Increased lipid production by heterologous expression of AtWRI1 transcription factor in *Nannochloropsis salina*

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**Abstract**

**Background:** Genetic engineering of microalgae is necessary to produce economically feasible strains for biofuel production. Current efforts are focused on the manipulation of individual metabolic genes, but the outcomes are not sufficiently stable and/or efficient for large-scale production of biofuels and other materials. Transcription factors (TFs) are emerging as good alternatives for engineering of microalgae, not only to increase production of biomaterials but to enhance stress tolerance. Here, we investigated an AP2 type TF Wrinkled1 in *Arabidopsis* (AtWRI1) known as a key regulator of lipid biosynthesis in plants, and applied it to industrial microalgae, *Nannochloropsis salina*.

**Results:** We expressed AtWRI1 TF heterologously in *N. salina*, named NsAtWRI1, in an effort to re-enact its key regulatory function of lipid accumulation. Stable integration AtWRI1 was confirmed by RESDA PCR, and its expression was confirmed by Western blotting using the FLAG tag. Characterizations of transformants revealed that the neutral and total lipid contents were greater in NsAtWRI1 transformants than in WT under both normal and stress conditions from day 8. Especially, total lipid contents were 36.5 and 44.7% higher in NsAtWRI1 2–3 than in WT under normal and osmotic stress condition, respectively. FAME contents of NsAtWRI1 2–3 were also increased compared to WT. As a result, FAME yield of NsAtWRI1 2–3 was increased to 768 mg/L/day, which was 64% higher than that of WT under the normal condition. We identified candidates of AtWRI1-regulated genes by searching for the presence of the AW-box in promoter regions, among which lipid metabolic genes were further analyzed by qRT-PCR. Overall, qRT-PCR results on day 1 indicated that AtWRI1 down-regulated *TAGL* and *DAGK*, and up-regulated *PPDK*, *LPL*, *LPGAT1*, and *PDH*, resulting in enhanced lipid production in NsAtWRI1 transformants from early growth phase.

**Conclusion:** AtWRI1 TF regulated several genes involved in lipid synthesis in *N. salina*, resulting in enhancement of neutral lipid and FAME production. These findings suggest that heterologous expression of AtWRI1 TF can be utilized for efficient biofuel production in industrial microalgae.

**Keywords:** Microalgae, *Nannochloropsis salina*, Transcription factor, Wrinkled1, Biofuels, TF engineering

**Background**

The declining availability of petroleum-based fuels and their environmental impact, including global warming, has increased interest in the use of biomass as an alternative energy source [1]. Microalgae are promising feedstocks for biofuel production, with advantages that include high photosynthetic efficiency and biomass production. These organisms can grow rapidly while fast consuming CO₂ and accumulate large amounts of lipids, which are converted to biofuels. In contrast to land plants, microalgae do not require large areas of arable land for cultivation and do not compete with food sources [2, 3]. However, high production costs limit the commercialization of microalgal biofuels, which requires significant genetic improvements for maximum production of lipids [4–6].
Conventional genetic engineering has focused on metabolic engineering of individual enzymes to increase lipid production of microalgae. For example, overexpression of glycerol-3-phosphate acyltransferase (GPAT) and glycerol-3-phosphate dehydrogenase (GPDH) in the diatom Phaeodactylum tricornutum increased lipid contents by 1.6-fold compared with wild type (WT) [7, 8]. Moreover, antisense knockdown of pyruvate dehydrogenase kinase (PDK) increased neutral lipid content in P. tricornutum by 82% [9], and overexpression of diacylglycerol acyltransferase (DGAT) increased lipid content by twofold in Scenedesmus obliquus [10]. However, overexpression of other enzymes such as acetyl-CoA carboxylase (ACCase) and fatty acid synthase (FAS) failed to enhance lipid production in microalgae [11], probably due to difficulties in genetic engineering and/or complicated metabolic feedback mechanisms in microalgae [12].

It is thus necessary to develop alternative strategies to regulate multiple genes in a metabolic pathway. Transcription factor (TF) engineering has been employed in global regulation of gene expression for certain metabolic pathways in plants and other eukaryotes, and is now emerging as a novel approach for improving microalgae for biomaterial production [13, 14]. In an effort to understand regulatory network of lipid metabolism, various TFs affecting triacylglycerol (TAG) accumulation have been identified through multi-omics (transcriptomic, proteomic and metabolomic) studies in Chlamydomonas [15]. For example, nitrogen-responsive regulator (NRR) plays an important role in TAG accumulation under nitrogen starvation condition [16]. Recently, a phosphorus starvation response 1 (PSR1) TF has been identified as a global regulator of carbon storage metabolism. The mechanisms underlying lipid and starch biosynthesis were confirmed by assessing the downstream target genes using PSR1 null and overexpressing mutants [17, 18].

Heterologous expression of plant TFs can be employed effectively in microalgae as TF engineering. TFs have conserved domains and can play similar roles in plants and microalgae [19, 20], resulting in improved regulatory systems in microalgae by heterologous expression of plant TFs with well-characterized functions. For instance, Dof (DNA binding with one finger) TFs containing a C2C2-type zinc finger-like motif play similar roles in different organisms [21–23]. In particular, GmDof from the soybean (Glycine max) is involved in lipid synthesis, and heterologous expression of GmDof has been shown to increase lipid contents in Arabidopsis [24]. Similar effects have also been reported in Chlorella and Chlamydomonas [25, 26].

Wrinkled 1 (WR1) is a master regulator of lipid accumulation in the seed of Arabidopsis. WR1 is a member of APETALA2-ethylene responsive element-binding protein (AP2-EREBP) family, and is controlled by LEAFY COTYLEDON2 (LEC2) [27]. WR1 in turn regulates fatty acid biosynthetic genes with promoters containing an AW-box sequence [CnInG(n)CG], such as PKp-β1, BCCP2, and KAS1, in Arabidopsis [28]. Genetic engineering of WR1 has shown that seed oil production can be increased by overexpression of WR1 homologs in plants, while decreased the null mutants [29, 30]. These results indicate that WR1 has a positive effect on lipid biosynthesis in plants.

Nannochloropsis spp. are considered industrial model strains for biofuels due to their rapid growth and high lipid content. Moreover, Nannochloropsis can produce high value products including eicosapentaenoic acid (EPA), a ω-3 long chain polyunsaturated fatty acid (PUFA), used in health supplements [31]. However, genetic manipulation is required to make Nannochloropsis economically feasible for biofuel production [32–36]. Genome and transcriptome data of Nannochloropsis spp. are available, enabling efficient genetic engineering of these microalgae [37–42]. Moreover, several attempts of genetic modification have been recently found to increase lipid contents in Nannochloropsis [43–45]. Above all, Nannochloropsis can be used for TF engineering: genomic analysis of TFs and TF binding sites (TFBSs) in Nannochloropsis oceanica has been carried out to facilitate prediction of the TFs related to lipid synthesis [46]. In addition, we reported overexpression of a bHLH TF in Nannochloropsis salina that showed enhanced fatty acid methyl esters (FAMEs) productivity [47].

In the present study, we conducted heterologous expression of Arabidopsis thaliana WR1 (AtWR1) in N. salina CCMP1776, and obtained N. salina transformants, named NsAtWR1, with high lipid content. To understand the mechanism by which AtWR1 enhanced lipid synthesis in N. salina, AtWR1-regulated genes involved in lipid synthesis were identified based on the presence of the AW-box, and the mRNA expression levels of selected target genes were confirmed by quantitative real-time polymerase chain reaction (qRT-PCR). These results provided an excellent proof of concept for heterologous expression of key regulatory TF from plants to improve lipid production in industrial microalgae.

**Results**

**Heterologous overexpression of AtWR1 in N. salina**

We constructed an AtWR1 overexpressing vector, pNsAtWR1 (Fig. 1a), and transferred linearized pNsAtWR1 into N. salina CCMP1776 by particle bombardment. We obtained two transformants expressing the AtWR1 protein, named NsAtWR1 2–3 and 1–31. Genomic PCR was performed to check the presence of the plasmid DNA (Fig. 1b). The primers W1 and W2 were
used to detect the *AtWRII* gene, and the primers SR6 and SR9 were used to detect *18S rDNA* (Additional file 1: Table S1). The *18S rDNA* was present in both *N. salina* WT and NsAtWRI1 transformants. In contrast, the *AtWRII* transgene was present in NsAtWRI1 transformants, but absent from WT, indicating that pNsAtWRI1 had been successfully transferred into the transformants. The integration site of transgene into genomic DNA was confirmed by restriction enzyme site-directed amplification (RESDA) PCR and homology search using genomic database of *N. gaditana* B-31 (http://www.Nannochloropsis.org/) (Additional file 2: Figure S1) [41]. The transgene was integrated near the beta-tubulin gene (homologous to Naga_10009g86 in *N. gaditana*) and a hypothetical gene (homologous to Naga_100450g4 in *N. gaditana*) in NsAtWRI1 2–3 and 1–31, respectively. Western blotting confirmed that AtWRI1 protein was expressed in both transformants (Fig. 1c). We used the FLAG tag that was attached to the C-terminus of AtWRI1. A specific band for AtWRI1 protein was present only in NsAtWRI1 transformants, and the F-type H-ATPase β subunit (Atpβ) that was used as a loading control. The expression level of AtWRI1-FLAG protein was higher in 2–3 than 1–31 under all culture conditions (Fig. 1d), suggesting that 2–3 may provide better phenotype than 1–31. Taken together, these results indicated that pNsAtWRI1 had been successfully integrated into the genome and stably expressed in the transformed cells.

We identified three endogenous proteins containing the AP2 domain in *N. salina*, named NsAP2-1, NsAP2-2, and NsAP2-3, and compared with that of *AtWRI1* (Additional file 3: Figure S2). All three showed very low sequence identity based on sequence analyses including BLASTp and CLCbio Main Workbench (Additional file 4: Table S2), suggesting that the endogenous AP2 TFs are not true orthologs, and may not interfere with AtWRI1.

**Growth analysis of NsAtWRI1 transformants**

The phenotype of NsAtWRI1 transformants was analyzed under various culture conditions: normal, N limitation, and osmotic stress. The cell density of NsAtWRI1 2–3 and 1–31 was higher than that of WT under normal and osmotic stress conditions (Fig. 2a, c). Under N limitation condition, the growth pattern and cell density of WT and 2–3 were similar, and 1–31 showed increased cell density (Fig. 2b). The maximum specific growth rate of NsAtWRI1 did not change significantly under normal and osmotic stress conditions. Although the specific growth rate was increased in NsAtWRI1 2–3 than in WT under N limitation, this difference did not affect their overall growth (Fig. 2b; Additional file 5: Table S3). Biomass yield of NsAtWRI1 transformants was similar to that of WT, indicating that AtWRI1 did not affect growth significantly (Additional file 5: Table S3).
Total lipid analysis of NsAtWRI1 transformants

We analyzed total lipid content by employing gravimetric measurements (Fig. 3a, b). Total lipid contents of the transformants were increased on day 8 (Fig. 3a), which were 33 and 31% higher in NsAtWRI1 2–3 and 1–31 than in WT under the normal condition, respectively. NsAtWRI1 transformants also showed moderate increase in total lipids under N limitation and osmotic stress conditions (Fig. 3c, d).
stress conditions. On day 12, transformants showed higher lipid contents under all culture conditions, and 2–3 showed higher increase in total lipids than 1–31 (Fig. 3b). Particularly, NsAtWRI1 2–3 showed 36.5 and 44.7% greater lipid contents compared to WT at day 12 under normal and osmotic stress conditions, respectively. However, under the N limitation condition, lipid contents were not significantly increased in NsAtWRI1 transformants compared with WT. This may be because lipid content was already saturated on day 12, regardless of WT and the transformants. We estimated total lipid yields by multiplying lipid contents and biomass yield as shown in Fig. 3c, d. Lipid yields were basically similar to lipid contents, because transformants did not show significant changes in the biomass. However, maximum increase in lipid yields was found in NsAtWRI1 2–3 showing approximately 70% increase under the normal condition on day 12 (Fig. 3d).

We also measured lipid composition on days 8 and 12 using HPLC that can separate lipids into neutral and polar lipids (Table 1). We identified hydrocarbons and TAG as neutral lipids, which were increased in NsAtWRI1 transformants under all culture conditions, but in general 2–3 was higher than 1–31, possibly reflecting higher expression levels of AtWRI-FLAG in 2–3 under all conditions than 1–31 (Fig. 1c, d). Membrane lipids (MGDG, DGDG, and PI) were also detected, but we did not find any substantial differences between transformants and WT, suggesting that AtWRI1 TF mainly affected accumulation of neutral lipids.

**FAME analysis of NsAtWRI1 transformants**

To evaluate whether NsAtWRI1 transformants can be used practically for biodiesel production, we examined their FAME contents and yields (Fig. 4). The FAME content of NsAtWRI1 transformants and WT was not enhanced on day 8, regardless of culture conditions (Fig. 4a). In contrast, the FAME content of NsAtWRI1 2–3 was 32 and 22% greater than WT on day 12 under normal and osmotic stress conditions, respectively (Fig. 4b). Under the N limitation condition, FAME contents of NsAtWRI1 2–3 did not increase compared to WT, possibly because FAME contents were already saturated on day 8. Although FAME yield, calculated by multiplying biomass yield and FAME content, was similar in all samples on day 8 (Fig. 4c), NsAtWRI1 2–3 was 64% higher than WT under normal condition on day 12 (Fig. 4d). Increased total lipid contents by AtWRI1

| Table 1 Lipid composition (percentage of dry weight biomass, %) in WT and NsAtWRI1 transformants under different culture conditions analyzed by HPLC |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Culture condition      | Day  | Strain | Neutral lipid (%) | Glycolipid (%) | Phospholipid (%) |
|------------------------|------|--------|------------------|----------------|------------------|
|                        |      |        | Hydrocarbon | TAG | Total | MGDG | DGDG | PI |
| Normal                 | Day 8 | WT     | 3.7 ± 2.3 | 1.0 ± 1.7 | 4.7 ± 3.2 | 19.3 ± 2.5 | 1.2 ± 1.4 | n.d. |
|                        | 2–3  | 7.1 ± 1.0* | 48 ± 0.7* | 11.9 ± 1.7* | 21.5 ± 2.0 | n.d. | n.d. |
|                        | 1–31 | 1.8 ± 3.1 | 56.6 ± 0.7** | 7.3 ± 2.4 | 20.4 ± 1.4 | 3.1 ± 0.3 | n.d. |
|                        | Day 12 | WT | 2.8 ± 0.3 | 150 ± 2.3 | 17.8 ± 2.6 | 11.1 ± 1.5 | 3.1 ± 1.8 | 0.5 ± 0.0 |
|                        | 2–3  | 3.3 ± 0.3* | 270 ± 2.9*** | 30.3 ± 2.8*** | 11.1 ± 1.2 | 3.3 ± 1.4 | 0.2 ± 0.2 |
|                        | 1–31 | 3.1 ± 0.2 | 186 ± 1.4* | 21.6 ± 1.4* | 11.0 ± 0.5 | 5.2 ± 0.4 | 0.8 ± 0.1 |
| N limitation           | Day 8 | WT | 5.8 ± 3.7 | 525 ± 6.9 | 583 ± 7.7 | n.d. | n.d. | n.d. |
|                        | 2–3  | 8.7 ± 1.3 | 634 ± 6.8* | 720 ± 7.9* | n.d. | n.d. | n.d. |
|                        | 1–31 | 9.3 ± 0.7 | 583 ± 2.4 | 676 ± 2.8* | n.d. | n.d. | n.d. |
|                        | Day 12 | WT | 6.6 ± 1.3 | 727 ± 3.4 | 793 ± 2.3 | n.d. | n.d. | n.d. |
|                        | 2–3  | 7.1 ± 0.3 | 791 ± 3.1* | 862 ± 3.2* | n.d. | n.d. | n.d. |
|                        | 1–31 | 5.7 ± 3.4 | 738 ± 6.3 | 795 ± 5.3 | n.d. | n.d. | n.d. |
| High salt              | Day 8 | WT | 10.9 ± 10.9 | 268 ± 7.6 | 37.7 ± 3.5 | n.d. | 2.6 ± 4.5 | n.d. |
|                        | 2–3  | 9.3 ± 11.6 | 317 ± 17.9 | 41.0 ± 14.1 | n.d. | 6.7 ± 11.7 | n.d. |
|                        | 1–31 | 20.5 ± 5.5 | 173 ± 4.3 | 378 ± 9.8 | n.d. | 4.9 ± 8.4 | n.d. |
|                        | Day 12 | WT | 5.7 ± 4.9 | 345 ± 4.9 | 402 ± 4.7 | 3.9 ± 3.9 | 4.2 ± 1.4 | 0.2 ± 0.4 |
|                        | 2–3  | 6.7 ± 0.6 | 569 ± 5.7** | 636 ± 6.2** | 4.4 ± 4.4 | 2.1 ± 2.8 | n.d. |
|                        | 1–31 | 6.4 ± 0.9 | 459 ± 11.0 | 522 ± 11.7 | 4.5 ± 4.5 | 3.6 ± 1.5 | 0.2 ± 0.4 |

* Significant differences against WT for the same condition, as determined by Student’s t test, are indicated by asterisks (*P < 0.05, **P < 0.01, ***P < 0.001)

**TAG** triacylglycerol, **MGDG** monogalactosyldiacylglycerol, **DGDG** digalactosyldiacylglycerol, **PI** phosphatidylinositol, n.d. not detected

*a* Total neutral lipid contents were calculated by sum of hydrocarbon and TAG contents
TF appeared to affect FAME production (Figs. 3, 4). Improvement of FAME yield limited to the normal condition may reflect the original functions of AtWRI1 in plants, which will be discussed further. Overall, NsAtWRI1 2–3 behaved better in lipid accumulation than 1–31, and we thus chose NsAtWRI1 2–3 for further analyses.

Identification and molecular analysis of AtWRI1-regulated candidate genes involved in lipid synthesis

To determine genes targeted by AtWRI1, we used matrix-scan provided by regulatory sequence analysis tool (RSAT), and obtained 475 genes with promoters containing an AW-box [48]. These 475 genes were classified by the gene ontology (GO) term enrichment test using Blast2GO software (https://www.blast2go.com/) into three domains: biological processes, cellular components, and molecular function (Additional file 6: Figure S3). Genes containing the AW-box represented a wide variety of functions. We noticed that many metabolic processes were included. Carbon utilization, lipid transport and localization processes were overrepresented compared with N. salina reference GO. These characteristics can be interpreted that many AW-box genes were involved in carbon and lipid metabolism. As lipid content was likely to be altered by AtWRI1 TF, we chose seven N. salina genes that may be involved in TAG and fatty acid synthesis (Table 2). These included genes encoding triacylglycerol lipase (TAGL) and lysophospholipase (LPL) that are involved in organic substance metabolic process; diacylglycerol kinase (DAGK) involved in phospholipid catabolic process; diacylglycerol acyltransferase family protein (DGAT) and lysophosphatidylglycerol acyltransferase 1 (LPGAT1) involved in transferring acyl groups; and pyruvate phosphate dikinase (PPDK) localized in chloroplasts and the dihydrolipoyllysine-residue acetyltransferase component of mitochondrial pyruvate dehydrogenase (PDH), which are involved in pyruvate metabolism. Gene structure and location of the AW-box are summarized in Additional file 7: Figure S4, and the

Fig. 4 FAME quantitation of NsAtWRI1 transformants under various culture conditions. FAME contents on day 8 (a) and day 12 (b). FAME yield on day 8 (c) and day 12 (d). The data points represent the average of samples and error bars indicate standard deviation (n = 4). Significant differences against WT for the same condition, as determined by Student’s t test, are indicated by asterisks (*P < 0.05, **P < 0.01, ***P < 0.001)
### Table 2  *N. salina* lipid synthesis-related genes with promoters containing AW-boxes

| Gene                                      | Abb. | Gene IDa | Protein IDa | GO Names (ID) listb                                     |
|-------------------------------------------|------|----------|-------------|--------------------------------------------------------|
| Triacylglycerol lipase                    | TAGL | Naga_100057g25 | EWM23354.1 | P: organic substance metabolic process (GO:0071704)    |
| Diacylglycerol kinase                     | DAGK | Naga_100089g7 | EWM25228.1 | P: phospholipid catabolic process (GO:0009395); P: organic substance metabolic process (GO:0071704); F: transferase activity (GO:0016740) |
| Acyl transferase/acyl hydrolase/lysophospholipase | LPL  | Naga_100156g4 | EWM20908.1 | P: organic substance metabolic process (GO:0071704)    |
| Acyl-lysophosphatidylglycerol acyltransferase 1 | LPGAT1 | Naga_100501g5 | EWM28984.1 | F: transferase activity; transferring acyl groups (GO:0016746); F: transferase activity (GO:0016740) |
| Diacylglycerol acyltransferase family protein | DGAT | Naga_100028g44 | EWM29372.1 | F: transferase activity; transferring acyl groups other than amino-acyl groups (GO:0016747); F: transferase activity (GO:0016740) |
| Dihydrolipoyllysine-residue acetyltransferase component of pyruvate dehydrogenase mitochondrial-like | PDC   | Naga_100006g108 | EWM29544.1 | P: pyruvate metabolic process (GO:0006090); P: organic substance metabolic process (GO:0071704); F: transferase activity (GO:0016740); C: cytoplasm (GO:0005737); C: organelle (GO:0043226); C: mitochondrial matrix (GO:0005759); C: cytoplasmic part (GO:0044444); C: intracellular organelle (GO:0043229) |
| Pyruvate phosphate dikinase                | PPDFK | Naga_100043g42 | EWM23135.1 | P: carbon utilization (GO:0015976); P: pyruvate metabolic process (GO:0006090); P: organic substance metabolic process (GO:0071704); F: transferase activity (GO:0016740) |

a  As *N. salina* CCMP1776 has not been annotated in NCBI, Gene ID and Protein ID refer to those of *N. gaditana* B-31

b  GO enrichment test was performed using Blast2GO software with Fisher's exact test and *P* value < 0.09
lipid synthesis pathway including roles of the selected AtWR1-regulated genes is presented in Additional file 8: Figure S5.

The mRNA expression levels of the seven lipid-related genes were analyzed by qRT-PCR (Fig. 5). The mRNA was obtained after cells were incubated for 24 h under normal, N limitation, and osmotic stress condition, and the endogenous actin gene was used for normalization. The TAGL expression level was lower in NsAtWR1 2–3 than in WT under normal and osmotic stress conditions. The DAGK expression level of the transformant also decreased under normal and N limitation condition compared with WT. On the other hand, the expression level of LPL was greater in the transformant than in WT under osmotic stress conditions. The LPGAT1 expression level of the transformant was also considerably higher than that of WT under N limitation and osmotic stress condition. The DGAT expression level of the transformant was lower in the transformant than in WT under normal conditions, but higher under osmotic stress conditions. The PDH expression level of the transformant was higher than that of WT under osmotic stress condition, and the PPDK expression level of the transformant was higher compared to that of WT under N limitation and osmotic stress conditions. Overall, these up- and down-regulation expression patterns could be reconciled by their possible contribution to lipid production from early growth phase, which led us to classify the target genes into positive and negative groups, respectively. Significance of the classification will be discussed further.

We also performed an extended qRT-PCR to reveal expression profiles on days 0, 1, and 8 (Additional file 9: Figure S6). Expression of LPL, PDH and PPDK remained high in NsAtWR1 2–3 compared to WT on day 8. Expression of DGAT was increased under the normal condition on day 8, which may have contributed to the increased lipid accumulation on day 12 under the normal condition (Fig. 3b, d). The expression pattern of DAGK was similar on day 8 compared to that of day 1. However, expression patterns of TAGL were changed on day 8, where they showed higher expression in NsAtWR1 2–3 than WT. It is not certain why these genes showed reversed expression pattern on day 8, but it appears that these genes might have contributed less significantly to lipid accumulation on day 12.

**Discussion**

Transcription factors’ engineering has emerged as an excellent strategy for improved productivity of biomass and lipids in plants, and is starting to be employed in microalgae. As manipulation of individual metabolic enzymes in microalgae may not be successful to enhance production of lipids effectively, global regulation of multiple enzyme in lipid metabolic pathways can be emphasized [13]. In an effort to improve industrial microalgae for lipid production employing TF engineering, we chose AtWR1 that is a key regulatory TF of lipid accumulation in Arabidopsis seeds [49]. Effects of this heterologous expression were positive on lipid content and yield, and we looked into revealing mechanisms for the enhancement. We identified its possible target genes by scanning the genome of N. salina for the presence of the AW-box containing promoters. Among the 475 genes, we selected seven genes involved in the lipid metabolism. We divided the seven genes into two groups: positive and negative ones based on their known functions, and their expression patterns were analyzed by qRT-PCR on day 1 (Fig. 5).

The negative group included TAGL and DAGK: TAGL degrades TAGs into diacylglycerol (DAG) and free fatty acids (FFA) [50], and DAGK catalyzes the phosphorylation of DAG leading to the formation of phosphatidic acid (PA) and indirectly decrease of TAG (Additional file 8: Figure S5) [51, 52]. It has been reported that down-regulation of TAGL results in TAG accumulation [50], and DAGK expression is lowered when a large amount of lipid is accumulated in microalgae [51]. NsAtWR1 2–3 showed lower mRNA expression of TAGL under normal and osmotic stress condition, compared to WT. The DAGK expression level of NsAtWR1 2–3 was also lower than that of WT under normal and N limitation. The down-regulation of TAGL and DAGK may contribute to accumulation of lipids under normal and stress conditions (Figs. 3, 5 and Table 1).

We classified PPDK, LPL, LPGAT1, and PDH into the positive group. Chloroplast PPDK plays an important role in the C4-like carbon-concentrating mechanism in addition to its classical role in reverse glycolytic pathway [53, 54]. As Nannochloropsis has C4-like carbon-concentrating mechanisms, high expression of chloroplast PPDK in NsAtWR1 transformants may increase carbon fixation leading to increased lipid content [39]. LPL metabolizes lysophosphatidylcholine (LPC), producing acyl groups that can contribute to TAG accumulation. LPGAT1 converts lysophosphatidylglycerol (LPG) to phosphatidylglycerol (PG), which is incorporated into cellular membranes of microalgae [55]. As fatty acid in membrane phospholipid such as PG can be used for TAG synthesis by remodeling [56], LPGAT1 enzyme can indirectly contribute to increase neutral lipid. In addition, mitochondrial PDH catalyzes the conversion of pyruvate to acetyl-CoA, which can be used for fatty acid and neutral lipid synthesis [9]. In this study, the LPL and PDH expression level was higher in NsAtWR1 than in WT under osmotic stress condition, and the expression level of LPGAT1 was higher in NsAtWR1 2–3 than in WT.
under N limitation and osmotic stress conditions. The increased expression level of \textit{LPL}, \textit{PDH}, and \textit{LPGAT1} was likely to enhance the neutral lipid content (Figs. 3, 5 and Table 1), as summarized in schematic metabolic pathways relevant to these genes (Additional file 8: Figure S5).

The above candidates showed expression patterns consistent with their functional annotations. However, the \textit{DGAT} gene that contained an AW-box did not show consistent expression pattern that varies depending on culture conditions on day 1. Whereas the mRNA expression level of \textit{DGAT} was higher in NsAtWRI1 than in WT under osmotic stress condition, the \textit{DGAT} expression level of NsAtWRI1 was lower than that of WT under normal condition. It is likely that this \textit{DGAT} gene may not contribute to the lipid accumulation observed in NsAtWRI1. Other genes described earlier might be more important for the phenotypes in NsAtWRI1. These expression patterns indicate that NsAtWRI1 induces lipid accumulation by regulating the possible target genes from day 1, resulting in early lipid accumulation from day 8 (Fig. 3).

We also analyzed the expression profiles of the AtWRI1-regulated genes with days 0, 1 and 8. Expression of \textit{LPL}, \textit{PDH}, and \textit{PPDK} on day 8 was increased in the NsAtWRI1 transformant, consistent with that on day 1, which may contribute to the increased lipid accumulation. We found that the expression pattern of \textit{DAGK} was similar on day 8 compared to day 1, and that of \textit{TAGL} was reversed. We are not certain of the significance of this reversal of gene expression for lipid metabolism. We speculate that the lipid accumulation would be combined effects of complex regulation of metabolic genes by the heterologous expression of AtWRI1 and endogenous regulatory mechanisms in \textit{Nannochloropsis} under different culture conditions and times. We showed that the target genes were regulated more predictably early on day 1, and became up-regulated later on day 8 in general, suggesting that the early regulation contributed the lipid accumulation from early growth phase, resulting in increased lipid contents from day 8 (Fig. 3).

Based on these results, we suggest that heterologous expression of AtWRI1 is applicable to industrial production of biofuels. AtWRI1 2–3 expressing more AtWRI1 TF accumulated more neural lipids. Total lipid yield of NsAtWRI1 2–3 was increased by 1.1 mg/L, which is about 70% greater than WT under normal condition (Fig. 3d). Moreover, FAME yield, which can be used for biodiesel, of NsAtWRI1 2–3 was improved under the normal condition, where it reached as high as 768 mg/L, about 64% greater than in WT (Fig. 4d). This improvement may reflect the original function of AtWRI1 in \textit{Arabidopsis}, where it is involved in lipid accumulation in seeds in response to developmental cues [28]. It is important to achieve improved lipid production under
normal conditions for industrial production, because it is costly to add additional process in large-scale cultivation schemes. We showed that AtWRI1 was capable of regulating certain lipid metabolic genes and increasing lipid production in *N. salina*. Taken together, our results provided proof of concept that heterologous expression of plant key regulatory gene can be a strategy to improve industrial microalgae for production of biofuels and other materials.

**Conclusion**

We demonstrated that heterologous expression of AtWRI1 TF could increase lipid contents and yield in *N. salina*. AtWRI1 seems to affect TAG synthesis and total lipid contents in *N. salina*. Total lipid content and yield of NsAtWRI1 started to increase on day 8. FAME yield of NsAtWRI1 2–3 was also 64% higher than that of WT on day 12 under normal condition. We identified seven possible AtWRI1-regulated genes involved in lipid metabolism in *N. salina* based on the presence of the AW-box in the promoter region, and analyzed their expression levels. In general, DAGK and TAGL were down-regulated, and LPL, LPGAT1, PDH, and PPDK were up-regulated, resulting in positive effects on lipid synthesis from early growth stage. This study provides proof of the concept that heterologous expression of AtWRI1 TF can be used for lipid production in industrial microalgae.

**Methods**

**Microalgal strains and culture conditions**

*Nannochloropsis salina* CCMP 1776 (Culture Collection of Marine Phytoplankton; now managed by the National Center for Marine Algae and Microbiota) was maintained in 250 mL Erlenmeyer baffled flasks containing 200 mL modified F2N medium [47], composed of 15 g/L sea salt (Sigma-Aldrich, USA), 10 mM Tris–HCl (pH 7.6), 427.5 mg/L NaNO₃, 30 mg/L NaH₂PO₄·2H₂O, 5 mL/L trace metal mixture (4.36 g/L Na₂EDTA·2H₂O, 3.15 g/L FeCl₃·6H₂O, 10 mg/L CoCl₂·6H₂O, 22 mg/L ZnSO₄·7H₂O, 180 mg/L MnCl₂·4H₂O, 9.8 mg/L CuSO₄·5H₂O, 6.3 mg/L Na₂MoO₄·2H₂O), and 2.5 mL/L vitamin stock (1 mg/L vitamin B₁₂, 1 mg/L biotin, 200 mg/L thiamine-HCl). The cells were incubated at 25 °C with agitation at 120 rpm under 120 µmol photons/m²/s of fluorescent light, while being supplied directly with air containing 2% CO₂ at 0.5vvm (volume gas per volume medium per minute). Normal medium in cultivation experiments was identical to modified F2N medium; in N limitation medium, the NaNO₃ concentration was 75 mg/L, whereas in osmotic stress medium, the sea salt concentration was 50 g/L.

**Vector construction**

The AtWRI1 gene from *Arabidopsis* was cloned into a plasmid, named pNsAtWRI1 (Fig. 1a). To optimize AtWRI1 expression in *N. salina*, codons were optimized based on the codon frequency of *Nannochloropsis* [57] and adjustment of GC contents below 60%. The sequence was synthesized by BIONEER Co. (South Korea). The coding sequence of AtWRI1 was PCR amplified using the W1 (forward) and W2 (reverse) primers (Additional file 1: Table S1), and the pNsAtWRI1 backbone was amplified using the W3 (forward) and W4 (reverse) primers. These two PCR products were combined by the Gibson assembly technique [58]. In the plasmid pNsAtWRI1, the codon-optimized AtWRI1 gene was expressed under the control of the endogenous TUB promoter and terminator. The Shble gene was used for selection, which confers resistance to Zeocin, expressed under the control of the UEP promoter and terminator (Fig. 1a) [47].

**Particle bombardment transformation**

The pNsAtWRI1 plasmid was linearized by digestion with NcoI and concentrated to 1 µg/µL for particle bombardment. The linearized plasmid was coated onto 0.6 µm microcarrier gold particles (Bio-Rad, USA) by mixing 3 mg of gold particles in 50 µL of 50% glycerol, 5 µL of the DNA, 50 µL of 2.5 M CaCl₂, and 20 µL of 0.1 M spermidine. The resulting DNA-coated particles were suspended in 60 µL of 100% ethanol, and 12 µL was used for each bombardment transformation [47].

*Nannochloropsis salina* WT cells cultivated under normal condition were harvested in mid-exponential phase. The cells were concentrated, and 10⁸ cells on cellulose acetate membrane filters (Sartorius Stedim Biotech, Germany) were placed on modified F2N agar medium. Particle bombardment transformation was performed using a GDS-80 low-pressure gene delivery system (Wealtec, USA), with 625 µg of coated gold particles being shot in 700 psi helium, and with a 3 cm target distance between the muzzle of the gene gun and the cells. After transformation, the cells were allowed to recover by incubation for 1 day in modified F2N broth medium at 25 °C with dim light (5 µmol photons/m²/s of fluorescent light). The cells were harvested, and plated onto agar plates of modified F2N agar containing 2.5 µg/mL Zeocin [47].

**Molecular analysis of NsAtWRI1 transformants by genomic PCR, RESDATA PCR, and Western blotting**

Transfer of the pNsAtWRI1 vector into *N. salina* was confirmed by genomic PCR. The genomic DNA of the cells was extracted using 200 µL of Instagene Matrix (Bio-Rad, USA) at 56 °C for 20 min and at 100 °C for
8 min. The mixture was centrifuged, and the supernatant was used as the template for genomic PCR. Genomic PCR was carried out using the W1 (forward) and W2 (reverse) primers to detect AtWRI1 and the SR6 (forward) and SR9 (reverse) primers to detect 18S rDNA. The PCR amplification was conducted under the following conditions; 95 °C for 5 min, 30 cycles of 95 °C for 1 min, 60 °C for 1 min, 72 °C for 1 min, and then 72 °C for 10 min. The PCR-amplified bands of AtWRI1 and 18S rDNA PCR were detected at 1.3 kb and 380 bp, respectively.

RESDA PCR was performed to determine if the transgene had integrated into genomic DNA [60]. RESDA PCR consists of two steps, amplification I (Amp I) and amplification II (Amp II). The Amp I was performed using RESDA fwd 1 primer and three degenerate primers (DegClal, DegNdel, DegSspl) (Additional file 1: Table S1) under the following conditions; 96 °C for 5 min, 20 cycles of 95 °C for 1 min, 60 °C for 1 min, 72 °C for 3 min, 10 cycles of 95 °C for 1 min, 40 °C for 1 min, and then 72°C for 10 min. The Amp II was performed using the RESDA fwd 2 and Q0 primers, and, as a template, 1 µL of the PCR product from the first amplification (Additional file 1: Table S1). The amplification was performed under the following conditions; 96 °C for 5 min, 30 cycles of 95 °C for 1 min, 60 °C for 1 min, 72 °C for 3 min, and then, 72 °C for 10 min. The PCR product from Amp II step was purified and sequenced (Additional file 2: Figure S1).

Western blotting was performed to confirm expression of FLAG-tagged AtWRI1. Protein was extracted from 3 × 10^8 cells by vigorous mixing with 1.5× modified Laemmli sample buffer [62.5 mM Tris–HCl, pH 7.6, 7% sodium dodecyl sulfate (SDS), 25% glycerol, 5% β-mercaptoethanol, and 0.02% bromophenol blue], followed by incubation at 100 °C for 5 min [47]. The samples were centrifuged, and 10 µL of each supernatant was electrophoresed in SDS-polyacrylamide gels (PAGE) (Any kD Mini-PROTEAN TGX Precast Protein Gels, Bio-Rad, USA), followed by transfer to polyvinylidene difluoride (PVDF) membranes using the Trans-Blot Turbo system (Bio-Rad, USA). The membranes were blocked by incubation with 5% skim milk and 0.1% Tween 20 in phosphate-buffered saline (PBS) and immunoblotted by incubation in a 1:1000 dilution of rabbit anti-FLAG-tag antibody (Cell Signaling Technology, USA) and a 1:5000 dilution of rabbit anti-AtpB antibody (Agrisera, Sweden) for loading control. After washing, the membranes were incubated with a 1:1000 dilution of horseradish peroxidase (HRP)-conjugated anti-rabbit secondary antibody (Cell Signaling Technology, USA). FLAG-tagged AtWRI1 was detected by enhanced chemiluminescence (ECL) reagents and the ChemiDoc system (Bio-Rad).

**Growth analysis of NsAtWRI1 transformants**

WT and NsAtWRI1 were cultivated under normal, N limitation, and osmotic stress conditions. Cell growth was analyzed by measuring cell density, specific growth rate, and dry cell weight (DCW). Cell density was estimated by the Cellometer Auto X4 Cell Counter (Nexcelom Bioscience, USA). Specific growth rate was calculated from cell density using the equation:

\[
\text{Specific growth rate (µ/day)} = \ln(X_2/X_1)/(t_2 - t_1),
\]

where \(X_1\) and \(X_2\) are the initial and final cell concentrations and \(t_1\) and \(t_2\) are the first and last days of culture. To determine DCW, cells were filtered with GF/C filter paper (Whatman, USA), washed with deionized water and dried at 105 °C overnight, after which cell weight was measured.

**Total lipid analysis by gravimetric method and HPLC**

Folch's method was used to extract total lipids [61]. Briefly, microalgal cells were harvested, and then lyophilized in a freeze drier (FD5508, IIShin BioiBase, South Korea). A chloroform–methanol mixture (2:1, v/v) was added to the lyophilized biomass and then sonicated for 1 h at room temperature. Deionized water was added to the samples and vortexed vigorously for 5 min. After centrifugation at 3500 rpm for 15 min, the separated lower layer (organic phase) was obtained and filtered using 0.20 µm RC-membrane syringe filters (Sartorius Stedim Biotech, Germany). Total lipid contents were calculated by measuring difference of lipid mass after evaporation of the chloroform. Lipid composition and their contents were determined by High Performance Liquid Chromatography (HPLC) Agilent 1260 and ELSD (Agilent technologies, USA) with a Chromolith®Performance-Si ((100 × 4.6 mm I.D.) column (VWR, USA). A gradient system was modified from a previous report [62].

**Fatty acid methyl ester (FAME) analysis**

Ten mg of lyophilized biomass was vigorously mixed with a chloroform–methanol mixture (2:1, v/v) for lipid extraction. Then, 0.5 mg of heptadecanoic acid (C17:0) was added as an internal standard. For transesterification, 1 mL of methanol and 300 µL of sulfuric acid were added to the samples, and incubated at 100 °C for 20 min. Each sample was then mixed with 1 mL of deionized water, followed by centrifugation to separate the organic (lower) phase, which was obtained by filtration using 0.20 µm RC-membrane syringe filters (Sartorius Stedim Biotech, Germany). FAMES were analyzed using a gas chromatograph (GC) (HP 6890, Agilent, USA) equipped with a flame ionized detector (FID) and an HP-INNOWax polyethylene glycol column (HP 19091N-213, Agilent, USA). The oven temperature of the GC was increased from
50 to 250 °C at a rate of 15 °C per min. The FAME contents in these samples were determined by reference to a 37-component mix of FAME standards (F.A.M.E. MIX C8-C24, Supelco, USA).

Identification of AtWRI1 targets and analysis of RNA expression by qRT-PCR
To select target genes containing AW-boxes [CnTnG(n)7CG], where ‘n’ represents any nucleotide [28], sequence patterns were predicted with RSAT ‘matrix-scan’ [48]. A total of 7494 sequence patterns that included AW-boxes were detected in the N. salina genome. We then obtained 7435 predicted promoter sequences located 500 bp upstream from the start codon of individual genes. Through this process, 475 genes containing AW-boxes in their promoter sequences were identified. These 475 genes could be classified by GO term enrichment test using Blast2GO software with Fisher’s exact test and \( P \) value < 0.05 (Additional file 6: Figure S3) [63]. The seven genes related to lipid metabolism (PPDK, LPGAT1, PDH, LPL, DGAT, TAGL, and DAGK) were selected for further experiments. To understand specific function of the seven genes, we repeated enrichment test using Blast2GO software with Fisher’s exact test and \( P \) value < 0.09 (Table 2; Additional file 7: Figure S4).

The expression levels of these selected target genes were analyzed by qRT-PCR in WT and NsAtWRI1. Cells were incubated for 1 and 8 days under normal, N limitation and osmotic stress conditions. The cells were harvested and RNA was extracted using NucleoZol reagent (Macherey–Nagel, Germany), according to the manufacturer’s protocol. DNA was removed from these RNA samples using DNA-free™ DNase kits (Ambion, USA), and the RNAs were reverse transcribed to cDNA using Superscript™ III Reverse Transcriptase (Invitrogen, USA) and an oligo (dT) 20 primer (Invitrogen, USA) [47].

qRT-PCR was performed using the CFX96 Real-Time system (Bio-Rad, USA). Used primers for the seven selected target genes and actin gene, used as a loading control for normalization, are summarized in Additional file 1: Table S1. Each 20 μL reaction volume contained 2 μL of cDNA (representing 20 ng of total RNA), 0.5 μL of each forward and reverse primer (at concentrations 2 μM), 7 μL of distilled water, and 10 μL of Universal SYBR Supermix (Bio-Rad, USA). The qRT-PCR was performed under the following conditions; 95 °C for 2 min; 40 cycles of denaturation at 95 °C for 10 s, annealing at 60 °C for 10 s, and extension at 72 °C for 20 s, followed by denaturation at 95 °C for 10 s and a final melting step at 65–95 °C. Gene expression level was analyzed by the \( 2^{-\Delta\Delta CT} \) method, and statistical significance was assessed by Student’s t test [47].
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Competing interests
The authors declare that they have no competing interests.

Availability of data and materials
All data generated or analyzed during this study are included in this published article.

Consent for publication
Not applicable.

Ethics approval and consent to participate
Not applicable.

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