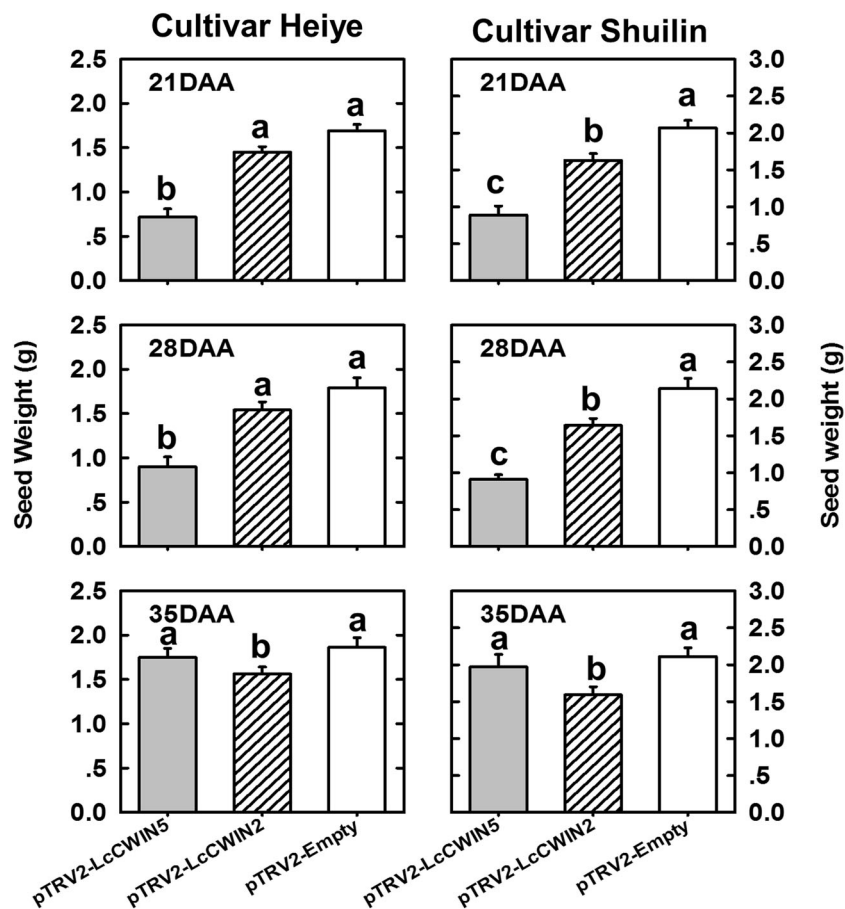


Fig. 8 Effects of VIGS-mediated *LcCWIN5* and *LcCWIN2* gene silencing at three different fruit developmental stages on the seed size of the big-seeded litchi cultivars, "Heiye" and "Shuilin." Injection was given at 21, 28, and 35 days after anthesis, and the seeds were harvested at fruit maturity (77 days after anthesis). Different letters above the bars represent significant differences at $P < 0.05$ ($n = 20$, Duncan's multiple range test)

from the amount of carbon import⁶. The filial tissue of a litchi seed accumulates about 90% dry mass during the filling stage (after 35 DAA), yet the silencing of *LcCWINs* at 35 DAA does not increase the seed abortion rate (Fig. 7). These results indicate that *LcCWIN5* functions in early litchi seed development when the endosperm and the embryo undergo active cell division.

In angiosperms, double fertilization leads to the formation of an embryo and an endosperm. The cellularized endosperm acts as nourishing tissue that is consumed by the embryo during maturation in dicotyledons¹. If endosperm development fails, it will ultimately cause the embryo to arrest its development. Both endosperm and embryo are known to display high mitotic activity during early seed development¹⁴. We hypothesize that the impaired endosperm that resulted from decreased *CWIN* expression might be a consequence of nuclear division proceeding earlier in the endosperm than in the embryo (Figs. 1 and 7; Table 2). Our finding that the silencing of *LcCWIN5* at 28 DAA, when there is an abundance of

liquid endosperm, resulted in failed embryo development (Fig. 8) indicates that the developing embryo can be directly controlled by *CWIN*.

Numerous studies have shown that sugar acts as a signaling molecule^{27–29}. For example, in *A. thaliana*, sugar provides an endogenous cue for the juvenile-to-adult phase transition³⁰. Sugar signals are perceived and transduced through a glucose sensor, HEXOKINASE1 (HXK1), which exerts its regulatory function through transcriptional activation, translational inhibition, mRNA decay, and protein degradation²⁷. The *CWIN*-elevated transgenic tomato plants exhibited higher activities of hexokinase and fructokinase in ovaries and enhances fruit set through suppressing programmed cell death in the placenta under heat stress³¹. Litchi seed abortion under reduced *CWIN* might also be associated with the programmed cell death of the filial tissues. The underlying networks in early seed development of litchi in response to *CWIN* availability will provide a subject for future investigation.

Different roles for the *LcCWIN* genes in seed development

In the present study, we identified and characterized five litchi *CWIN* genes. Except for *LcCWIN1*, the deduced amino sequences of the other four *LcCWIN* proteins all contain the highly conserved β -fructosidase motif and cysteine catalytic sites (Fig. 3). A phylogenetic tree of *CWIN* sequences from other plants and these four putative litchi *CWIN* proteins showed four clusters (monocot-I, monocot-II, dicot-I, and dicot-II), which appeared to be associated taxonomic relationship of the associated plants (Fig. 4). The *LcCWIN2* gene belonged to dicot-I, while *LcCWIN3-5* grouped with other dicot-II sequences, suggesting functional divergence. *LcCWIN2* was predominantly expressed in the funicle and seed coat, the site of seed assimilate unloading. *LcCWIN5* was specifically expressed in flowers (anther and pistil) and in young fruit (Figs. 5 and 6), which is consistent with of the expression pattern of *LIN5*, a cell wall invertase critical for seed development and fruit set in tomato⁸.

We adopted a VIGS approach to silence the expression of *LcCWIN2/5* genes. As mentioned earlier, the silencing of *LcCWIN5* before 28 DAA significantly inhibited seed development and increased the rate of seed abortion of normal seed cultivars, whereas silencing of *LcCWIN5* after 28DAA did not affect seed development (Table 2, Figs. 7 and 8). These results suggest that *LcCWIN5* regulates litchi seed development at the early stage, when filial tissues undergo active cell division. The filial tissues (embryo and endosperm) are symplastically isolated from the maternal seed coat, and so sugars are secreted from the maternal layers and imported into the filial tissues. In maize seeds, an in situ fluorometric assay of glucose in tissue sections suggested that glucose might enter the filial tissues during the very early stages of seed development, 0–2 days after pollination³². *LcCWIN5* was expressed in very young fruits (Fig. 6a). In litchi, *CWIN* activity in the critical sugar transportation site (funicle and seed coat) during the early stage of seed development is likely due to *LcCWIN5* expression, which helps the entrance of hexose, a prerequisite for normal seed development.

We observed high expression of *LcCWIN1*, which has deletions in an important functional motif, but low expression of *LcCWIN2* in tissues of the FZX cultivar (Fig. 6). The RT-PCR analysis showed that *LcCWIN2* transcript levels in the funicle of FZX were much lower than in NMC and HY at 28–42 DAA, when early seed filling occurred (Figs. 1 and 6). This was consistent with the low *CWIN* activity in the funicle of FZX during this period (Fig. 2g). The *CWIN* activities in the seed coat and cotyledon of FZX during this period were also much lower than those in cultivar HY (Fig. 2g). In rice, *GIF1* (GRAIN INCOMPLETE FILLING 1, also known as *OscIN2*) is a cell wall invertase required for carbon partitioning during early grain-filling⁷. The *gif1* mutant is

morphologically normal, with normal seed setting, but reduced grain weight and loosely packed starch granules⁷. Similarly, the *LcCWIN2* reduced cultivar FZX displayed normal liquid endosperm development, but reduced seed size (Figs. 1 and 6). And in addition, the silencing of *LcCWIN2* reduced the seed size without increased abortion rate (Table 2, Fig. 8). Taken together, these results suggest that low expression of *LcCWIN2*, and thus low *CWIN* activity, during the critical seed filling stage might be associated with the small seed size of the FZX cultivar.

In the present study, the *CWIN* activities in the funicle, seed coat, and cotyledon of small-seeded FZX were all significantly lower than the big-seeded HY (Fig. 2g). In paralleled with the reduced seed size in response to the silencing of *LcCWIN2*, the *CWIN* activity in the cotyledon, but not in the funicle or seed coat, was significantly downregulated (Fig. S2). These results suggested that the *CWIN* in the filial tissue might play an important role in the seed development of litchi. This is consistent with the result of maize. A loss-of-function mutation at the *Mn1* locus results in only 1% *CWIN* activity in the endosperm and a loss of >70% weight at maturity but non-lethal seed⁶.

In summary, we propose that the litchi *CWIN* genes have different roles in seed development. The maternal early-expressed *LcCWIN5* is involved in the early liquid endosperm and embryo development, whereas the late-expressed *LcCWIN2* is associated with seed filling.

Interestingly, the expression patterns of *LcCWIN5* among cultivars during the early stage of fruit development did not correlate with *CWIN* activity. Much lower *CWIN* activity, but comparable *LcCWIN5* expression, was detected in the funicle of the NMC cultivar (Figs. 2 and 6). This suggests that post-translational regulation may be responsible for the decreased *CWIN* activity and abnormal early seed development in NMC. This is in agreement with reports of post-translational regulation of invertase activity by inhibitory proteins in tobacco³³, maize³⁴, and tomato¹². Future studies will focus on a cell wall invertase inhibitor gene (*LcCIF*, KX981211) that we have identified, which is predominantly expressed in NMC.

Materials and methods

Plant materials

The experiments were conducted at the experimental orchard of the South China Agricultural University, Guangzhou, China. The litchi trees used for the experiments received standard horticultural practices and were open pollinated. Mature leaves, male and female flowers, and fruits from three trees were collected from big-seeded cultivar “Heiye,” small-seeded cultivar “Feizixiao,” and seed-aborted cultivar “Nuomici.” The funicle, seed coat, and cotyledon tissues of different litchi cultivars were harvested to measure the *CWIN* activities and the

expression of *LcCWIN* genes from anthesis until 49 DAA when cotyledons filled the seed coat.

For VIGS assay, the cultivars were chosen on the basis of their seed types. In 2015, big-seeded cultivars, “Heiye” and “9911,” was used to investigate the effects of *LcCWIN5* on seed development through fruit-stalk injection at 21 DAA. In 2016, the big-seeded cultivar, “Heiye,” and the partly aborting cultivar, “Baitangying,” were used to evaluate the effects of *LcCWIN2* and *LcCWIN5* silencing through panicle dipping at female blooming. Furthermore, the big-seeded cultivars, “Heiye” and “Shuilin,” were used to investigate the effects of *LcCWIN2* and *LcCWIN5* silencing on seed development through fruit-stalk injection at 21, 28, and 35 DAA.

Observation of seed development and the measurement of liquid endosperm volume and seed weight

Fruit at different developmental stages were cut longitudinally into two parts and photographed. A syringe (100 μ L) was used to remove and quantify the volume of liquid endosperm after cutting the tip of the seed. More than 50 fruits from each sampling tree were used for the liquid endosperm volume measurement. The seed weights of cultivars “Heiye,” “Feizixiao,” and “Nuomici” were measured at fruit maturity using 1/1000 electronic balance.

CWIN activity assays and immunocolloidal gold labeling

Proteins were extracted and CWIN activity was assayed, as previously reported³⁵. For the immunogold labeling assay, litchi funicles were cut into 2–3 mm³ cubes that were immediately fixed with 4% (w/v) paraformaldehyde and 2.5% (v/v) glutaraldehyde in 100 mM precooled PBS (8 mM Na₂HPO₄, 1.5 mM KH₂PO₄, 3 mM KCl, and 500 mM NaCl, pH 7.2) for 4 h. Dehydration and subsequent infiltration was conducted according to Zhang et al³⁶. Ultrathin sections (~50 nm) were cut and mounted on 100-mesh nickel grids coated with 0.3% Formvar films for subsequent immunolabeling. The ultrathin sections were first blocked by floating the grids on droplets of PBS (pH 7.4) supplemented with 50 mM glycine for 30 min at room temperature and continuously blocked with PBS (pH 7.4) supplemented with 0.1% (w/v) gelatin, 0.5% (w/v) bovine serum albumin (BSA), and 0.1% (v/v) Tween 20. Antiserum against CWIN was custom made in Bioleaf Biotechnology Co., Ltd (Shanghai, China). Goat anti-rabbit IgG antibody conjugated with 10 nm gold was purchased from Sigma.

Isolation and cloning of CWIN genes

Fourteen sequences annotated as putative invertase were obtained from the *L. chinensis* genome database (<http://litchidb.genomics.cn/page/species/index.jsp>). Among them, five sequences showed high similarity to the

reported CWIN genes from other plant species and were predicted to encode proteins with conserved CWIN functional motifs. These were targeted as putative litchi CWIN genes and were named *LcCWIN1* to *LcCWIN5*. Their coding sequences were amplified from cDNA libraries using gene-specific primers (Table S1). Full-length *LcCWIN1*, *LcCWIN3*, and *LcCWIN4* fragments were obtained from a cDNA library that was reversed transcribed from litchi leaf mRNA, while full-length *LcCWIN2* and *LcCWIN5* were obtained from cDNA libraries that were reversed transcribed from funicle and flower mRNA extracts.

Quantitative real-time PCR (qRT-PCR) analysis

Total RNA from different litchi organs/tissues was extracted using the RNA_{OUT} kit (Tiandz, Beijing, China), and cDNA was synthesized from total RNA (2 μ g) using oligo dT primers and M-MLV reverse transcriptase, according to the manufacturer’s instructions (Invitrogen, USA), in a total volume of 20 μ L. *LcCWIN* transcript levels were measured using qRT-PCR, as previously described³⁷. The specific real-time PCR primers are listed in Table S2. We analyzed the expression in biological triplicate samples. Real-time PCR reactions were normalized to Ct values for litchi *LcActin* (HQ615689) and *LcGAPDH* (JF759907). The relative expression levels of the target genes were calculated using the $2^{-\Delta\Delta Ct}$ method³⁸.

VIGS-mediated *LcCWIN* silencing

VIGS was carried out as previously described, with modifications in treatments and bacterial concentration²⁵. Air temperature and humidity is critical for high infection rate, and therefore successful for VIGS assays. The orchard air temperature is typically 22–25 °C and the relative humidity is usually > 70% during litchi flowering and young fruit development, which is optimal for VIGS assays in the orchard. TRV1 and TRV2 vectors containing *LcCWIN2* (400 bp) or *LcCWIN5* (446 bp) fragments were transformed separately into *Agrobacterium tumefaciens* strain GV3101 and then grown separately overnight in LB liquid media containing 50 mg L⁻¹ kanamycin and 1 mg L⁻¹ rifampicin. The rest of the protocol followed our previous report²⁵. After optimization, two ways of treatment, panicle dipping and fruit-stalk injection, were employed for silencing of the genes, with a bacterial culture concentration of OD₆₀₀ \approx 1.0. For the dipping treatments, panicles were dipped into the bacterial suspension for about 30 s at female flower blooming. For the injection treatments, a 1 mL syringe was used to inject bacterial suspension into the fruit stalk. Thirty inflorescence or clusters existing in different parts of the canopy of each tree were treated. Among them, 20 clusters were used to investigate the fruit retention, seed weight, and abortion

rate, and the rest ten clusters were used to take samples to measure liquid endosperm volumes and silencing effects. The number of fruitlets was counted both at 2 weeks after panicle dipping or at the day of injection and 49–56 DAA, and the fruit retention rate was calculated. The liquid endosperm volume was investigated at around 28 DAA or 10 days after injection. The seed weight and abortion rate were investigated around 49–77 DAA, when seed filling had finished or at fruit maturity. Samples were collected ~5 weeks after dipping to test the silencing effects. Three fruits from three individual clusters were pooled into a single replication.

Sequence and statistical analyses

Multiple sequence alignment was performed using ClustalX 1.83 (<http://www.ebi.ac.uk>) and MEGA5³⁹. The signal peptide, putative cleavage site, and putative glycosylation sites were predicted using the Signal P 3.0 server, NetNGlyc 1.0 Server (<http://www.cbs.dtu.dk/services/NetNGlyc/>), and the DictyOGlyc server (<http://www.cbs.dtu.dk/index/shtml/>), respectively.

Data were processed with SigmaPlot 10.0. Statistical analyses were performed using the statistical DPS (v3.0) package. A two-tailed *t*-test or Duncan multiple range test were used to determine significance at the 5% level.

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Conflict of interest

The authors declare that they have no conflict of interest.

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