Structural Analysis of the Wheat Genes Encoding NADH-Dependent Glutamine-2-oxoglutarate Amidotransferases and Correlation with Grain Protein Content

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Abstract

Background: Nitrogen uptake and the efficient absorption and metabolism of nitrogen are essential elements in attempts to breed improved cereal cultivars for grain or silage production. One of the enzymes related to nitrogen metabolism is glutamine-2-oxoglutarate amidotransferase (GOGAT). Together with glutamine synthetase (GS), GOGAT maintains the flow of nitrogen from NH₄⁺ into glutamine and glutamate, which are then used for several aminotransferase reactions during amino acid synthesis.

Results: The aim of the present work was to identify and analyse the structure of wheat NADH-GOGAT genomic sequences, and study the expression in two durum wheat cultivars characterized by low and high kernel protein content. The genomic sequences of the three homoeologous A, B and D NADH-GOGAT genes were obtained for hexaploid Triticum aestivum and the tetraploid A and B genes of Triticum turgidum ssp. durum. Analysis of the gene sequences indicates that all wheat NADH-GOGAT genes are composed of 22 exons and 21 introns. The three hexaploid wheat homoeologous genes have high conservation of sequence except intron 13 which shows differences in both length and sequence. A comparative analysis of sequences among di- and mono-cotyledonous plants shows both regions of high conservation and of divergence. qRT-PCR performed with the two durum wheat cvs Svevo and Ciccio (characterized by high and low protein content, respectively) indicates different expression levels of the two NADH-GOGAT-3A and NADH-GOGAT-3B genes.

Conclusion: The three hexaploid wheat homoeologous NADH-GOGAT gene sequences are highly conserved – consistent with the key metabolic role of this gene. However, the dicot and monocot amino acid sequences show distinctive patterns, particularly in the transit peptide, the exon 16–17 junction, and the C-terminus. The lack of conservation in the transit peptide may indicate subcellular differences between the two plant divisions - while the sequence conservation within enzyme functional domains remains high. Higher expression levels of NADH-GOGAT are associated with higher grain protein content in two durum wheats.

Introduction

Nitrogen uptake is an essential element in crop improvement, either directly for grain protein content or indirectly for photosynthetic production. Thus, nitrogen utilization is fundamental to crop productivity. Over the past 50 years nitrogen (N) fertilizers have been extensively used to increase both grain yield (GY) and grain protein content (GPC) in cereals - helping to support a vastly increased world population. Despite that GY and GPC are genetically negatively correlated, it has been shown in various cereals (including wheat) that this correlation can be broken down by adequate nitrogen supply in late plant development [1,2]. However, this requires that growers must optimize the use of nitrogen fertilizers to avoid pollution, while maintaining reasonable profit margins. Improved crops should make better use of nitrogen fertilizer supplies giving higher yields with improved protein contents. Therefore, selecting new crop varieties exhibiting improved nitrogen use efficiency (NUE; the yield of grain per unit of available nitrogen in the soil), and adapting agricultural practices to reduce the use of nitrogen fertilizers represent challenges for both breeders and farmers [3].

Whether nitrogen is derived from soil reserves, from nitrogen fertilizer, or from N₂ fixation, it is incorporated into the organic form via the assimilation of ammonia. However, the primary assimilation of ammonia from external inorganic nitrogen is only part of the process. Nitrogen is also released from plant metabolism as ammonia and reassimilated many times during
the movement of nitrogen around the plant, from seed reserve, through transport to vegetative organs, to eventual redepotition in a new crop of seeds. There is also a major release and reassimilation of nitrogen during the process of photorespiration in C3 plants. The process of ammonia assimilation is thus of crucial importance to crop growth and productivity.

Two major enzymes are responsible for cyclic assimilation of ammonium into amino acids in the biochemical pathway of NH₃₄- ammonia; i.e., glutamine synthetase (GS) and glutamine-2-oxoglutarate amidotransferase (GOGAT). These two enzymes are involved in assimilation and recycling of mineral nitrogen catalyzing ATP-dependent conversion of glutamine into glutamate using ammonia as substrate [4,5]. Glutamine synthetase exists in multiple enzyme forms, the chloroplastic isozyme being encoded by one gene (GS2) and the cytosolic form organized in sub-family of genes. Studies have shown that both chloroplastic and cytosolic GS isozymes are developmentally regulated in leaves [6,7,8,9] and have different metabolic roles. The GOGAT enzyme catalyzes the reductive transfer of the amide group of glutamine to 2-oxoglutarate to form two glutamate molecules [1]. Together with GS, it maintains the flow of nitrogen from NH₃₄ into glutamine and glutamate, which are then used for several other amino transferase reactions during the synthesis of amino acids [5]. Kinetic and inhibitory studies have suggested that GOGAT is the rate-limiting step in amino acid production [10,11]. In rice, two different GOGAT enzymes have been identified based on the electron donor; i.e., a ferredoxin (Fd)-dependent GOGAT and a NADH-dependent GOGAT. Also in rice, NADH-GOGAT is active in developing organs, such as unexpanded non-green leaves and developing grains [12]. NADH-GOGAT has been proposed to be involved in the use of remobilized nitrogen, because it is located in the specific cell types which are important for solute transport from the phloem and xylem elements [13].

A recent report showed, through cereal genome paleo-history studies, that a NUE QTL has been conserved at the same orthologous loci as the GOGAT gene on wheat chromosome 3B, rice chromosome 1, sorghum chromosome 3 and maize chromosomes 3 and 8, despite 50–70 million years of separate evolution associated with considerable sequence shuffling (50 rearrangement events within the NUE locus region) [14]. For these reasons, NADH-GOGAT is one of the potential candidate genes involved in the control of the complex character trait GPC.

The aims of the present work were the identification of wheat NADH-GOGAT A, B and D genomic sequences, the determination of their structure, a comparison to GOGAT sequences to other plants, and the study of NADH-GOGAT expression in two durum wheat cvs characterized by low and high GPC.

Results
Assembly of Hexaploid and Tetraploid NADH-GOGAT Gene Sequences

A previously reported rice genomic sequence of NADH-GOGATI [15] was used to extract 487 454-sequences of hexaploid wheat (A-, B-, and D-genomes) cv Chinese Spring (cerealsdb.uk.net/search_reads.htm) and 82 454-reads of the D-genome from the diploid Triticum tauschii (avena.pw.usda.gov/RHmapping/blast/). These two groups of 454 reads were separately assembled. The Chinese Spring assembly produced a single contig comprising of the three distinct sequences. By comparing this contig with the reported chromosome 3B genomic sequence containing a NADH-GOGAT gene (FN564429) and the sequence obtained by assembling D-genome 454-sequences, we were able to separate the 487 Chinese Spring 454-reads into three sequences assignable to each of the three genomes: 178 reads to the A-genome, 162 to the B-genome, and 147 to the D-genome (the A-genome assignment was by elimination). The three hexaploid wheat homoeologous NADH-GOGAT gene sequences are given in File S1.

Analysis of the sequences finds that the wheat NADH-GOGAT gene is comprised of 22 exons and 21 introns for all the three homoeologous genes (Figure 1). The previously reported sequence for rice showed one more 5' exon entirely within the 5' UTR and is not translated into protein. The rest of the rice gene shows a very high similarity in exon/intron length and organization with the three genes derived for wheat. The intron borders matched the canonical plant intron borders (GT...AG) for all 21 introns. Consensus exon/intron boundaries were determined using grass ESTs sequences aligned with the genomic sequence. When there were no overlapping wheat EST sequences, as happened for the first 5 kb at the 5' end of the genomic sequence, we used other grasses EST sequences to determine the boundaries. In regions with few or no wheat ESTs, exon/intron boundaries matched rice and maize sequences in all cases.

Although the three genes show the same exon/intron number and size, an exception is intron 13, which shows major differences in both length and sequence for all of three homoeologues (Figure S1).

Since there is neither information about wheat NADH-GOGAT DNA coding nor amino acidic sequences, the precise the 5' UTR start is uncertain. However, from the starting codon ATG to the stop codon TAG the three genes show a similar length: the NADH-GOGAT-3B gene is the longest at 10,151 bp, the NADH-GOGAT-3A is 9,996 bp long, and the gene located on chromosome 3D is the shortest at 9,985 bp (Figure 1).

The genomic sequence and structure of the 3A and 3B NADH-GOGAT genes were also obtained for the two durum wheat cultivars Svevo and Ciccio using primers designed from the Chinese Spring sequences. No polymorphisms were found for either gene between the two durum varieties. The only differences between the hexaploid and tetraploid wheats are few single nucleotide polymorphisms and single nucleotide indels, all located in introns. The identical tetraploid Svevo and Ciccio DNA sequences are available at NCBI under accessions KC960545 for the 3A gene and KC960544 for the 3B gene.

Comparison of the Three Hexaploid NADH GOGAT Homoeologous Genes

The three hexaploid NADH-GOGAT genes are highly conserved as seen in the dot plot example of the entire A-genome GOGAT vs the B-genome GOGAT genes using the criterion of an 80% match over a 20 base window (Figure 2A). The conservation extends extensively into the introns as seen by increasing the match criterion to 100% over a 20 base window (Figure S2). Only intron 13 shows both major sequence and length differences. The other pairwise comparisons (A vs B, B vs D) show the same pattern (not shown).

Focusing on exon sequences only, the alignments between two homoeologues at a time showed a total of 84 SNPs spread all over the 22 exons in the NADH-GOGAT-3A and NADH-GOGAT-3B comparison. Fewer SNPs are found in the NADH-GOGAT-3A and NADH-GOGAT-3D comparison (71 SNPs) or the NADH-GOGAT-3B and NADH-GOGAT-3D comparison (67 SNPs). The same situation is found when only introns sequences are compared: NADH-GOGAT-3A and NADH-GOGAT-3B showed a total of 73 SNPs spread all over the 20 introns (intron 13 excepted), with NADH-GOGAT-3A and NADH-GOGAT-3D introns different in the same regions for 67 SNPs, NADH-GOGAT-3B and NADH-GOGAT-3D introns different by only 48 SNPs. In addition, the
three pairwise comparisons find intron sequence differences to include 17 short indels (Figure S1). Intron 13 is the longest intron at 706, 831, and 688 bases for the A, B, and D genomes, respectively - with the central approximately 300 bp of the three sequences differing among the three homoeologues (Figure S1).

A high conservation is also seen in the derived amino acid sequences as shown aligned in Figure 3. There are 41 total sequence differences among the three homoeologues -39 residue changes and two fewer amino acids. The latter occurs in the transit peptide portion of NADH-GOGAT-3B and near the C-terminal end of NADH-GOGAT-3D. Of the 39 single residue changes in the 3-way comparison of Fig. 3, 22 occur in the B-genome sequence compared to 7 and 10 differences in the A- and D-genome proteins, respectively.

Comparison of Amino Acid Sequences from Other Species

The degree of conservation of amino acid sequences was shown among a selection of plant genomic sequences. The coding sequence from the wheat NADH-GOGAT-3B gene was used to derive the complete 3B amino acid sequence, and a similar analysis was carried out for available sequences from Brachypodium, grape, poplar, soybean, Arabidopsis and rice. The seven derived amino acid sequences are aligned in Figure 4. The wheat exon-encoding boundaries are indicated by vertical red lines. There is an additional 5' exon in rice and grape that is entirely 5' UTR sequence; e.g., exon 2 in wheat would be exon 3 in rice and grape.

Highlighted in yellow are conserved amino acid positions on all analyzed plants and positions unique to monocots are highlighted in blue. Exon encoded positions are conserved among all plants analyzed with the previously mentioned exception for the first exon and the exon 16-17 junction point in comparing dicot and monocot plants. Although much of the sequence is conserved among all plants, there is a distinctive pattern of sequence differences between the dicots and monocots as seen in the blue highlighting of Figure 4. In addition, several sequence regions are characterized by both residue differences and sequence length variation. The major examples of differences between monocots and dicots include the junction of exons 16 and 17, and the beginning and ends of the polypeptides.

The only other notable difference in the wheat sequences are deletions of five and eight amino acid residues at positions 840 and 2205, respectively, in Figure 4. The latter deletion was reconfirmed by rechecking the 454 reads and with wheat ESTs, and the former confirmed only with the 454 reads as there are no Triticeae ESTs in that region. Both sequence indels occur within non-conserved regions.

The conservation of DNA sequence extends into the intron sequences, but this conservation decays with evolutionary distance as seen in Figure 2B where the diagonals of four dot blots are arrayed horizontally for comparison. The diagonals compare the gene sequences (from start to stop codons) of the 3A vs 3B genes shown in Figure 2A and the wheat 3A gene vs NADH-GOGAT genes of Brachypodium, rice, and Arabidopsis (Figure 2B). The Brachypodium result shows the similarity to wheat extends extensively into the introns, whereas the matching sequence in rice is largely exonic with limited intron similarity. By the evolutionary distance represented by Arabidopsis, the sequence similarity is restricted to exons and short sequences at the intron ends.

Below the alignment of the NADH-GOGAT enzymes in Figure 4 are noted the four known functional domains – indicated by names, lines, and green vertical lines at the domain ends [16].

Chromosome Mapping of the NADH-GOGAT Genes

Since the genes in the two durum cultivars (Ciccio and Svevo) had no SNPs, the Ciccio/Svevo mapping population would not be used to map the chromosome positions. However, the positions in the hexaploid cv Chinese Spring were mapped using genetic stocks including nulli-tetrasomic, di-telosomic and a set of wheat deletions bin lines (described in Material and Methods).

NADH-GOGAT-3A and NADH-GOGAT-3B genes were physical located to the 3A and 3B chromosomes, respectively, using nulli-tetrasomic lines of cv Chinese Spring (Figure 5) – then further localized to the long arms of those chromosomes (using

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**Figure 1.** Exon/intron organization of wheat NADH-GOGAT genes. A. A genome NADH-GOGAT, B. B genome NADH-GOGAT and C. D genome NADH-GOGAT. Exons are indicated by numbered boxes and introns by intervening lines. Start and stop codon positions of the full-length coding regions are marked.

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ditelocentric 3AS lines missing the long arms) and then to bins 3AL-0.42-0.78 and C-3BL2-0.50. Only the 3A results are shown in Figure 5 for the arm and bin mappings.

**Expression Profile of NADH-GOGAT Genes in Wheat**

General patterns of NADH-GOGAT gene expression were determined by two approaches. The first approach was to estimate if there was differential expression in hexaploid wheat among the three homoeologous genes. To determine this, 67 wheat ESTs encoding wheat NADH-GOGAT sequences were identified at NCBI. When matched to the three wheat homoeologous NADH-GOGAT sequences, 23 match to the 3A gene, 19 to 3B, and 15 to 3D. A Chi-square test for random distribution of the ESTs results in $p = 0.43$, indicating the distribution provides no support for preferential distribution and therefore no support for differential gene expression.

The second approach to examine expression was to determine NADH-GOGAT-3A and NADH-GOGAT-3B genes expression levels of the two durum wheat cvs Svevo and Ciccio from roots (collected at seedling stage) and leaves at three different phenotypic stages (first leaf, flowering and grain filling). Total RNA was extracted from tissues of plants grown in field conditions and

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**Figure 2. GOGAT sequence comparisons.** NADH-GOGAT DNA sequences from start to stop codons, including both exons and introns are compared to show relative conservation of sequences. A. Sequences of THE NADH-GOGAT genes from the A- and B-genomes. Match criterion of 80% match over a 20 residue window. B. DNA sequences from start to stop codons were compared between indicated plants using a criterion on 80% match over a 20 base window. Dot blot diagonals are arrayed horizontally for comparison. In both frames the color-coded heat map represents a composite of the degree and length of sequence similarity. doi:10.1371/journal.pone.0073751.g002
reverse-transcribed for qRT-PCR analyses. To test if the homoeologous genes show differential expression patterns, qRT-PCR were performed using specific primers designed to preferentially amplify the A and B sequences (Figure 6). In roots, significant differences (P<0.01) were observed in the expression of both homoeologous tetraploid genes (NADH-GOGAT-3A and NADH-GOGAT-3B) between the two cultivars Ciccio and Svevo, characterized by different grain protein content values (about 14% and 15%, respectively [17]). Roots of cv Svevo, the durum cultivar with the higher final kernel protein content [17], showed the highest level of both genes transcript (Figure 7A). In particular, the relative ratio of NADH-GOGAT-3A and NADH-GOGAT-3B transcript abundances were both 1.50 in cv Svevo, while the transcript abundances were 0.75 and 0.80 in cv Ciccio, for A and B homoeologous, respectively.

In leaves, transcript levels in the first leaf and grain filling stages showed similar expression levels, while a significantly higher value of transcripts was observed during flowering (P<0.01). NADH-GOGAT-3A and NADH-GOGAT-3B relative mRNA abundances at flowering were 1.78 and 1.00, respectively, in cv Ciccio, and 1.71 and 1.91, respectively, in cv Svevo.

A similar trend was observed for both the homoeologous genes and in both cultivars (Figure 7B–C). Statistical analysis of all data with ANOVA did not show significant differences among biological and technical replicas demonstrating a stability of genes expression even in field conditions. In cv Svevo, a durum cultivar with the higher final kernel protein content [17], showed the highest level of both genes transcript (Figure 7A). In particular, the relative ratio of NADH-GOGAT-3A and NADH-GOGAT-3B transcript abundances were both 1.50 in cv Svevo, while the transcript abundances were 0.75 and 0.80 in cv Ciccio, for A and B homoeologous, respectively.

Discussion

The current report describes the determination of the wheat NADH-GOGAT gene sequences in hexaploid and tetraploid wheat. This gene is one of the candidate genes involved in the control of the complex character grain protein content (GPC). The recent and constant increase in partial or complete genome sequence data opens up interesting perspectives for the use of such large data resources. Through comparative genomics and using new generation sequencing data, it will be easier to obtain information about metabolisms and/or genes of interest in very large and high repetitive genomes such as wheat. Positional cloning of genes from the numerous sequenced genomes in a so-called ‘cross-genome map-based cloning’ is also a useful tool to gain information in lesser known genomes. In this study, we used the single reported sequence for rice [15] to extract from public websites all Roche 454 sequences of wheat cv Chinese Spring (http://www.cerealsdb.uk.net/search_reads.htm) and 454 reads of Triticum tauschii accession (http://avena.pw.usda.gov/RHmapping/blast/), the diploid ancestor of the D-genome of hexaploid wheat (ABD). The latter data was generated in our laboratory. We were able to assemble the three complete orthologous genes for the three hexaploid wheat genomes and a partial sequence from T. tauschii. PCR primer pairs were designed and used to identify genome assignments using DNA from Chinese Spring nulli-tetrasomic, ditelosomic, and bin-deletion lines - resulting in assigning the NADH-GOGAT genes to the group 3 chromosome short arm. The three homoeologous genes have the same intron/exon structure with similar SNP differences in both introns and exons. The conservation in sequence is especially seen in the amino acid sequences, which even when compared to other plants show regions of high conservation – presumably related to regions of critical functionality essential for plant metabolism.

Interestingly, there is major conservation even with intron sequences compared to more divergence of introns in another nitrogen-metabolism related gene, lysine ketoglutarate reductase saccharopine dehydrogenase (LKR-SDH) [18]. Both genes contain a large centrally located intron with the least sequence conservation. In the LKR-SDH gene, this region is referred to as a ‘hinge’ region separating the two major functional domains of the enzyme, LKR and SDH. Neither is known the exact role of this region, nor if it is also involved in separating functional domains in the NADH-GOGAT enzymes.

The conservation among NADH-GOGAT sequences can also be seen when comparing amino acid sequences. The coding sequence from the wheat NADH-GOGAT genes (encoded sequences indicated in Figure 4) were used to derive the complete amino acid sequences and compared with model and crop species such as Brachypodium, grape, poplar, soybean, Arabidopsis and rice. Despite the presence of residue differences among species, what is clear is that some regions of amino acid sequence are highly conserved among the species examined, most of all in the regions corresponding to the enzyme functional domains. Five different protein domain are present in NADH-GOGAT, which are in order: 1) a glutamine amidotransferases class-II (376 aa); 2) a glutamate synthase central domain (288 aa) which connects the amino terminal amidotransferase domain with the FMN-binding domain and has an alpha/beta overall topology; 3) a conserved region containing a putative FMN binding site and Fe-S cluster (369 aa); 4) a GXGXG motif, found only in glutamate synthase and few other enzymes, and characterized by a repeated G-XX-G-XXX-G motif (192 aa); 5) and a pyridine nucleotide-disulphide oxidoreductase domain, which is actually a small NADH binding domain within a larger FAD binding domain (314 aa) [16]. Length and position of all of these domains are conserved in all the species examined, as shown in Figure 4. The wheat NADH-GOGAT sequence shows a higher similarity with the Brachypodium gene sequence in pairwise alignments and phylogenetic trees generated from multiple species alignments (not shown). This is consistent with the finding of the close sequence relationship between Brachypodium and the Triticeae species [19].

The comparison of this set of plant NADH-GOGAT genes (wheat, Brachypodium, rice, sorghum, poplar, Arabidopsis, and grape) suggests regions of greater sequence and structure conservation likely related to critical enzymatic functions and metabolic control. The three main differences in amino acid sequence among the aligned proteins in Figure 4 are the N- and C-termini, and the junction between sequences encoded by exons 16 and 17. The N-termini of NADH-GOGAT proteins is reported to encode an approximately 100 bp presequence/transit/targeting peptide in both rice [15] and bean [17]. In contrast, the alignment of Figure 4 shows that whatever the exact structure of this ‘presequence,’ there must be an approximately 25 amino acid residue difference between the aligned dicot and monocot polypeptides. In addition, the sequences in this region are also very different between

Figure 3. Comparison of three hexaploid NADH-GOGAT amino acid sequences. The three cv Chinese Spring NADH-GOGAT polypeptide sequences are aligned. Genomes are indicated on the right by letter of the genome (A, B, D). Differences in amino acid residues are indicated by yellow shading. doi:10.1371/journal.pone.0073751.g003
monocots and dicots. It is not yet determined if these variations in length and sequence have any different functional metabolic roles in the two plant branches. Similar uncertainty exists as to whether the dicot/monocot differences in NADH-GOGAT structure at the C-terminus and junction of exon 16 and 17 encoded amino acid sequences have functional significance or are more random drift of relatively non-functional sequences since no role in metabolism is yet assigned.

Together with GS, GOGAT maintains the flow of nitrogen from NH₄⁺ into glutamine and glutamate, which are then used for several other aminotransferase reactions during the synthesis of amino acids [5]. Kinetic and inhibitory studies have suggested that NADH-GOGAT is the rate-limiting step in amino acid production [10,11].

A recent cross-genome ortho-meta QTL study of NUE in cereals identified a GOGAT gene as a major candidate for cereal NUE [14]. In rice, the suppression of both GOGAT genes reduced yield per plant and thousand kernel weight, phenotypic indications of nitrogen starvation [20]. Transgenic wheat cv Kasalath lines over-expressing the NADH-GOGAT rice orthologue (accession no. AB001916 corresponding to LOC_Os01g48960) under the control of its own promoter showed an increase in grain weight (80% maximum), supporting that NADH-GOGAT is indeed a key step in nitrogen utilization and grain filling in rice [12]. In the present study, the transcriptional level of the NADH-GOGAT-3A and NADH-GOGAT-3B genes was assessed in two durum wheat cultivars Svevo and Ciccio previously analyzed for grain protein content in several years and environmental conditions [17] and characterized by a high and low final grain protein content. qRT-PCR conducted with specific primers in order to analyze individually the two homoeologous genes revealed a differential pattern for the two genes in the two cultivars. A significantly different expression was observed between the two cultivars in roots with an higher level of transcript in the cv Svevo for both genes. This results suggest that the NADH-GOGAT genes play a crucial role in differencing the final grain protein content since these genes are principally active in non-photosynthetic germinating seed, roots and etiolated

![Figure 4. Dicot vs Moncot NADH-GOGAT amino acid sequences.](image)

The derived amino acid sequence of the wheat B-genome NADH-GOGAT protein is compared to sequences derived from DNA sequences of other plants (poplar, grape, Arabidopsis (Arab.), maize, rice, and Brachypodium (Brachy.). Amino acid residues conserved in all aligned proteins are shaded in yellow. Amino acids unique to monocots are shaded blue. Vertical red lines indicate exon boundaries. Exon numbers, such as ‘XS’ for exon 5, are to the right of the exon boundary lines. Known or presumptive functional domains of the NADH-GOGAT enzyme are below the alignment – with approximate domain boundaries [16] indicated by vertical green lines.

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![Figure 5. Physical mapping of the NADH-GOGAT gene on chromosome 3A.](image)

A gene specific marker was amplified in cvs Ciccio (C), Svevo (S), Chinese Spring (CS) and a set of nullitetrasomic and bin deletion lines. The 1 kb fragment was absent in the nulli-3A-tetra-3D line, in the ditelo-3AS, and in the bin line 3AL-4-0.42, as indicated by arrows.

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Results indicate that, in roots, the alleles encoded by cv Svevo are more transcribed than those of cv Ciccio. In leaves, the two cultivars reverse patterns of NADH-GOGAT transcript titers, both cultivars with peaks at the flowering stage and lowest titers at first leaf stage for Svevo and grain filling stage for Ciccio. These patterns are similar for both the 3A and 3B NADH-GOGAT genes, and the higher titers for Svevo at grain filling is consistent with the higher Svevo titers in roots.

The physical chromosome position of the NADH-GOGAT-3B gene co-localize with Meta QTLs for high protein content [14]. Besides a QTL for grain protein content also has been reported on shoots [21].

Figure 6. NADH-GOGAT primers. NADH-GOGAT-3A and NADH-GOGAT-3B specific primer combinations used in qRT-PCR analysis. The boxed base indicates the change in the NADH-GOGAT-3B forward primer to increase specificity. Primers are listed in Table 2.

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Figure 7. qRT-PCR of NADH-GOGAT. qRT-PCR conducted for NADH-GOGAT-3A and NADH-GOGAT-3B genes with specific probes. The figure shows: A. The expression levels of the two genes in the roots of the two durum wheat cvs Ciccio and Svevo (C and S, respectively); B. Comparison of the two homoeologous gene expressions in cv Ciccio during the stage of first leaf, flowering and grain filling; D. Comparison of the two homoeologous gene expressions in cv Svevo during the stage of first leaf, flowering and grain filling. The error bars indicate the ± SE of the mean.

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chromosome 3AL in the same genetic material used in the present paper in the homoeologous position of the \textit{NADH-GOGAT-3B} gene [17]. In conclusion, our results support the proposition that \textit{NADH-GOGAT} may be a major candidate gene driving NUE on chromosome group 3 in bread and durum wheat. Future experiments might address this proposition via specific absence or overexpression of the \textit{NADH-GOGAT} genes. Among the potential options for the former is transgene mediated silencing, deletion lines such as used in the present report [31], or use of wheat tillig populations to identify mutations [32]. Gene Overexpression could be accomplished by transformation [33] with gene \textit{NADH-GOGAT} gene constructs. These strategies are now available with the gene sequences reported in the current report.

**Materials and Methods**

**Plant Material and DNA Analysis**

The durum wheat cultivars Ciccio and Svevo were used for \textit{NADH-GOGAT} genomic sequence determination. These cultivars are the parentals of a mapping population represented by a set of 120 recombinant inbred lines (RILs) as previously described [22]. The two parents were chosen for differences in important qualitative and quantitative traits; i.e., grain yield components, grain protein content, yellow pigment, and adaptive traits.

Nulli-tetrasomic lines (NTs) of \textit{Triticum aestivum} cv Chinese Spring [23,24] were used to physically localize \textit{NADH-GOGAT} markers to specific chromosomes. Chinese Spring di-telosomic lines [25] were used for further assignment of markers to each chromosomal arm. Physical location on chromosome bins of each PCR fragment was obtained using a set of eight common wheat deletions lines dividing the 3A and 3B genome chromosomes into bins [26].

Genomic DNA was isolated from fresh leaves using the method as described [27] and subsequently purified by phenol-chloroform extraction. DNA amplifications were carried out as described previously [22]. For qRT-PCR, leaf and root samples were collected from 5 plants of both cultivars in different phenological stages (first leaf, flowering, grain filling).

**Gene Sequence Determination for Chinese Spring and \textit{Triticum tauschii}**

To isolate the complete sequences of the three \textit{NADH-GOGAT} genes in hexaploid wheat, we used a reported rice genomic sequence of \textit{NADH-GOGATI} [15] as the initial query. The rice sequence was used to probe public databases to extract the 454 sequences of the hexaploid wheat cv Chinese Spring (cerealsdb.uk.net/search_reads.htm) and the 454 reads of \textit{Triticum tauschii} (avena.pw.usda.gov/RHmapping/blast/). These two groups of 454 reads were separately assembled using the SeqMan software by DNAsStar (Lasergene). Using the public database at NCBI (http://www.ncbi.nlm.nih.gov/) we also identified a \textit{GOGAT} gene from the wheat B-genome within the previously reported wheat genomic DNA clone FN564429.

**Gene Characterization in the Italian Durum Wheat cvs Ciccio and Svevo**

A set of genome specific primer pairs were designed using OligoExplorer and Primer3 (frodo.wi.mit.edu/primer3/) software for the three distinct cv Chinese Spring \textit{GOGAT} hexaploid sequences previously obtained. Single PCR fragments were cloned from cultivar Chinese Spring using the vector pCR4-TOPO and transformed into DH10B electroMAX cells (Invitrogen, Cloning Kit) following the manufacturer’s instructions, and then sequenced at both insert ends with T3 and T7 primers and BigDye chemistry (Applied Biosystems) using a 96 capillary automatic sequencer ABI PRISM 3730XL. Gaps and uncertain sequence were resolved by primer walking. Regions of less coverage or ambiguous reads were rechecked with additional primers.

**Analysis of Sequences**

NCBI (ncbi.nlm.nih.gov) was used for annotation of the new wheat BAC sequence by BLAST analyses and total EST analyses by direct querying to NCBI. Exon/intron junctions are predicted

| Table 1. Primer pairs sequences for housekeeping genes [28]. |
|---|
| **Gene name** | **Orientation** | **Primer 5’-3’ sequence** | **Product length (bp)** | **Melt peak (°C)** |
| CDC | Forward | CAAATAGGCACATCCAGGAGACATC | 227 | 86.5 |
| | Reverse | CGTCGCGAAGACGACGACG | |
| RLI | Forward | CGATTAGAGGAGCGATATGTGG | 242 | 80 |
| | Reverse | AGTGTGCAGGGTCTCTATTAATG | |
| ADP-RF | Forward | GACCACATCTCTACAAG | 276 | 84.5 |
| | Reverse | AGACGACAGACCACACC | |

| Table 2. Primer pairs specific for the two isoforms of the \textit{NADH-GOGAT} gene. |
|---|
| **Gene name** | **Orientation** | **Primer 5’-3’ sequence** | **Product length (bp)** | **Melt peak (°C)** |
| \textit{NADH-GOGAT-3A} | Forward | GCCGCGATCATCGAGGAGGC | 113 | 86 |
| | Reverse | GCCGCGATCATCGAGGAGGC | |
| \textit{NADH-GOGAT-3B} | Forward | CAGCGGGCGATCATCGAGGAGGC | 108 | 83 |
| | Reverse | GCCGTGATCGAGCGCGTGGC | |
by alignment with Triticeae EST sequences, when available, or with other monocot EST if no Triticeae ESTs covered those sequences.

DNA sequences were compared using the Megalign (alignments and dot plots) and Seqman (assemblies) modules of the Lasergene suite (DNASTAR, Inc.). Sources of genomic sequences were as follows: Arabidopsis thaliana NADH-GOGAT, Genbank.ncbi.nlm.nih.gov (AT5G33460.1); Brachypodium distachyon, http://brachypodium.org; Bradi2g46670.1; poplar (Populus trichocarpa), genome.jgi-psf.org (POPT_00115s01950.1); grape (Vitis vinifera), genoscope.cns.fr/externe/GenomeBrowser/Vitis/GSVIT1038741001; rice (Oryza sativa), gramene.org (LOC_Os01g48960.1); maize (Zea mays), http://plantgdb.org/ZmGDB (GRMZM2G077054).

For ease of reading, it will be understood that common names and genus names will be used unless referring to specific species than noted above; e.g., Brachypodium instead of B. distachyon and rice instead of O. sativa. Plant ESTs were searched at Genbank, except for Brachypodium ESTs that were found at brachypodium.org.

RNA Isolation for Quantitative Real-time PCR

Tissues of each sample were harvested in each phase, frozen in liquid nitrogen and stored at -80°C until RNA extraction. Total RNA was extracted with RNeasy Plant Mini Kit (QIAGEN®) and checked on 1.5% denaturing agarose gels. The total amount of RNA and its purity was determined using Nano-Drop ND1000 spectrophotometer (Thermo Scientific, Walthman, MA, USA). All RNA samples were adjusted to the same concentration (1 mg/ml) for subsequent treatment with DNase I recombinant (Roche Applied Science, Mannheim, Germany) in order to remove genomic DNA, and then were reverse-transcribed into double stranded cDNA with Transcriptor First Strand cDNA Synthesis Kit (Roche Applied Science, Mannheim, Germany).

Reference Genes and Specific Gene Primer Design

Genes coding for Cell Division Control AAA-Superfamily of ATPases (CDC), ADP-Ribosylation Factor (ADP-RF) and RNase L Inhibitor-like protein (RLI) were used as reference genes as suggested by Zhang and Yuan [28,29] and checked on 1.5% denaturing agarose gels. The three cv Chinese Spring reference genes were evaluated by NormFinder software [30].

DNA sequences of NADH-GOGAT-3A and NADH-GOGAT-3B from the two durum cvs Svevo and Ciccio were aligned with Brachypodium, genome.ucdavis.edu (Brachypodium distachyon), genome.jgi-psf.org (POPT_00115s01950.1); grape (Vitis vinifera), genoscope.cns.fr/externe/GenomeBrowser/Vitis/GSVIT1038741001; rice (Oryza sativa), gramene.org (LOC_Os01g48960.1); maize (Zea mays), http://plantgdb.org/ZmGDB (GRMZM2G077054), and by sequencing of the amplified fragments (BMR Genomics, Padova, Italy).

The specificity of the amplicons was confirmed by the presence of a single band of expected size for each primer pair in agarose gels (2% w/v), by single peak melting curves of the PCR products, and by sequencing of the amplified fragments (BMR Genomics, Padova, Italy).

The specificity of the amplicons was confirmed by the presence of a single band of expected size for each primer pair in agarose gels (2% w/v), by single peak melting curves of the PCR products.

qRT-PCR data for both NADH-GOGAT and endogenous controls genes reported in this paper derive from the mean values of three independent amplification reactions carried out on five different plants harvested in the same phenotypic stage. All calculations and analyses were performed using CFX Manager 2.1 software (Bio-Rad Laboratories) using the 2^(-ΔΔCt) method which uses the Relative Quantity (RQ) calculated on ratio from relative quantity of the target gene and the relative expression of the reference gene (that includes the three reference targets in each sample). Standard deviation was used to normalize the expression values for the highest or lowest individual expression levels (CFX Manager 2.1 software user manual). All the results were analyzed by ANOVA.

Supporting Information

Figure S1 Alignment of 3A, 3B, and 3D NADH-GOGAT gene sequences. The three cv Chinese Spring NADH-GOGAT gene sequences are aligned from 5’ to 3’ regions common to all three genes using the Megalign module of Lasergene (DNASTAR, Inc.). Red brackets enclose exons whose numbers are in blue above the alignment. Introns are numbered according to the preceding exon. Intron 13 is indicated in red above the intron. Start and stop codons are boxed in red.

Figure S2 Comparison of 3A and 3B NADH-GOGAT genes. The NADH-GOGAT genes from the A- and B-genomes were analyzed by dot plot at a criterion of 100% match over a 20 base window.

File S1 Durum wheat NADH-GOGAT gene sequences. Fasta sequences of hexaploid cv Chinese Spring NADH-GOGAT genes and encoded polypeptides.

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Author Contributions

Conceived and designed the experiments: DN YG IM AB AG OA. Performed the experiments: DN NH IM. Analyzed the data: DN YG NH IM AG OA. Wrote the paper: DN AG OA.

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