Cloning and Functional Analysis of TaWRI1Ls, the Key Genes for Grain Fatty Acid Synthesis in Bread Wheat

Fengping Yang 1,2, Guoyu Liu 1, Ziyan Wu 1, Dongxue Zhang 1, Yufeng Zhang 1, Mingshan You 1, Baoyun Li 1, Xiuhai Zhang 2,* and Rongqi Liang 1,*

1 Key Laboratory of Crop Heterosis and Utilization (MOE) and Beijing Key Laboratory of Crop Genetic Improvement, China Agricultural University, Beijing 100193, China; yangfengping@baafs.net.cn (F.Y.); guoyuliu16@163.com (G.L.); wuziyan1125@163.com (Z.W.); zhangdx27@163.com (D.Z.); zhangyufeng@cau.edu.cn (Y.Z.); msysyou67@cau.edu.cn (M.Y.); baoyunli@cau.edu.cn (B.L.)

2 Beijing Academy of Agriculture and Forestry Sciences, Beijing 100097, China

* Correspondence: zhangxiuhai@baafs.net.cn (X.Z.); liangrq@cau.edu.cn (R.L.)

Abstract: WRINKLED1 (WRI1), an APETALA2 (AP2) transcription factor (TF), critically regulates the processes related to fatty acid synthesis, storage oil accumulation, and seed development in plants. However, the WRI1 genes remain unknown in allohexaploid bread wheat (Triticum aestivum L.). In this study, based on the sequence of Arabidopsis AtWRI1, two TaWRI1Ls genes of bread wheat, TaWRI1L1 and TaWRI1L2, were cloned. TaWRI1L2 was closely related to monocotyledons and clustered in one subgroup with AtWRI1, while TaWRI1L1 was clustered in another subgroup with AtWRI3 and AtWRI4. Both were expressed highly in the developmental grain, subcellular localized in the nucleus, and showed transcriptional activation activity. TaWRI1L2, rather than TaWRI1L1, promoted oil body accumulation and significantly increased triglyceride (TAG) content in tobacco leaves. Overexpression of TaWRI1L2 compensated for the functional loss of AtWRI1 in an Arabidopsis mutant and restored the wild-type phenotypes of seed shape, generation, and fatty acid synthesis and accumulation. Knockout of TaWRI1L2 reduced grain size, 1000 grain weight, and grain fatty acid synthesis in bread wheat. Conclusively, TaWRI1L2, rather than TaWRI1L1, was the key transcriptional factor in the regulation of grain fatty acid synthesis in bread wheat. This study lays a foundation for gene regulation and genetic manipulation of fatty acid synthesis in wheat genetic breeding programs.

Keywords: bread wheat; TaWRI1; fatty acid synthesis; functional analysis; transcription factor

1. Introductions

Bread wheat (Triticum aestivum L.) is one of the most important staple crops in China and in the world. Triacylglycerols (TAGs) are one of the main classes of storage components in wheat grains, accounting for about 1.5–3% of the dry weight of the grains [1–3]. They are mainly stored in the embryo and aleurone layer and can be extracted from milling byproducts such as the bran and germ of the wheat. TAGs are not only necessary for wheat seed germination, plant growth, and stress resistance [4,5], they are also nutritious for human beings [1].

WRINKLED1 (WRI1), discovered by Focks and Benning in Arabidopsis seed epidermis shrunken mutants, directly regulates fatty acid synthesis and glycolysis in the fatty acid synthesis pathway of plant seeds [6]. As an AP2 gene family member, its protein sequence contains two conserved AP2/ERF domains [7–9], and its conserved domain sequences have higher homology with those of the AINTEGUMENTA (ANT)-like group and are therefore considered part of the ANT-like group [7,10].

The WRII transcription factor acts as a regulator and binds to the conserved AW-box [CnTrg(n)CG] element in the proximal upstream region of these genes coding for proteins that function in various metabolic steps of fatty acid synthesis, and directly promotes the expression of these genes [9,11–15]. It is now clear that WRII directly binds the proximal
upstream region of pyr dehydrogenase (PDHE1α), plastidic pyruvate kinase (PI-PKα, PI-PKβ1), ketoacyl-ACP synthases (KAS1), acetyl-CoA carboxylase (BCCP2), enoyl-ACP reductase 1 (ENR1), acyl-carrier protein (ACP1), and sucrose synthase (SUS2), and regulates the expression of these targets [9]. Therefore, WRI1 is regarded as a distributor of the direction of carbon flow in seeds. WRI1 also stabilizes root auxin levels by activating the auxin transporter PIN (PIN-FORMEDs) and the auxin-degrading protein Gretchen Hagen 3.3 (GH3.3) [5]. LEAFY COTYLEDON1 and 2 (LEC1 and LEC2) and FUSC3A (FUS3) are upstream regulators of WRI1 [16–19]; LEC2 directly acts on WRI1 and regulates fatty acid metabolism during seed maturation [16].

In Arabidopsis, three genes, AtWRI2, AtWRI3, and AtWRI4, belonging to the AP2/EREBP family, with high homology to the AtWRI1 gene, have been cloned [20]. AtWRI1, AtWRI3, and AtWRI4 are all involved in triggering acyl chain production, but the expression profiles of these three genes are quite different. Unlike AtWRI3 and AtWRI4, AtWRI1 is mainly responsible for regulating the synthesis of fatty acids in seeds [20]. At present, the WRI1 genes have been cloned in Arabidopsis, castor, coconut, camellia, corn, Brassica napus, and other plants [6,21–27]. These WRI1 genes play a key regulatory role in promoting the synthesis and accumulation of fatty acids in seeds.

Overexpression of WRI1 was used to drive carbon flux towards fatty acid synthesis in leaves or seeds in some crops, such as tobacco, maize, rice, and camellina [22,24–27]. Overexpression of exogenous AtWRI1 in rice not only increased the content of fatty acids in rice seeds but also increased the content of starch in endosperm [26]. Maize ZmWRI1a and ZmWRI1b, derived from gene replication during evolution, encoded functional proteins and could restore the wrinkle phenotype and low fatty acid content of Arabidopsis WRI1-4 mutant seeds [27]. The expression of ZmWRI1 in maize seeds increased the oil content of seeds by 46% compared with the control [22].

The synthesis and accumulation of fatty acids in grains play a crucial role not only in grain germination and seedling morphogenesis [4] but also in human nutrition and grain storage. Unlike dicotyledonous plants such as Arabidopsis, the fatty acids in wheat seeds are mainly accumulated and stored in the embryo. As bread wheat is an allohexaploid crop, the WRI1 genes may have gene redundancy and functional differentiation. The purpose of this study was to clone the wheat WRI1 genes from the wheat genome, preliminarily analyze their function, and therefore provide an important reference for further research on the regulation of wheat seed fatty acids to meet genetic breeding objectives.

2. Results
2.1. Two Orthologs of AtWRI1, as Candidate Genes, Were Isolated from Wheat Genome

According to the deduced amino acid sequence of Arabidopsis AtWRI1 (At3g54320) and by BLASTP alignment with wheat genome, two candidate AP2 genes containing AP2/ERF domain sequences with the highest homology (82.1% and 84.2%, respectively) to AtWRI1 were screened out; these were the orthologs of AtWRI1. These two genes were selected as candidate TaWRI1Ls genes in this study, named TaWRI1L1 (TraesCS5A02G229400, TraesCS5B02G221600, TraesCS5D02G229400) and TaWRI1L2 (TraesCS5A02G141700, TraesCS5D02G141700, TraesCS5D02G150500), respectively.

Using two pairs of primers (TaWRI1L1-F/TaWRI1L1-R and TaWRI1L2-F/TaWRI1L2-R) and JSM1 (jingshengmai1) cDNA, RT-PCR was performed to obtain the target products. The cDNA sequence alignment showed that JSM1TaWRI1L1-5A, -5B, and -5D were 98.24%, 95.91%, and 100% homologous to TraesCS5A02G221600, TraesCS5B02G220400, TraesCS5D02G229400, and TaWRI1L2 (TraesCS5A02G141700, TraesCS5D02G150500), respectively.

Using two pairs of primers (TaWRI1L1-F/TaWRI1L1-R and TaWRI1L2-F/TaWRI1L2-R) and JSM1 (jingshengmai1) cDNA, RT-PCR was performed to obtain the target products. The cDNA sequence alignment showed that JSM1TaWRI1L1-5A, -5B, and -5D were 98.24%, 95.91%, and 100% homologous to TraesCS5A02G221600, TraesCS5B02G220400, TraesCS5D02G229400, and TaWRI1L2 (TraesCS5A02G141700, TraesCS5D02G150500), respectively.
of different wheat varieties had high homology in their nucleotide and deduced amino acid sequences.

The N- and C-terminus sequences of the TaWRI1Ls were quite different. TaWRI1L1 started with a continuous non-polar amino acid P (Pro), but TaWRI1L2 was rich in multiple non-polar amino acids A (Aln) at the N-terminus (Figure S1A,B). That could be used as a sequence feature to distinguish these TaWRI1Ls. The N-terminus of TaWRI1L2 was enriched with multiple polar S (Ser) amino acids, which shared the same sequence characteristics with AtWRI1 and WRI1 of other monocotyledonous plants, such as maize and rice, suggesting that TaWRI1L2 may have a similar functional mechanism to that of AtWRI1.

The deduced amino acid sequences of eight monocotyledonous, eight dicotyledonous, and six oil plant WRI1s were selected to construct an evolutionary tree (Figure 1). TaWRI1L2 was closely related to the monocotyledonous plants’ BdWRI1, ZmWRI1a, ZmWRI1b, and OsWRI1, and clustered into an independent branch. Thus, TaWRI1L2 had the sequence characteristics of monocotyledonous WRI1. However, TaWRIL1 showed higher homology to CeWRI1, AtWRI3, and AtWRI4, and clustered in a subgroup. Therefore, TaWRIL1 may have functions similar to AtWRI3 or AtWRI4.

Figure 1. The evolutionary tree of TaWRI1Ls and WRI1s of other plants.

The bootstrap parameter was 1000.

2.2. TaWRI1L1 and TaWRI1L2 Were Predominantly Expressed in the Developmental Grain, Were Subcellular Localized in the Nucleus, and Showed Transcriptional Activation Activity

Using transcriptome data for JSM1 at six grain developmental stages, the expression profiles of TaWRI1Ls and 21 paralogs were analyzed. Two homoeologs of TaWRI1L2 were significantly expressed, and expression of TaWRI1L2-5A was the highest, followed by TaWRI1L2-5D (Figure 2A). Three homoeologs of TaWRI1L1 had generally lower expression levels than the two homoeologs of TaWRI1L2 but the levels were significantly higher than those of the paralogs (Figure 2B). This indicated that TaWRI1L1 and TaWRI1L2 play an
important role in the development of grain. Based on the expression trend, the expression levels of these two candidate genes decreased continuously with grain development. However, at DAF13, the middle-to-late leaf stage of embryonic development, their expression levels rebounded and appeared as a small peak.

Figure 2. Expression levels of TaWRI1Ls and their paralogs in Chinese Spring and Fielder. (A) TaWRI1Ls of Chinese Spring grains at different filling stages; (B) the 21 paralogs of Chinese Spring grains at different filling stages; (C) TaWRI1Ls of Fielder root, stem, leaf, and grain at different filling.

Bars represent the means ± SD of three biological replicates.

Our qPCR results (Figure 2C) revealed that the homologs of TaWRI1L1 were expressed at three time points (DAF 4, DAF 7, and DAF 10) and showed a downward trend. The homoeologs of TaWRI1L2 were expressed at all time points, reaching a peak at DAF 7. The homoeologs of TaWRI1L2 were expressed significantly—dozens of times—more than those of TaWRIL1. These results indicate that TaWRIL2 plays a major role in fatty acid synthesis in wheat developmental grains.

The data from the Wheat eFP Browser (Figure S2) also showed that TaWRI1L1 was mainly expressed in the wheat panicle, flag leaf, glume, and early stage of grain development, while TaWRI1L2 was mainly expressed in the wheat grain and root tip. TaWRI1L2 is mainly expressed during embryonic development, while TaWRIL1 is strongly expressed in endosperm and weakly expressed in embryos. From the perspective of embryonic development, both genes showed peak expression in the leaf late embryo stage, but the expression level of TaWRIL2 was dozens of times that of TaWRIL1.

The transient expression vectors pYBA1132-W1 (TaWRI1L1-5B) and pYBA1132-W2 (TaWRI1L2-5A) were constructed, respectively (Figure 3A), and used for transient transformation of onion epidermis for subcellular localization of TaWRI1Ls. The results (Figure 3B) showed that the expression of both candidate genes was localized in the nucleus of onion epidermal cells, illustrating that both belonged to the nuclear genes and accorded with the typical characteristics of transcription factors.

The pBD-W1 and pBD-W2 yeast transformation experiments (Figure 3C) showed that the yeast cells transformed with TaWRI1L1-5B or TaWRI1L2-5A genes appeared blue, indicating that the two TaWRI1Ls were expressed as fusion genes with GAL4 activity and activated the expression of the reporter gene LacZ in yeast AH109. These results confirmed that both TaWRI1L1 and TaWRI1L2 genes showed transcriptional activity.
2.3. TaWRI1L2, Instead of TaWRI1L1, Promoted Oil Body Accumulation and Significantly Increased Triglyceride Contents in N. benthamiana Leaves

After Agrobacterium GV3101 injection for 5–6 days, the clear outline of N. benthamiana leaf epidermal cells could be found under a laser confocal microscope (Figure 4A). The obvious accumulation of oil bodies stained with Nile red (indicated by the arrow) could be observed in the transient expressing TaWRI1L2 leaves, while almost no oil bodies were formed in the control and TaWRI1L1-transformed leaves.

To further determine the enhancement of oil body accumulation in TaWRI1Ls transient expression leaves, TAG contents in dried leaves were detected with an ORACLE universal fat tester; consistent with the Nile red staining result, expression of TaWRI1L2 can significantly increase the content of TAG in N. benthamiana leaves by 9–10% (Figure 4B). However,
transient expression of TaWRIL1 could not play an active role in promoting TAG increase in N. benthamiana leaves, which was not significantly different from that of the control.

In conclusion, the transient expression results showed that TaWRIL2 could positively regulate fatty acid synthesis in N. benthamiana leaves, while the regulatory effect of TaWRIL1 was not obvious, indicating that the functions of these two TaWRILs genes were significantly different.

2.4. Overexpression of TaWRIL2 Compensated for the Functional Loss of AtWRI1 in an Arabidopsis Mutant and Restored the Wild-Type Phenotypes of Seed Shape, Generation, and Fatty Acid Synthesis and Accumulation

To further confirm the effect of TaWRIL2 on fatty acid synthesis, the wild-type Arabidopsis Col-2 and wri1-1 mutant were transformed with TaWRIL2-5A using the dip flower method. The results showed that the wild-type Col2 seeds were round and plump, and the seed plumpness of Col2-OE-W2 transgenic lines was not significantly different from wild-type Col2 (Figure 5A). The wri1-1 mutant had shriveled seeds, a shrunken seed coat, and a darker seed coat color than wild-type Col2, while the wri1-1-OE-W2 transgenic lines almost recovered the seed shrinkage phenotype of the wri1-1 mutant and were not significantly different from wild-type Col2 with respect to seed shrinkage, plumpness, and the epidermis (Figure 5A). These results indicated that the expression of the TaWRIL2 gene in the wri1-1 mutant made up for the functional loss of the WRI1 gene in the mutant and restored the normal level of fatty acid synthesis and accumulation in seeds, thus showing the wild-type seed phenotype.

The transgenic seeds were sterilized and inoculated on MS medium with 3% sucrose to observe the growth of seedlings. The 10-day-old seedlings were collected and stained with Sudan 7B to record the distribution of fatty acids by anatomical microscope. The results showed that wild-type Col2 seeds germinated on the culture medium for 1–2 days and grew into seedlings normally; after Sudan 7B staining, only the hypocotyl and root system of the plant turned light red (Figure 5B). Compared with the Col-2 seedling, the Col2-OE-W2 seedlings grew slightly slower, with expanded cotyledons and hypocotyls, dark color, and relatively strong plants; after Sudan 7B staining, cotyledons, hypocotyls, and the roots of Col2-OE-W2 seedlings were dyed purplish red (Figure 5B), showing that the Col2-OE-W2 seedlings accumulated significantly more fatty acids than wild-type Col2. The wri1-1-W2 transgenic seeds germinated slightly more slowly than the wild-type Col-2, and their seedlings grew normally after germination. After Sudan 7B staining, the wri1-1-W2 transgenic seedlings had light red hypocotyls and roots and accumulated fatty acids similarly to Col2 (Figure 5B). In a word, the overexpression of the exogenous TaWRIL2 gene could make the wri1-1 mutants restore the wild-type phenotype and significantly improve the utilization of sucrose to activate fatty acid synthesis and accumulation in plants.

Using the seedlings of T3 homozygous lines overexpressing TaWRIL2 for 10 days after germination, the expression levels of the target genes Pl-PKβ1, BCCP2, and KASI were studied. Each target gene had a significantly higher expression level in Col2-OE-W2 lines than in the wild-type Col2 (Figure 5C), while three target genes expressed in the wri1-1-OE-W2 transgenic line could restore the expression level in the wri1-1 mutant or surpass the expression level in the wild-type Col2 (Figure 5D), indicating that TaWRIL2 could activate the three downstream target genes of the wri1-1 mutant and play a regulatory role in the fatty acid synthesis pathway.

The seeds of the T3 homozygous Col2-OE-W2 lines were analyzed for eight fatty acid components (C16:0, C18:0, C18:1, C18:2, C18:3, C20:0, C20:1, and C22: 1). Compared with wild-type Col2, eight fatty acid components in the transgenic lines were increased to different degrees, and the increases in Col2-OE-W2-4 and Col2-OE-W2-9 reached significant levels (Figure 5E,G). Compared with the wri1-1 mutant, three transgenic lines (wri1-1-W2-1, wri1-1-W2-2, and wri1-1-W2-3) overexpressing TaWRIL2 showed increases in eight fatty acid components and total fatty acid content (Figure 5F,H), similar to the content of wild-type Col2. Taken together, overexpression of exogenous TaWRIL2 could
compensate for the functional loss of AtWRI1 in mutants and increase the fatty acid content of Arabidopsis seeds.

Figure 5. Functional analyses of TaWRI1Ls in the Arabidopsis mutant. (A) Seeds of Col2, wri1-1 mutant, and their respective OE lines. Bar = 100 µm. (B) Upper: seedlings of Col2, wri1-1 mutant, and their respective OE lines stained with Sudan Red 7B. Bar = 500 mm. (C) Expression of Pl-PKβ1, BCCP2, and KASI in Col2 and its OE lines. (D) Expression of Pl-PKβ1, BCCP2, and KASI in wri1-1 and its OE lines. (E) Seed fatty acid contents of Col2 and its OE lines. (F) Seed fatty acid contents of wri1-1 and its OE lines. (G) Seed fatty acid components of Col2 and its OE lines. (H) Seed fatty acid components of wri1-1 and its OE lines. Data are represented as means ± SD from independent biological replicates, and p-values are indicated by Student’s t-test (* p < 0.05, ** p < 0.01).

2.5. Knockout of TaWRI1L2 Reduced the Grain Size, 1000 Grain Weight, and Grain Fatty Acid Synthesis in Bread Wheat

To further study the function of the TaWRI1L2 gene in wheat, we knocked out this gene using CRISPR. The gRNA sequences for TaWRI1L2 were screened out on the website https://crispr.bioinfo.nrc.ca/WheatCrispr/ (accessed on 12 November 2018) (Figure 6A), and a vector pBUE411-TaU3p-W2 containing two editing sites was constructed for the transformation of young embryos. Twelve T\textsubscript{0} plants were positive for the bar gene. After
two generations of pedigree planting and screening, two homozygous lines, KO-1 and KO-2, derived from two independent T₀ plants, were obtained. The KO-1 line was edited at the first site by adding a base, and the KO-2 line was edited at the second site by deletion of four bases (Figure 6A,B).

Figure 6. Phenotypes of wheat TaWRI1L2 knock-out mutants. (A) The location of gRNA. (B) The edited sites of the KO mutants. (C) Mature grains and their generation of WT and KO mutants. Bars = 5 mm. (D) Generation of WT and KO mutants. Bars = 5 mm. (E) The grain lengths of WT and KO mutants. n = 3. (F) The grain widths of WT and KO mutants. n = 3. (G) The 1000 grain weight of WT and KO mutants. n = 3. (H) Grain fatty acid components of WT and its KO mutants. Data in (C,D) are represented as means ± SD from independent biological replicates, and p-values are indicated by Student’s t-test (** p < 0.01, *** p < 0.001).

Grains of the KO lines became smaller (Figure 6C) than the receptor Fielder. Meanwhile, grain lengths, grain widths, and 1000 grain weights were also reduced to different degrees (Figure 6E–G). There were no significant differences in imbibition capacity, germination vigor, germination speed (Figure 6D), and developmental process between the KO lines and Fielder.

Five fatty acid components (C16:0, C18:0, C18:1, C18:2, and C18:3) and total fatty acid analyses were performed on the seeds of the KO lines as well as the control Fielder, and the results showed that the contents of oleic acid (C18:1) and linolenic acid (C18:3), were lower in Fielder and not detected in the transgenic lines. The contents of palmitic acid (C16:0) and linoleic acid (C18:2) were significantly lower in KO lines than in Fielder, which resulted in a significant reduction in total fatty acid content in seeds of the two KO lines—only up to 18–20% of the control (Figure 6H). These results illustrated that functional loss of TaWRI1L2 reduced fatty acid synthesis in wheat grain.

Conclusively, the above functional analyses showed that TaWRI1L2 had a biological activity and functions similar to the A1WRII gene, such as activating the expression of key genes in the fatty acid synthesis pathway and promoting the synthesis and accumulation of seed fatty acids. The regulatory effect of TaWRI1L1 is not obvious, indicating that there are significant differences in the functions of the two TaWRI1Ls genes.

3. Discussion

The fatty acids in plant seeds are essential for seed germination and plant growth [4,5]. Studying the coordination mechanism between the synthesis of fatty acids and the synthesis of starch, protein, and other components in wheat grains could promote directional breeding to meet wheat end-product demand.
Dicotyledonous and monocotyledonous seed structures are distinctly different morphologically due to different endosperm/embryo volume ratios during differentiation. In Arabidopsis thaliana, the embryo accumulated lipids and proteins in the cotyledons by depleting nutrients in the endosperm tissue during maturation, and the endosperm cells subsequently degenerated [28]. In cereal crops, embryos are small in size and mainly accumulate lipids and globulins [29], while endosperms persist after cellularization and mainly accumulate starch and storage proteins and undergo programmed cell death (PCD) without degradation [30].

Due to obvious differences in the developmental trends of monocotyledonous and dicotyledonous plants, the expression patterns of homologous genes may be different. The expression of AtWRI1 and its regulated target genes started at 4 days after flowering and then increased rapidly to a maximum at day 8 [6,7,11]. TaWRIL1 and TaWRIL2 were expressed at high levels in the developmental grain, while TaWRIL1 was mainly expressed in the embryo and TaWRIL2 in the endosperm. However, TaWRIL2 expression was significantly—dozens of times—higher than that of TaWRIL1. TaWRIL6 had the sequence characteristics of the monocotyledonous plant WRI1, and TaWRIL1 may have functions similar to WRI3 or WRI4. Therefore, TaWRIL2 and its function was further analyzed in tobacco, Arabidopsis, and wheat in this study.

WRI1 is the regulator of carbohydrate metabolism during the stage of seed filling [6], so its functional loss might affect grain starch synthesis and grain size as well. The Arabidopsis endosperm degenerated and disappeared at the late stage of maturity, and the cotyledons were the main place for oil preservation. The WRI1 mutation led to a significant decrease in TAG content, so the Arabidopsis seeds shrank significantly [6]. The embryo of bread wheat was the main place for oil storage and could not cause obvious grain shrinkage when grain fatty acid synthesis was affected. Therefore, grain shrinkage was not the phenotype of TaWRI1. We found that there was no difference in germination and subsequent growth and development between the KO grains and wild-type grains, probably because the starch stored in the endosperm was the major carbon source and energy provider during the germination stage. The overexpression of AtWRI1 in rice can lead to an increase in starch content and a significant decrease in fatty acid content in WRI1 rice endosperm [26]. In our study, the knockout of TaWRIL2 in wheat resulted in a significant decrease in grain size and weight. Thus, it can be seen that WRI1 acts in different ways in various plant and organs or tissues, the mechanisms of which need to be further explored in future research.

Maize ZmWRI1a and ZmWRI1b, derived from gene replication during evolution, shared 85.32% amino acid sequence similarity [27]. ZmWRI1a was mainly expressed in the maize embryo and had two expression peaks at the early stage of grain development and at the end of the grain filling stage [27], which was consistent with the expression pattern of TaWRIL1. ZmWRI1b was highly expressed in the endosperm and had a peak during grain development [27] which resembled the expression pattern of TaWRIL1L1. However, TaWRIL1L1 and TaWRIL1L2 had 38.64% amino acid sequence similarity, and TaWRIL1L2 had higher homology with OsWRI1 and ZmWRI1 than TaWRIL1L1. Therefore, TaWRIL1L2, along with OsWRI1 and ZmWRI1, had a functional mechanism similar to AtWRI1.

The expression of ZmWRI1 increased the oil content of seeds by 46% without affecting seed germination or plant growth, and there was no significant difference in yields between transgenic lines and control lines [22]. The C16 and C18 fatty acid contents of maize seeds were significantly increased in the lines overexpressing ZmWRI1a compared to the control [27]. Our results showed that TaWRIL1L2-5A was more expressed than TaWRIL1L2-5B and TaWRIL1L2-5D in wheat seeds, and that its over-expression increased oil production and accumulation in tobacco leaves, while its knockout reduced the oil contents in wheat seeds. Therefore, we can manipulate the key regulator TaWRIL1L2-5A in future breeding programs to regulate TAG synthesis to produce more vegetable oil in modified biomass for increased oil nutritional value or less oil for the longer storage life of wheat seeds.
4. Materials and Methods

4.1. Cloning and Sequence Analysis of Target TaWRI1Ls Gene

Using the induced amino acid sequence of the AP2 protein domain of the Arabidopsis thaliana AtWRI1 gene (At3g54320), BLASTP alignment was performed on the Chinese Spring genome data using the website database at http://plants.ensembl.org/index.html (accessed on 16 April 2017) to screen out the homology genes. According to the ortholog data of the gene provided by the database, many candidate TaWRI1Ls (TaWRI1-like) genes containing AP2/ERF domain sequences were found in the wheat genome. Two pairs of primers, TaWRI1L1-F/ORF-TaWRI1L1-R and TaWRI1L2-F/TaWRI1L2-R, were designed in the conserved regions at the 5’ and 3’ ends of TaWRI1Ls to amplify the ORF region (Table 1).

Table 1. The primers for RT-PCR and qPCR.

| Primer Names | Sequence (5′–3’) | Length of Product |
|--------------|-----------------|------------------|
| TaWRI1L1-F   | CACGAGCGCAATGGCAAAG | 1204 bp |
| TaWRI1L1-R   | TCACAAGTCCAGCTCGAA | 1204 bp |
| TaWRI1L2-F   | CCACCATTGACACAGCACAG | 1352 bp |
| TaWRI1L2-R   | AAACCTGAAACCTCTTGTGG | 1352 bp |
| PI-PKb1-F    | CCATATGAGTGAGATTTGC | 119 bp |
| PI-PKb1-R    | GCCATATGAGTGAGATTTGC | 119 bp |
| BCCP2-F      | GCAAGCTGTGGACTTGTGAC | 100 bp |
| BCCP2-R      | TGAAGGCAGAGAAGGCGACTAC | 100 bp |
| KAS1-F       | ACAAGCTGTGGACTTGTGAC | 119 bp |
| KAS1-R       | TGAAGGCAGAGAAGGCGACTAC | 119 bp |

The bread wheat variety “Jingshengmai1 (JSM1)”, bred by the Beijing Agricultural Biotechnology Research Center, was used to extract total RNA from grains 4–22 days after flowering (DAF) for RT-PCR amplification and RNA-sequence analysis.

The evolutionary relationship between TaWRI1Ls candidate genes and WRI1 genes of model plants (such as Arabidopsis thaliana, maize, and cabbage) was analyzed with MEGA7.0 (Mega Limited, Auckland, New Zealand). The evolutionary tree was constructed using the neighbor-joining method (NJ), and the bootstrap parameter was 1000.

4.2. Expression Analysis of TaWRI1Ls

According to the description of the morphological and developmental stages of wheat grains [31], the grains of “Jingshengmai 1” was collected at 4DAF (pre-embryo), 6DAF (transition), 10DAF (leaf early), 13DAF (leaf mid), 16DAF (leaf late), and 19DAF (mature) for qPCR and RNA-seq expression analysis. The expression of TaWRI1Ls in JSM1 grains was investigated by qPCR using an SYBR Premix ExTaq Kit (Takara Branch, Beijing, China) with three biological replicates. The relative expression levels were evaluated by the $2^{-\Delta\Delta Ct}$ method [32].

The expression data for the TaWRI1Ls gene were downloaded from the Wheat eFP Browser (http://bar.utoronto.ca/efp_wheat/cgi-bin/efpWeb.cgi (accessed on 8 May 2017)) for expression profile analysis [33].

4.3. Subcellular Localization and Yeast Activation Analysis of TaWRI1Ls

The ORF fragments of TaWRI1L1-5B (TraesCS5B02G220400) and TaWRI1L2-5A (TraesCS5A02G14170) of the candidate TaWRI1Ls gene (with restriction sites) were cloned into the downstream of CaMV 35S promoter in the GFP fusion vector pYBA1132, and the recombinant vectors, named pYBA1132-W1 and pYBA1132-W2 (Figure 3A), respectively, were used for subcellular localization by transient expression. The onion epidermis was bombarded with a BIO-RAD high-pressure helium PDS1000 gene gun (BIO-RAD Co., Hercules, CA, USA), and the subcellular localization of the target genes was detected by confocal laser observation 24 h later.
The ORF fragments of the TaWRI1L1-5B and TaWRI1L2-5A genes were cloned into the yeast BD expression vectors and the recombinant vectors, named pBD-W1 and pBD-W2, were introduced into the yeast strain AH109 to observe the color of the yeast cells after staining with X-gal to determine whether the target genes had transcriptional activation activity.

4.4. Functional Analysis of TaWRI1Ls in N. benthamiana Leaves

The TaWRI1L1-5B and TaWRI1L2-5A fragments were respectively constructed into the overexpression pYBA1302 vector containing the herbicide selection marker, and the recombinant vectors pYBA1302-W1 and pYBA1302-W2 were respectively transferred into Agrobacterium strain GV3101. Taking the Agrobacterium containing the empty vector as the control, the Agrobacterium solution containing the recombinant vector was injected into the back of N. benthamiana leaves at the same position.

After 3–4 days, oil accumulation could be observed in injected leaves. After 6–7 days, the injected leaves were harvested for intracellular oil analysis. The oil bodies inside the cells were observed by means of a laser scanning microscope (red fluorescence at wavelengths 560 nm and 615 nm) after the leaves were stained with 10 µg/mL Nile red solution (Sigma-N3013) for 30 min at room temperature and washed with 0.1 M Tris-HCl buffer (pH 8) for 10 min [24].

Triglyceride (TAG) contents in tobacco leaves were detected with an ORACLE universal fat tester (CEM Co., Matthews, NC, USA). One gram of dried leaves of each sample was measured with three biological replicates to calculate the percentage (w/w) of TAGs in the dry leaves.

4.5. Functional Analysis of TaWRI1Ls in Arabidopsis

The wild-type Arabidopsis thaliana Columbia 2 (Col2) and its wri1-1 mutant (obtained from the Arabidopsis Information Resource) were used as receptors, respectively, and the transformation was performed by the floral dip method using the Agrobacterium strain GV3101 containing the overexpression vectors pYBA1302-W1 and pYBA1302-W2. The transgenic Arabidopsis seeds were sterilized and sown on ½ MS solid medium with 3% sucrose. One week after germination, the seedlings were observed and stained with Sudan Red 7B to observe the degree of oil accumulation. Red staining indicates the accumulation of neutral lipids in Arabidopsis seedlings [18].

After harvest, the content of each fatty acid component in the seeds was determined with three biological replicates by gas chromatography [34], according to the manufacturer’s instructions.

Three target genes, PI-PKβ1 (At5g52920), BCCP2 (At5g15530), and KASI (At5g46290), were selected [9] to analyze the regulatory effects of the overexpressed TaWRI1L2 gene. The sequences of the qPCR primer were shown in Table 1.

4.6. Functional Analysis of TaWRI1Ls in Bread Wheat

The gRNA fragment of the TaWRI1L2-5A gene was screened on the website https://crispr.bioinfo.nrc.ca/WheatCrispr/ (accessed on 12 November 2018) and used to construct the CRISPR vector pBUE411-TaU3p-W2. The spring wheat cultivar “Fielder” embryos were transformed with Agrobacterium strain EHA105 containing this CRISPR vector, and the transgenic plants were obtained after callus induction, callus differentiation, and plant regeneration.

T0–T2 transgenic plants and their recipient variety Fielder were grown in the greenhouse. The leaves of the transgenic plants were used to extract genomic DNA. The PCR was performed using specific PCR primers and the high-fidelity Taq PCR mix (SuperStar Co., Beijing, China), and the target fragments separated by 1% agarose gel electrophoresis were sequenced (Shanghai Shenggong Co., Shanghai, China) to determine the editing sites.

After harvest, the contents of each fatty acid component in the seeds were measured as mentioned above. The grain lengths, grain widths and 1000 grain weights of Fielder and the knockout lines were recorded with three biological replicates using the rapid SC-G
grain appearance quality image analysis system (Hangzhou WSeen Detection Technology Co., Ltd., Hangzhou, China) [35].

4.7. Statistical Analysis

Means and standard deviations were calculated for each treatment and the statistical differences were analyzed by Student’s t-test at p < 0.05 using SPSS 16.0 (SPSS, Inc., Chicago, IL, USA).

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/ijms23105293/s1.

Author Contributions: R.L., F.Y. and X.Z. conceived the project. F.Y. performed the experiments. G.L. performed qPCR of TaWRi1. D.Z., Y.Z. and M.Y. helped with planting the materials. F.Y., R.L. and Z.W. wrote the manuscript. R.L., B.L., G.L. and X.Z. revised the manuscript. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by National Natural Science Foundation, grant number 31290213.

Institutional Review Board Statement: The study did not require ethical approval.

Informed Consent Statement: Not applicable.

Data Availability Statement: Data supporting this work are available within the paper and its supplementary files. RNA-seq data were uploaded to the NCBI with the accession number PRJNA791126.

Acknowledgments: We thank Na Song (China Agricultural University) for wheat transformation.

Conflicts of Interest: The authors declare no conflict of interest.

References
1. Chung, K.O. Relationship of polar lipid content to mixing requirement and loaf volume potential of hard red winter wheat flour. Cereal Chem. 1982, 59, 14–20.
2. Zhang, Y.P. Prospect on the research & development of good quality wheat varieties in the turn of the new century. Cereal Feed. Ind. 1998, 9, 4–6. (In Chinese with English abstract).
3. Chi, X.Y.; Tian, J.C. Advances in the Research of Wheat Lipids. J. Triticeae Crops 2005, 5, 120–125. (In Chinese with English abstract).
4. Cernac, A.; Andre, C.; Hoffmann-Benning, S.; Benning, C. WR1 is required for seed germination and seedling establishment. Plant. Physiol. 2006, 141, 745–757. [CrossRef] [PubMed]
5. Kong, Q.; Ma, W.; Yang, H.; Ma, G.; Mantyla, J.J.; Benning, C. The Arabidopsis WRINKLED1 transcription factor affects auxin homeostasis in roots. J. Exp. Bot. 2017, 68, 4627–4634. [CrossRef]
6. Focks, N.; Benning, C. wrinkled1: A novel, low-seed-oil mutant of Arabidopsis with a deficiency in the seed-specific regulation of carbohydrate metabolism. Plant Physiol. 1998, 118, 91–101. [CrossRef]
7. Cernac, A.; Benning, C. WRINKLED1 encodes an AP2/EREB domain protein involved in the control of storage compound biosynthesis in Arabidopsis. Plant J. 2004, 40, 575–585. [CrossRef]
8. Ohto, M.A.; Fischer, R.L.; Goldberg, R.B.; Nakamura, K.; Harada, J.J. Control of seed mass by APETALA2. Proc. Natl. Acad. Sci. USA 2005, 102, 3123–3128. [CrossRef]
9. Maeo, K.; Tokuda, T.; Ayame, A.; Mitsui, N.; Kawai, T.; Tsukagoshi, H.; Ishiguro, S.; Nakamura, K. An AP2-type transcription factor, WRINKLED1, of Arabidopsis thaliana binds to the AW-box sequence conserved among proximal upstream regions of genes involved in fatty acid synthesis. Plant J. 2009, 60, 476–487. [CrossRef]
10. Shigyo, M.; Hasebe, M.; Ito, M. Molecular evolution of the AP2 subfamily. Gene 2006, 366, 256–265. [CrossRef]
11. Baud, S.; Lepiniec, L. Regulation of de novo fatty acid synthesis in maturing oilseeds of Arabidopsis. Plant Physiol. Biochem. 2009, 47, 448–455. [CrossRef] [PubMed]
12. Baud, S.; Lepiniec, L. Physiological and developmental regulation of seed oil production. Prog. Lipid Res. 2010, 49, 235–249. [CrossRef] [PubMed]
13. Liu, H.; Zhai, Z.; Kuczynski, K.; Keerewetawee, J.; Schwender, J.; Shanklin, J. WRINKLED1 Regulates BIOTIN ATTACHMENT DOMAIN-CONTAINING Proteins that Inhibit Fatty Acid Synthesis. Plant Physiol. 2019, 181, 55–62. [CrossRef]
14. Fukuda, N.; Ikawa, Y.; Aoyagi, T.; Kozaki, A. Expression of the genes coding for plastidic acetyl-CoA carboxylase subunits is regulated by a location-sensitive transcription factor binding site. Plant Mol. Biol. 2013, 82, 473–483. [CrossRef]
15. Chen, L.; Zheng, Y.; Dong, Z.; Meng, F.; Sun, X.; Fan, X.; Zhang, Y.; Wang, M.; Wang, S. Soybean (Glycine max) WRINKLED1 transcription factor, GmWR11a, positively regulates seed oil accumulation. Mol. Genet. Genom. 2018, 293, 401–415. [CrossRef] [PubMed]
16. Baud, S.; Mendoza, M.S.; To, A.; Harscoet, E.; Lepiniec, L.; Dubreucq, B. WRINKLED1 specifies the regulatory action of LEAFY COTYLEDON2 towards fatty acid metabolism during seed maturation in Arabidopsis. Plant J. 2007, 50, 825–838. [CrossRef] [PubMed]

17. Mu, J.; Tan, H.; Zheng, Q.; Fu, F.; Liang, Y.; Zhang, J.; Yang, X.; Wang, T.; Chong, K.; Wang, X.J.; et al. LEAFY COTYLEDON1 is a key regulator of fatty acid biosynthesis in Arabidopsis. Plant Physiol. 2008, 148, 1042–1054. [CrossRef]

18. Zhang, M.; Cao, X.; Jia, Q.; Ohlrogge, J. FUSCA3 activates triacylglycerol accumulation in Arabidopsis seedlings and tobacco BY2 cells. Plant J. 2016, 88, 95–107. [CrossRef]

19. Song, J.; Xie, X.; Chen, C.; Shu, J.; Thapa, R.K.; Nguyen, V.; Bian, S.; Kohalmi, S.E.; Marsolais, F.; Zou, J.; et al. LEAFY COTYLEDON1 expression in the endosperm enables embryo maturation in Arabidopsis. Nat. Commun. 2021, 12, 3963. [CrossRef]

20. To, A.; Joubes, J.; Barthole, G.; Lecureuil, A.; Scagnelli, A.; Jasinski, S.; Lepiniec, L.; Baud, S. WRINKLED transcription factors orchestrate tissue-specific regulation of fatty acid biosynthesis in Arabidopsis. Plant Cell 2012, 24, 5007–5023. [CrossRef]

21. Liu, J.; Hua, W.; Zhan, G.; Wei, F.; Wang, X.; Liu, G.; Wang, H. Increasing seed mass and oil content in transgenic Arabidopsis by the overexpression of wr1-like gene from Brassica napus. Plant Physiol. Biochem. 2010, 48, 9–15. [CrossRef] [PubMed]

22. Shen, B.; Allen, W.B.; Zheng, P.; Li, C.; Glassman, K.; Ranch, J.; Nübel, D.; Tarczynski, M.C. Expression of ZmLEC1 and ZmWRI1 increases seed oil production in maize. Plant Physiol. 2010, 153, 980–987. [CrossRef] [PubMed]

23. Wu, X.L.; Liu, Z.H.; Hu, Z.H.; Huang, R.Z. BnWRI1 coordinates fatty acid biosynthesis and photosynthesis pathways during oil accumulation in rapeseed. J. Integr. Plant Biol. 2014, 56, 582–593. [CrossRef] [PubMed]

24. An, D.; Kim, H.; Ju, S.; Go, Y.S.; Kim, H.U.; Suh, M.C. Expression of Camelina WRINKLED1 Isoforms Rescue the Seed Phenotype of the Arabidopsis wr1 Mutant and Increase the Triacylglycerol Content in Tobacco Leaves. Front. Plant Sci. 2017, 8, 34. [CrossRef]

25. Sun, R.; Ye, R.; Gao, L.; Zhang, L.; Wang, R.; Mao, T.; Zheng, Y.; Li, D.; Lin, Y. Characterization and Ectopic Expression of CoWRI1, an AP2/EREBP Domain-Containing Transcription Factor from Coconut (Cocos nucifera L.) Endosperm, Changes the Seeds Oil Content in Transgenic Arabidopsis thaliana and Rice (Oryza sativa L.). Front. Plant Sci. 2017, 8, 63. [CrossRef]

26. Yang, J.; Tian, R.; Gao, Z.; Yang, H. Characterization of AtWRI1 in fatty acids and starch synthesis in rice. Biosci. Biotechnol. Biochem. 2019, 83, 1807–1814. [CrossRef]

27. Pouvreau, B.; Baud, S.; Vernoud, V.; Morin, V.; Py, C.; Gendrot, G.; Pichon, J.; Rouster, J.; Paul, W.; Rogowsky, P.M. Duplicate maize wrinkled1 transcription factors activate target genes involved in seed oil biosynthesis. Plant Physiol. 2011, 156, 674–686. [CrossRef]

28. Olsen, O.A. Nuclear endosperm development in cereals and Arabidopsis thaliana. Plant Cell 2004, 16 (Suppl. S1), S214–S227. [CrossRef]

29. Sreenivasulu, N.; Wobus, U. Seed-development programs: A systems biology-based comparison between dicots and monocots. Annu. Rev. Plant Biol. 2013, 64, 189–217. [CrossRef]

30. Domínguez, F.; Cejudo, F.J. Programmed cell death (PCD): An essential process of cereal seed development and germination. Front. Plant Sci. 2014, 5, 366. [CrossRef]

31. Xiang, D.; Quilichini, T.D.; Liu, Z.; Gao, P.; Pan, Y.; Li, Q.; Nilsen, K.T.; Venglat, P.; Esteban, E.; Pasha, A.; et al. The Transcriptional Landscape of Polyploid Wheats and Their Diploid Ancestors during Embryogenesis and Grain Development. Plant Cell 2019, 31, 2888–2911. [CrossRef]

32. Livak, K.J.; Schmittgen, T.D. Analysis of relative gene expression using real-time quantitative PCR and the 2(-Delta Delta C(T)) Method. Methods A Companion Methods Enzymol. 2001, 2888, 2911. [CrossRef]

33. Winter, D.; Vineagar, B.; Nahal, H.; Ammar, R.; Wilson, G.V.; Provart, N.J. An “Electronic Fluorescent Pictograph” browser for exploring and analyzing large-scale biological data sets. PLoS ONE 2007, 2, e718. [CrossRef]

34. Li, X.L.; Zhang, L.Q.; Zhang, X.H. Improvement of unsaturated fatty acid composition of maize by AcD12 gene with dual function desaturase. Mol. Plant Breed. 2017, 15, 4805–4812. (In Chinese with English abstract).

35. Guan, P.F.; Di, N.; Mu, Q.; Shen, X.Y.; Wang, Y.F.; Wang, X.B.; Yu, K.H.; Song, W.J.; Chen, Y.M.; Xin, M.M.; et al. Use of near-isogenic lines to precisely map and validate a major QTL for grain weight on chromosome 4AL in bread wheat (Triticum aestivum L.). Theor. Appl. Genet. 2019, 132, 2367–2379. [CrossRef] [PubMed]