Genetic Dissection Uncovers Genome Wide Marker-Trait Associations for Plant Growth, Yield and Yield Related Traits Under Varying Nitrogen Levels in Nested Synthetic Wheat Introgression Libraries

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Genetic dissection uncovers genome wide marker-trait associations for plant growth, yield and yield related traits under varying nitrogen levels in nested synthetic wheat introgression libraries

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**Key message**

To meet future wheat production demands, improving nitrogen use while maintaining grain yield is vital. We identified marker-trait associations affecting plant growth, yield and yield related traits under varying nitrogen levels. We also identified promising breeding lines with significant genetic variations and carrying the trait-associated markers or candidate genes. These may serve as potential donors to be exploited further in genomics-assisted breeding programs targeting improved NUE while maintaining grain yield in wheat.

**Abstract**

Nitrogen is one of the most important macronutrients for crop growth and metabolism. To identify marker-trait associations for complex NUE-related agronomic traits, field experiments were conducted on nested synthetic wheat introgression libraries at three nitrogen input levels across two seasons. The introgression libraries were genotyped using the 35K Axiom® Wheat Breeder’s Array and genetic diversity and population structure were examined. Significant phenotypic variation was observed across genotypes, treatments and their interactions across seasons for all the 22 traits measured. Significant positive correlations were observed among grain yield and yield attributing traits and root traits. Across seasons, a total of 233 marker-trait associations (MTAs) associated with fifteen traits of interest at differential levels of nitrogen (N0, N60 and N120) were detected using 9,474 genome-wide single nucleotide polymorphism (SNP) markers. Of these, 45 MTAs for 10 traits in the N0 treatment, 100 MTAs for 11 traits in the N60 treatment and 88 MTAs for 11 traits in the N120 treatment were detected. We identified putative candidate genes underlying the significant MTAs which were associated directly or indirectly with various biological processes, cellular component organization and molecular functions involving improved plant growth and grain yield. In addition, the top 10 lines based on N response and grain yield across seasons and treatments were identified. The identification and introgression of superior alleles/donors improving NUE while maintaining grain yield may open new avenues in designing next-generation nitrogen efficient high yielding wheat varieties.

**Keywords:** nitrogen, genome wide association studies, marker-trait association, wheat, yield
Introduction
The global demand for nitrogen currently stands at about 117 million metric tons with a projected annual increase of approximately 1.5% expected in the near future (FAO 2019). Farmers generally apply high doses of nitrogenous fertilizers to ensure good yields. The high input of commercially available fertilizers has led to the degradation of air, soil, and water quality (Hickman et al. 2014; Russo et al. 2017). In addition, when the supply of nitrogen is in excess of crop nitrogen demand, it increases the susceptibility of plants to various diseases and insect pests (Reddy 2017). Therefore, it is necessary to optimize and improve the nitrogen use efficiency (NUE) of cereal crops to maximize yield in addition to minimizing the negative impact of increase in nitrogen use on the environments and natural resources. Identification of marker-trait associations can be used to make effective targeted introgressions and is one possible genetic method to address the challenge of developing nitrogen efficient wheat varieties with stable yield under nitrogen limited environments.

Wheat varieties that maintain yield under moderate or intense nitrogen deficiency can adapt to low input systems. To breed such varieties, genetic variation for adaptation traits to nitrogen deficiency is required. To date, limited quantitative trait loci (QTL) for both yield and its response to N deficiency in wheat under field conditions have been documented. Detection of genotypes and underlying QTLs for maintaining yields at low nitrogen levels are of value in wheat breeding programs designed to increase nitrogen-deficiency tolerance. Some QTLs influencing nitrogen uptake have been genetically mapped in wheat under different doses of fertilizer application using bi-parental populations (An et al. 2006; Laperche et al. 2007; Xu et al. 2014; Deng et al. 2017; Mahjourimajd et al. 2016). A number of genetic loci for agronomic traits related to nitrogen use and grain yield have also been mapped to the chromosomal regions containing the GS2 gene in wheat and rice (Prasad et al. 1999; Obara et al. 2001; Yamaya et al. 2002; Fontaine et al. 2009; Habash et al. 2006; Laperche et al. 2007). This suggests the role of the genomic region surrounding GS2 is favourable in breeding wheat and rice varieties with improved agronomic performance and nutrient use efficiency. Other genetic regions associated with nutrient uptake have also been detected in rice (Wissuwa et al. 1998; Ming et al. 2000), wheat (Su et al. 2006; 2009), maize (Zhu et al. 2005), common bean (Liao et al. 2004; Yan et al. 2004), and soybean (Li et al. 2005; Liang et al. 2010). The NRT2.1, NRT2.2, and NAR2.1 gene have been reported to be the important contributors to the high affinity transport system in Arabidopsis roots (Orsel et al. 2006). Sixteen genes were identified in wheat homologous to characterized Arabidopsis low-affinity nitrate transporter NPF genes,
suggesting a complex wheat NPF gene family (Buchner and Hawkesford 2014). The regulation of wheat NFP genes by plant N-status indicated involvement of these transporters in substrate transport in relation to N-metabolism.

The phenotypic traits reported to be associated with NUE in cereal crop so far include root number, length, density and branching (Morita et al. 1988; Yang et al. 2012; Steffens and Rasmussen 2016), dense and erect panicle (Sun et al. 2014), plant height (Gaju et al. 2011), and leaf width (Zhu et al. 2020). The colocation of QTLs for N-uptake and root architecture traits have suggested that breeding for better and efficient root systems is a way to improve NUE (Coque et al. 2008; Sandhu et al. 2015).

Diverse accessions, landraces, breeding populations, and next-generation mapping populations, including nested-association mapping (NAM) and multi-parent advanced generation inter-cross (MAGIC) populations have shown potential for mining novel genetic variation in rice (Zhao et al. 2011; Subedi et al. 2019; Sandhu et al. 2019), wheat (Mackay et al. 2014), maize (Yu et al. 2008) and soybean (Xavier et al. 2015). NAM and MAGIC populations have proven advantageous over biparental populations as they capture additional recombination breakpoints thus increasing the allelic diversity and improving the power of QTL detection (Yu et al. 2008; Scott et al. 2020). Further, the availability of high throughput genotyping platforms to generate uniformly distributed genome wide molecular markers are critical for the high-resolution genetic dissection of polygenic traits, and the tracking of favourable alleles in breeding populations (Pandey et al. 2012; Varshney et al. 2013; Pandey et al. 2016). To date, a series of high-density wheat SNP arrays such as the Illumina 9K iSelect SNP array (Cavanagh et al. 2013), Illumina 90K iSelect SNP genotyping array (Wang et al. 2014), 15K SNP array (Boeven et al. 2016), Axiom® 660K SNP array, 55K SNP array, Axiom® HD 820K genotyping array (Winfield et al. 2016), 35K Axiom array (Allen et al. 2017) and 50K Triticum TraitBreed array (Rasheed and Xia 2019) have been developed and their utility has been demonstrated across a range of applications.

In the present study we developed nested synthetic wheat introgression libraries capturing novel genetic variation. The libraries were genotyped using a high-density SNP array and phenotypically assessed for root traits and agronomic performance under three nitrogen input conditions in the field. Genome-wide association mapping was used to identify marker-trait
associations for the root and agronomic traits and lines carrying favourable genetic combinations were also identified for use in future breeding for improved nitrogen use.

**Material and Methods**

**Plant material**
A total of 31 cultivated and 12 synthetic wheats were evaluated at 6 nitrogen levels (N0, N40, N80, N120, N160 and N200) in 3 replications in 2016-2017 and 2017-2018 during the rabi seasons at Punjab Agricultural University, Ludhiana, India. The synthetic wheats PDW233/Ae. tauschii acc. pau 14135 and PBW114/Ae. tauschii acc. pau 14170 produced high grain yields as well as high agronomic efficiency at low fertilizer N doses (unpublished data). These synthetic wheats were used to develop a nested synthetic hexaploid wheat (N-SHW) introgression library constituting a set of 352 lines derived from four sub-populations. The N-SHW library was made up of subsets from four populations (Pop1: 75 lines from PDW233/Ae. tauschii acc. pau 14135 amphiploid //2*BWL4444; Pop2: 106 lines from PDW233/Ae. tauschii acc. pau 14135 amphiploid //2*BWL3531; Pop3: 88 lines from PBW114/Ae. tauschii acc. pau 14170 amphiploid //2*BWL4444; Pop4: 83 lines from PBW114/Ae. tauschii acc. pau 14170 amphiploid //2*BWL3531 along with the two common parents (BWL3531, BWL4444) and other unique parents (PDW233, PBW114, Ae. tauschii acc. pau 14135 amphiploid and Ae. tauschii acc. pau 14170 amphiploid). The breeding scheme used to develop the N-SHW introgression library is summarised in Fig. 1.

**Agronomic practices and management of experiments**
The N-SHW library, six parents and two synthetic hexaploid wheats were assessed at the experimental farms of School of Agricultural Biotechnology, PAU Ludhiana (30° 54’ N latitude, 75° 48’ E longitude, and 247 m above sea level) over 2 years in 3 nitrogen level (6-year x N combinations). Details of the number of lines tested and experimental design is provided in Table 1. The breeding material was sown on 21th of November and 18th of November in 2018 and 2019, respectively. In both years the experiments were conducted at three nitrogen levels [i.e. zero N (0 Kg ha⁻¹), half N (60 Kg ha⁻¹) and full N (recommended, 120 Kg ha⁻¹)], referred to as N0, N1 and N2, respectively. The recommended dose of phosphorus, potassium and manganese was applied at the time of sowing. Half of the nitrogen (N) was applied at the time of sowing while the other half was applied in two equal splits, the first at crown root initiation stage and the remaining at the maximum tillering stage in both the
N1 and N2 experiments. N0 was treated as a control. Recommended fungicides and insecticides were applied to control stripe rusts, brown rusts, and aphids at jointing, booting and 10 days after anthesis to prevent diseases and pests. Weeds were controlled manually.

**Characterization of phenotypic traits**

A total of twenty-two traits were assessed in all experiments across both seasons except the maximum root length and root angle which were measured in 2018 only. The details of the NUE related traits, root and plant morphological traits, grain yield and yield attributing traits are presented in Supplementary Fig. S1. Destructive sampling of six plants per plot was done at 60 DAS to evaluate early root and shoot traits (Supplementary Fig. S2). Shoots were separated from the roots, fresh root weight (FRW; g) and fresh shoot weight (FSW; g) were measured. The root and shoot samples were dried at 70ºC in an oven until constant shoot dry weight (g) was observed, while the roots were cleaned thoroughly and stored in 70% alcohol at 4 ºC for root trait evaluation. MRL (maximum root length) and RA (root angle) were measured using ImageJ software. TRL (total root length), RSA (total root surface area), RD (total root diameter), RV (total root volume), NF (number of forks) and Ntips (number of tips) were recorded using WinRhizo STD4800 (Supplementary Fig. S2). The roots were then dried at 70ºC in the oven until constant RDW was observed. The data on nitrogen uptake related traits was recorded using chlorophyll meter (SPAD502) and leaf color chart (LCC). The LCC provides a decision support system to the farmers for sustaining the high yields with optimum nitrogen dose in the field crops. It measures the leaf color variations of 6 SPAD (Soil Plant Analysis Development Meter) units comprising 3, 3.5, 4.0, 4.5, 5.0 and 6.0 and provides nitrogen recommendation in the field crops. Flag leaf length (FLL) and flag leaf width (FLW) were recorded using a centimeter scale. Days to 50% flowering (DTF) was recorded when 50% of the plants in a plot exerted their panicles. Spikelets per spike (SPS) was counted manually from five random plants. NPT (number of productive tillers) was counted manually in 0.5 m row length and SB (shoot biomass) at harvesting was measured from 0.5 m row length. PHT (plant height) in cm was measured as the mean height of five random plants for each entry measured from the base of the plant to the tip of the panicle during maturity stage. The plants were harvested at physiological maturity or when 80-85% of the panicles turned to golden yellow and the panicles at the base were already at the hard dough stage; harvested grains were threshed, dried and weighed to determine the GY (grain yield).
Phenotypic data analysis

Analysis of variance (ANOVA), experiment and experiment-wise mean for each season was calculated using mixed model analysis in PBTools V 1.4.0 for augmented design and in STAR Version: 2.0.1 for the split plot design. In split plot design the nitrogen levels were considered as the main plot and the breeding lines as subplot. Fisher’s t-test was used to determine the significant difference among the breeding lines, treatments and to estimate the interactions. The correlation analysis among traits was performed in R. v.1.1.423.

To evaluate the phenotypic stability and grain yield adaptability of the breeding lines across seasons and treatments, the GGE biplot analysis was performed, considering the effects of genotype (G) and genotype by environment (GE) as random. The best linear unbiased prediction (BLUP) values of the G and GE effects were calculated. The multiplicative model in PB tool version 1.3 (bbi.irri.org) was used to explain the relationship between genotype and seasons.

Genotypic data

High-density genotyping was performed using the 35K Axiom® Wheat Breeder’s Array (Affymetrix UK Ltd., United Kingdom). The quality pre-processing of 35,143 markers obtained from the 35K chip was done using PLINK software (Purcell et al. 2007). A total of 9,474 SNPs with MAF (minor allele frequency) of >5%, maximum heterozygote proportion of 0.1 and missing rates < 0.1 were used to estimate genetic relationships and for the mapping of marker-trait associations for different traits associated with plant growth, yield and yield related traits. APCA was carried out to detect and correct for population structure.

Population structure and association analysis

The model-based STRUCTURE V. 2.3.4 software was used to test K values from 1 to 10, with a burn-in period to 10,000 and 1,000,000 MCMC reps after burn-in in order to assess population structure in the 352 breeding lines using a total of 9,474 SNPs. The consistency and accuracy of the results was validated across 10 runs for each K. The K value with maximum likelihood over the 10 runs was used to estimate the most appropriate number of clusters (Pritchard and Wen 2004). The population structure was determined by plotting the proposed number of subpopulations against the delta k (Earl and Vonholdt 2012). Principal components analysis (PCA) was performed in R/GAPIT and added iteratively to the fixed model, ranging from PC1 to PC10.
Significant marker-trait associations were identified using CMLM (compressed mixed linear model)/P3D (population parameters previously defined) in GAPIT (Genome Association and Prediction Integrated Tool) executed in R. Identity by state (IBS) values and a relatedness matrix were used to estimate the random effect and genetic similarity of the accessions, respectively. The statistical power of the association studies was further improved by considering the population structure ($Q$ value) and kinship matrix ($K$) estimated from the genotyping data. The Bonferroni correction method was used to correct for false positives in the analysis, using the stringent p-value benchmark. The Bonferroni multiple test correction was performed (0.05/9474; significance level of 5%/total number of markers used in analysis) and the calculated threshold value was $5.28 \times 10^{-6}$. The allelic effect of all the significant markers associated with the measured traits was determined by comparing the mean phenotypic values and the significant allelic variants for the trait/s using a Kruskal–Wallis test in R.

Candidate gene analysis and functional annotation of putative candidate genes

SNPs that exhibited a false discovery ratio (FDR) corrected p-value <0.05 for a particular trait of interest were evaluated as markers for the potential putative candidate genes. A window of 1Mb adjacent to each significant SNP was examined for candidate genes and annotations were identified through the Ensemblplants database (http://plants.ensembl.org/index.html).

The functional annotation and gene ontology of identified putative candidate genes was performed using OMIX box software. Blast (E-value≤10^{-5}) was performed using the CloudBlast tool against Triticum (nr_subset)[monocots_triticum, taxa:4564] and NCBI non-redundant database (http://www.ncbi.nlm.nih.gov), followed by the InterPro using CloudIPS, followed by GO mapping (Gene Ontology), and annotation configuration. GO terms were then used to generate the semantic similarity-based scatterplots/interactive graphs/tag clouds by using REVIGO (http://revigo.irb.hr/).

Defining N-insensitive and N-sensitive lines

The genotypes that showed more or equal/stable grain yield with the minimal application of nitrogen fertilizer when compared to the recommended or standard nitrogen fertilizer application, were considered as the nitrogen insensitive genotypes (NIS) or the top grain yielders across seasons and treatments. On the other way around, the genotypes that were low yielding or not able to maintain the grain yield with the minimal application of nitrogen
fertilizer when compared to the recommended or standard nitrogen fertilizer application, were considered as the nitrogen sensitive genotypes (NS) or the poor grain yielders across seasons and treatments.

Results

**Significant phenotypic trait variation and correlations detected across nitrogen treatments**

The 352 N-SHW lines, six parents and two synthetic hexaploid wheat donors were screened for twenty-two traits in six growing conditions (2 years x 3 nitrogen level). Analysis of variance (ANOVA) revealed significant genetic variation for the root, plant morphological and agronomic traits among genotypes, treatments, seasons and their interactions (genotype x treatment, genotype x season, treatment x season and genotype x treatment x season) (Table 2). The detailed information on trial means, LSD and heritability for all the traits measured are presented in Supplementary Table S1. The results revealed significant genetic variations across genotypes, treatments and interactions in 2018-2019 and 2019-2020 seasons for all the traits measured (Supplementary Table S2). The phenotypic data of the traits measured in the present study were averaged across two seasons and are presented as mean values in Supplementary Table S3.

GY increased with applied N level. In the N0 treatment, the average GY of the tested breeding lines across seasons was 2022 kg ha⁻¹ and ranged from 564 to 4092 kg ha⁻¹ (Supplementary Table S3). In the N60 treatment, the GY varied from 882 to 4685 kg ha⁻¹ with average GY of 2357 kg ha⁻¹ and, while in N120 treatment, the GY varied from 1332 to 4270 with an average of 2579 kg ha⁻¹ (Supplementary Table S3). Across seasons, N in the limited conditions (N0) resulted in the 14% and 22% GY reduction compared to N60 and N120 treatments, respectively. The N application also significantly increased the SB by 8% in N60 and 52% in N120 treatment across seasons. The average NPT across experiments was higher in N120 (28) compared to N60 (24) and N0 (22) (Supplementary Table S3). Under the N0 treatment, the average LCC value varied from 3.3 - 4.8, and ranged from 3.5 - 5.1 in the N60 treatment and from 4.1 - 5.3 in the N120 treatment (Supplementary Table S3). The response of lines in terms of average DSW across seasons increased from 3.28 in N0 to 3.62 in N60 to 3.75 in N120 treatment (Supplementary Table S3). The minimum and maximum value of DRW under N0 were 0.187 g and 2.425 g; 0.298 g and 2.001 g under N60 and 0.338 g and 2.333 g under N120,
respectively (Supplementary Table S3). The average root diameter was highest under N60 (0.610 g) compared to N0 (0.560 g) and N120 (0.409 g) (Supplementary Table S3). Across seasons, average flowering was delayed by 2 days under the N0 treatment compared to the N60 and N120 treatments. Average PHT was lower (92 cm) in N0 compared to N60 (95 cm) and N120 (99 cm).

We calculated the Pearson’s correlation coefficients between all the traits measured in N0 (Fig. 2A), N60 (Fig. 2B) and N120 (Fig. 2C) treatments. The Pearson correlation coefficients across all treatments considering pooled mean data for all traits measured in the present study is presented in Supplementary Fig. S3. The strongest and most significant positive correlation among grain yield and yield attributing traits and root traits were observed in N60 treatment. The grain yield was significantly and positively correlated with SB (r = 0.23, p <0.001), NPT (r = 0.18, p <0.01), FRW (r = 0.16, p <0.01), FSW (r = 0.24, p <0.001), DSW (r = 0.23, p <0.001) and with RSA = (r = 0.23, p <0.001). Across treatments GY showed negative correlation with DTF, SB showed positive correlation with GY.

**Population structure analysis detected three genetic sub-populations**

The population structure of the N-SHW lines was assessed to understand the genetic structure of the 352 lines based on 9,474 SNPs distributed across all 21 wheat chromosomes. The most appropriate K explaining the population structure was K=3 at MAF ≥ 5% (Fig. 3A). The kinship heatmap indicated a weak relatedness in the panel (Fig. 3B). The first three principal components (PCs) were most informative gradually decreasing (Fig. 3C, Fig. 3D) until the tenth PC. The kinship and PCs were considered during the GWAS analysis to correct for population structure. The appropriate number of sub-populations was determined from the largest delta K value of 3 (Fig. 3E).

**Mapping reveals significant marker-trait associations for all traits**

GWAS was performed exploiting the phenotypic variability in the 352 N-SHW lines using 9,474 SNPs from the 35K Axiom® Wheat Breeder’s Array. Using the -log(P) ≥0.001 at 5% significance level, a total of 233 marker-trait associations (MTAs) were detected across seasons associated with fifteen traits of interest at differential N levels (N0, N60 and N120; Table 3). Of these, 45 MTAs for the 10 traits in the N0 treatment, 100 MTAs were associated with 11 traits in the N60 treatment and 88 MTAs were associated with 11 traits in the N120 treatment.
Across seasons and nitrogen treatments, a total of 53 MTAs associated with more than one trait/treatment were detected (Table 3). In addition to these 53 MTAs, another 41 MTAs associated with single trait only were detected across seasons (Supplementary Table S4). All MTAs detected in the present study either in one season, both seasons, each treatment or across treatments are compiled in Supplementary Table S5. Mapping detected MTAs on all subgenomes (A: 42, B: 18 and D: 34) across seasons and treatments. The highest number of MTAs were detected on chromosome 6A (26) followed by 2D (25), 3B (12), 4A (9), 6D (8), 2A (6), 2B (4) with 1 MTA on each of 1B, 4B, 5A and 7D. Considering all assessed traits, significant MTAs were reported for FRW (2A, 2D), FSW (2A, 2B, 2D, 5A, 7A), DRW (2A, 2B, 2D, 7A), DSW (2A, 2B, 2D), FLW (4A, 4B), NT (3A), SPS (1B), DTF (3A, 3B, 6A), SB (6A), TRL (6D), RSA (6A, 6D), RV (6D), tips (6A, 6D) and forks (6A, 6D).

The Manhattan plots depicting the significant -log (p-values) for the MTAs associated with NUE-related traits, root traits and yield/yield related traits measured in the present study at differential level of N are presented in Supplementary Fig. S4, Supplementary Fig. S5 and Fig. 4, respectively. Location of significant MTAs and SNP marker density distributed across 21 wheat chromosomes is presented in Fig. 5. The SNPs for positively correlated traits such as GY, BY, tips, RSA, RV and forks appeared to be collocated on chr 6A at differential level of N (Table 3). A genomic region on 2D (ranging from 576749639 - 702726797 bp) contained 25 detected MTAs for a range of traits (FRW, DRW, FSW and DSW) across seasons and treatments (Table 3; Fig. 5). A cluster of 17 SNPs spanning a 7.7 Mb region on the short arm of 6A showed association with GY at N60 and N120 (Fig. 5). Across seasons and treatments, significant association in a 198 kb region on the long arm of chr 6A were detected for root traits (RSA, RV, tips and forks). The SNP, AX-94565231 at 683.64 Mb on the long arm of 6D showed association with different root traits (RSA, RV, tips and forks) across seasons and treatments. In the N60 treatment, significant associations for FLW were detected in a 5.6 Mb region (549799824 - 544201748 bp) on the long arm of 4A. Interestingly, the association of the trait DTF with SNP AX-95136655 on chr 3B at 234.49 Mb was common under N0, N60 and N120 treatments (Fig. 5). In the N0 treatment, significant associations harbouring three strongly associated SNPs (AX-94593608, AX-94786978 and AX-95134564) spanning the genomic region 76 bp on the long arm of 3A were detected for NPT (Table 3). Further, single SNPs were identified in association with different traits at different N levels. For example, the SNP AX-94914391 (36.43 Mb, 6A) was significantly associated with SB at N0 and with GY
at both N60 and N120 (Fig. 5). The SNP AX-94705680 (598.80 Mb, 2B) showed association with FSW at N60 and with DRW at N120.

**Candidate gene identification and functional annotation**

In order to identify candidate genes underlying the consistent MTAs, we surveyed putative candidates in a 1Mb upstream and 1 Mb downstream region the identified significant SNPs using EnsemblPlants (http://plants.ensembl.org/index.html). Detailed information on the identified candidate genes is presented in Table 3.

The GO term of identified putative candidate genes were categorised into four groups according to their trait relatedness; NUE uptake related (LCC, SPAD, FSW, DSW), root morphological (MRL, TRL, RSA, RD, RV, NF, Ntips, FRW, DRW), plant morphological (FLL, FLW, PHT) and grain yield/yield attributing traits (DTF, S/S, NPT, SB, GY). Most of the putative candidate genes in NUE uptake related traits across treatments were associated with protein phosphorylation/proteolysis, recognition of pollen, molybdoprotein cofactor biosynthetic process, and transmembrane transport (Supplementary Table S6). Some were part of the cellular component organization and molecular functions of binding molecules and ions, catalytic activity, peptidase activity and transmembrane transport activity (Supplementary Table S6; Supplementary Fig. S6). The putative candidate genes for the root morphological traits were associated with nitrogen compound metabolic processes, phosphorylation, proteolysis, catabolic processes, response to stresses, regulation of flower development by delineating the composition and architecture of gene regulatory network underlying flower development, and carbohydrate metabolism (Supplementary Table S6; Supplementary Fig. S7). The cellular components include chloroplast, ribosome, membrane, cytoplasm, nucleus and mitochondria (Supplementary Table S6; Supplementary Fig. S7). The primary molecular functions related to these genes were catalytic activity (protease, peptidase, hydrolase, transferase, ligase, and oxidoreductase), and binding activity (small molecule binding, ion binding, lipid binding, and carbohydrate derivative binding) (Supplementary Table S6; Supplementary Fig. S7). The putative candidate genes for the plant morphological traits were mainly associated with phosphorylation, response to light-intensity, stress-related responses, and metabolic processes. They were related to the molecular functions of metal ion binding, catalytic activity, kinase activity, and DNA/RNA/ATP binding (Supplementary Table S6; Supplementary Fig. S8). The yield and yield attributing traits related putative candidate genes were associated with phosphorylation, metabolic process, protein folding, catabolic process, response to water-stress and light, flower development and pollen
recognition (Supplementary Table S6; Supplementary Fig. S8). The molecular functions include catalytic activity (peptidase, hydrolase, lyase, oxidoreductase, transferase), binding activity (ion, metal, ATP/GTP, polysaccharide, protein, DNA) and metabolic activity (Supplementary Table S6; Supplementary Fig. S9).

**Selection of promising lines with stable performance for use in breeding**

To identify stable breeding lines across treatments and seasons, a GGE biplot method was used. The first two PCs (principal components) explained 77.7% (PC1=50.3%, PC2=27.4%) of the total GGE variation in the data (Fig. 6). The ranking of breeding lines based on their mean GY and stability across seasons and treatments (Supplementary Table S7) was used to identify 20 breeding lines with high and stable yield across seasons and treatments (Supplementary Table S8; Fig. 7). Based on GY data across seasons and treatments, the top 10 N-insensitive (NIS-top grain yielders) and 10 N-sensitive (NS-poor grain yielders) breeding lines were identified (Table 4).

Further analysis was undertaken to assess the significant differences between the mean values of the allelic classes of MTAs for root growth and grain yield using the Kruskal–Wallis test. The presence of favourable alleles with significant differences was checked in promising breeding lines. This allowed the selection of 20 promising breeding lines possessing the favourable allele combinations for improving plant root growth (Fig. 8A) and grain yield under N limitation (Fig. 8B).

**Discussion**

Increase in crop production by development of high-yielding varieties is largely dependent on the supply of N fertilizers. Excessive application of nitrogenous fertilizer is becoming very expensive which accounts for the great loss of economic profit to the farmers in addition to the negative impacts on the environment (Hawkesford and Griffiths 2019). The reliable phenotyping under low nitrogen input is very challenging and affected by genotype (G), environment (E), and the G x E interactions (Rao et al. 2018). Proper understanding of the genotype behaviour, identification and development of nitrogen efficient genotypes without compromising the GY is a paramount need for improving the NUE. Notably, very few wheat breeding programs are targeting the development of nitrogen efficient genotypes. In crop plants such as wheat, the efforts are constrained due to the lack of variation in the cultivated germplasm for NUE. The narrow genetic diversity and fewer recombination events in the biparental mapping populations
may result in poor QTL detection power (Gangurde et al. 2019). The next generation high-resolution mapping populations such as nested synthetic wheat introgression libraries used in the present study may provide a vast and untapped source of genetic variations for the nitrogen use efficiency related traits due to high numbers of recombination events. The use of synthetic hexaploid wheat in the present study presenting an effective genetic resource for transferring the agronomically important genes from wild relatives to the common wheat (Li et al. 2018). The introgression of favourable alleles associated with root traits and grain yield from *Ae. tauschii* wild accessions to cultivated wheat (Fig. 8) indicated the potential of synthetic wheat providing new sources for improving yield potential and nutrient-use efficiency when bred with the modern wheat varieties.

The different traits associated with nitrogen uptake and nitrogen use efficiency were studied in nested synthetic wheat introgression libraries at three different nitrogen levels. The ANOVA results revealed the native variation across the genotypes toward the nitrogen response which had given the possibility to identify the nitrogen use efficient lines under differential levels of nitrogen. The genotypic variations purely reveal the phenotypic plasticity of the breeding lines toward traits. The diverse responses have been observed among the breeding lines across different level of nitrogen, despite similar growth conditions and an equal amount of nitrogenous fertilizer application in a given N level as indicated by significant differences among the genotypes within and across treatments and non-significant differences among the replications. Significant G x E, G x S, G x E, G x T x S interactions indicated that the seasons and environments under different level of N application was a critical factor in explaining the genotypic variance for the traits measured in the present study. The results reported in the present study concurred with other reported studies in rice (Srikanth et al. 2016) and wheat (Sial et al. 2005; Belete et al. 2018).

In general, the increase in GY was correlated with the increase in the rate of N fertilizer application, which might be due to availability of sufficient nitrogen for proper growth and development of the plants. Šarˇcevi´c et al. (2014) reported 10% reduction of GY at low N condition compared to normal condition in wheat. The significant and positive correlation among different root traits and GY and yield attributing traits indicated complementary functional roles of the root traits in improving grain yield by improving nutrient acquisition from the soil. The collocation of MTAs for the correlated traits strengthens the significance of MTAs. A significant positive correlation between GY and NUE related traits in wheat, maize and oilseed
rape (He et al. 2017; Belete et al. 2018; Fageria et al. 2010) signified the importance of NUE related traits in improving GY under limited N conditions.

Different mapping approaches using NAM populations successfully exploited the genetics of complex traits and facilitated the discovery of candidate genes in rice (Fragoso et al. 2017), wheat (Hu et al. 2018; Jordan et al. 2018), maize (McMullen et al. 2009; Yu et al. 2008) and soybean (Song et al. 2017; Xavier et al. 2018). For NUE-related traits, significant genetic variations in hybrids, open-pollinated populations, large germplasm panels, backcross and recombinant inbred line populations in different cereal crops such as rice, wheat, maize and oilseed rape were observed (Chen et al. 2014; Vijayalakshmi et al. 2015; Li et al. 2015; He et al. 2017; Ertiro et al. 2017; Rao et al. 2018). Mapping for NUE related traits using different populations and mapping approaches highlight the complex nature of the trait.

In the present study, the nested synthetic wheat introgression libraries were designed for the identification of genomic regions associated with traits related to NUE using GWAS approach keeping into account the genetic effects produced in each genetic background. The associated SNPs were used to track the potential candidate genes associated with a particular trait of interest. The presence of high phenotypic variability in the nested synthetic introgression libraries coupled with the high marker density across the whole genome provided a strong base to the association mapping.

Interestingly, the genes responsive to nutrient uptake under water stress (Wang et al. 2017; Diédhiou et al. 2008; Janicka-Russak and Kabala 2015), shoot growth, root and plant development (Wang et al. 2017), nutrient uptake and transport of various nutrients (Wang et al. 2014; Weng et al. 2020; Takahashi et al. 2012) reported to be collocated with 126 Mb genomic region on chr 2D constituting 25 MTAs which stood out as hot-spot for different traits (FRW, DRW, FSW and DSW) in the present study. This indicates the positive interactions between root traits, nutrient uptake and plant growth and development. The 7.7 Mb region on short arm of Chr 6A constituting 17 SNPs associated with GY showed collocation with the genes that were directly or indirectly involved in improving grain yield in different cereal crops. These include the genes controlling flowering (Kania et al. 1997), panicle and seed development (Jain et al. 2007; Li et al. 2011), grain yield (Terao et al. 2010), resistance to pathogenesis (Taniguchi et al. 2013; Wang et al. 2014; Niño et al. 2020) and abiotic stress tolerance (Palusa et al. 2007; Brands and Ho, 2002). The MTAs associated with different root
traits such as RSA, RV, tips and forks in the present study were located near the earlier reported genes involved in regulating abscisic acid sensitivity and root growth development in Arabidopsis (Rodriguez et al. 2014) and adaptation under water stress conditions in wheat (Singh et al. 2017). Interestingly, the gene accelerating flowering in Arabidopsis (Hwang et al. 2019) was observed to be collocated with the SNP AX-95136655 associated with DTF on chr 3B in the present study. The colocation of identified MTAs with earlier reported genes controlling the photosynthetic traits, root development, plant growth, nutrient uptake and transport, flowering, resistance to pathogenesis and stress-responsive genes further confirms the contribution of these identified traits/MTAs in improving nitrogen uptake/utilization and grain yield under N limited conditions. The identified nitrogen insensitive breeding lines with favourable alleles in combination for the multiple traits might serve as potential donors for the development of nitrogen efficient wheat varieties.

**Conclusions**

The nested synthetic introgression libraries covering extensive phenotypic variability coupled with huge genome coverage was used to identify the significant MTAs associated with NUE related traits in wheat. Significant phenotypic variations for the NUE related traits, yield and yield related traits among genotypes, treatments, seasons and their interactions (genotype x treatment, genotype x season, treatment x season and genotype x treatment x season) were observed. Stable MTAs identified for different traits measured in the present study co-migrating with various genes associated with nitrogen uptake/utilization and improving grain yield may help to harness their benefits in genomics-assisted breeding programs. The identification of nitrogen efficient breeding lines may serve novel donors in genomics-assisted introgression programs. The identification and introgression of superior haplotype improving NUE while maintaining grain yield using haplotype-based breeding may open new avenues in designing next-generation nitrogen efficient high yielding wheat varieties.

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**Author contribution statement**

NS and PC designed this study; AK provided the genotypic data of two populations and contributed to the development of nested introgression libraries; NS and MS conducted the
field experiments, NS analysed the data; NS, SK and PC provided resources; NS wrote the manuscript, and all co-authors revised the manuscript.

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**Data availability**

The all-supported information’s are available in supplementary material.

**declarations**

**Conflicts of interest**

The authors declare that they no conflict of interests.

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### Table 1 Details on experiments conducted in 2018-2019 and 2019-2020 rabi season

| Pop  | Pedigree                      | Total no lines | Design                                                                                                                                 |
|------|-------------------------------|----------------|--------------------------------------------------------------------------------------------------------------------------------------|
| Pop1 | PDW233- *Ae. tauschii* acc. pau14135 amphiploid // BWL4444 | 75             | Augmented/ Split plot design, nitrogen level main plot, breeding lines as Subplots, 2 replications, 2 rows plot (1.5 m long with 20 cm row to row spacing) |
| Pop2 | PDW233- *Ae. tauschii* acc. pau 14135 amphiploid // BWL3531    | 106            | Augmented design/ Split plot design, nitrogen level main plot, breeding lines as Subplots, 2 replications, 2 rows plot (1.5 m long with 20 cm row to row spacing) |
| Pop3 | PBW114- *Ae. tauschii* acc. pau 14170 amphiploid//BWL4444       | 88             | Split plot design, nitrogen level main plot, breeding lines as Subplots, 2 replications, 1.5 m x 2 rows plot                           |
| Pop4 | PBW114- *Ae. tauschii* acc. pau 14170 amphiploid//BWL3531       | 83             | Split plot design, nitrogen level main plot, breeding lines as Subplots, 2 replications, 1.5 m x 2 rows plot                           |
Table 2: Analysis of variance (ANOVA) for the NUE related, root, plant morphological, yield and yield related traits among G (genotypes), (T) treatments, (S) seasons and their interactions (G x T, genotype x treatment; G x S, genotype x season; T x S, treatment x season; and G x T x S, genotype x treatment x season)

| Population          | LCC   | SPAD  | FRW   | DRW   | FSW   | DSW   | DTF   | NPT   | PHT   | SPS   | SB    | FLL   | FLW   | GY    | TRL   | RSA   | AD    | RV    | Tips  | Forks |
|---------------------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| PDW233/ Ae. tauschii| G     | 7.45  | 4.47  | 2.44  | 1.89  | 5.84  | 7.2   | 58.79 | 3.22  | 4.9   | 4.92  | 2.27  | 35.01 | 88.48 | 5.14  | 1.56  | 3.90  | 0.87  | 3.81  | 8.84  | 3.13  |
|                     | S     | 612.8 | 637.2 | 117.9 | 18.49 | 447.45| 394.06| 605.42| 30.88 | 4.04  | 2.08* | 7.22  | 24196 | 44138 | 109.4 | 124.8 | 4.93  | 90.05 | 1.269 | 1062  | 277.8 |
| T x S               |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |
|                     | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   | ***   |
|                     |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |
| PBW114/ Ae. tauschii| G     | 2.24  | 2.3   | 1.94  | 3.15  | 1.42  | 1.63  | 3.40  | 1.60  | 3.22  | 1.45  | 1.54  | 1.65  | 1.49  | 1.81  | 2.01  | 1.31  | 11.06 | 2.78  | 2.15  | 1.66  |
|                     | S     | 42.69 | 2.94  | 565.7 | 2.69  | 11.69 | 248.86| 8439.4| 1.24  | 98.01 | 893   | 59.26 | 1026  | 1364  | 2.10* | 1122  | 516   | 653.4 | 4.69  | 449.5 | 682.7 |
|                     |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |       |
|     |  G x T  |  G x S  |  T x S  |  G x T x S  |
|-----|---------|---------|---------|------------|
|     | 2.65    | 2.65    | 90.59   | 2.10       |
|     | 2.58    | 2.55    | 95.61   | 2.14       |
|     | 2.22    | 2.0     | 51.14   | 1.89       |
|     | 1.35    | 1.54    | 13.09   | 1.25       |
|     | 1.43    | 2.99    | 18.81   | 1.53       |
|     | 1.47    | 2.13    | 13.57   | 1.80       |
|     | 3.78    | 1.94    | 69.76   | 3.75       |
|     | 1.67    | 1.47    | 21.72   | 1.58       |
|     | 2.80    | 1.29    | 584     | 3.22       |
|     | 1.75    | 1.99    | 17.82   | 1.26       |
|     | 1.34    | 1.32    | 43.22   | 1.26       |
|     | 1.26    | 1.71    | 40.5    | 1.41       |
|     | 1.68    | 1.48    | 120.5   | 1.65       |
|     | 1.74    | 1.98    | 21.72   | 1.71       |
|     | 2.46    | 1.49    | 4.05    | 1.36       |
|     | 1.37    | 2.94    | 120.5   | 2.94       |
|     | 2.94    | 0.97    | 25.83   | 2.90       |
|     | 1.42    | 2.94    | 31.73   | 1.46       |
|     | 1.79    | 2.94    | 20.1    | 1.79       |

**Significance Levels:**

- ***: Significant at <0.001 level
- **: Significant at <0.01 level
- *: Significant at <0.05 level

*PBW114/ Ae. tauschii 14170 amphiploid/
BWL4444*
| SNP          | Chr | Position (bp) | Trt/Trait          | p-value     | R²  | FDR  | Gene stable ID       | Gene end (bp) | Gene start (bp) | Description                                                                 | Function                                                                 |
|-------------|-----|---------------|--------------------|-------------|-----|------|----------------------|---------------|-------------------|-----------------------------------------------------------------------------|--------------------------------------------------------------------------|
| AX-95136668 | 3A  | 690432670     | N60, N120 (GY)     | 5.85E-07    | 0.202 | 0.001 | TraesCS3A02G452300   | 690460596     | 690459736         | flowering-promoting factor 1-like protein 2                               | regulates flowering (Kania et al. 1997) and gibberellin signalling pathway |
| AX-94415776 | 6A  | 28700804      | N60, N120 (GY)     | 1.54E-06    | 0.2   | 0.001 | ENSRNA050010222      | 28845847      | 28845775          | putative disease resistance protein At3g14460                               | defence response to fungus (Bianchet et al. 2019)                         |
| AX-94978974 | 6A  | 29876500      | N60, N120 (GY)     | 1.20E-06    | 0.202 | 0.001 | TraesCS6A02G056800   | 29879453      | 29877038          | putative F-box protein At3g16210                                         | regulates gibberellin signalling (McGinnis 2003); panicle and seed development in rice (Jain et al. 2007, Li et al. 2011) |
| AX-94737868 | 6A  | 29876631      | N60, N120 (GY)     | 6.07E-07    | 0.204 | 0.001 | TraesCS6A02G057000   | 29969466      | 29967087          | L-type lectin-domain containing receptor kinase IX.1 LURP-one-like protein (DUF567) | promotes cell death (Wang et al. 2015), resistance response to pathogens (Wang et al. 2014) defence and resistance to H. parasitica mediated by the R-proteins RPP4 and RPP5 (Gallego-Giraldo et al. 2018) |
| AX-95210745 | 6A  | 29967076      | N60, N120 (GY)     | 1.27E-07    | 0.202 | 0.001 | TraesCS6A02G058500   | 30876811      | 30873143          | F-box protein At5g03970-like                                             | APO gene in rice improved grain yield per plant (Terao et al. 2010)       |
| AX-95631197 | 6A  | 30030973      | N60, N120 (GY)     | 5.94E-07    | 0.206 | 0.001 | TraesCS6A02G057100   | 30036467      | 30032329          | LURP-one-like protein (DUF567)                                           | APO gene in rice improved grain yield per plant (Terao et al. 2010)       |
| AX-95011132 | 6A  | 30031026      | N60, N120 (GY)     | 7.51E-07    | 0.202 | 0.001 | TraesCS6A02G058500   | 30876811      | 30873143          | L-type lectin-domain containing receptor kinase IX.1 LURP-one-like protein (DUF567) | APO gene in rice improved grain yield per plant (Terao et al. 2010)       |
| AX-95255669 | 6A  | 30873212      | N60, N120 (GY)     | 8.32E-07    | 0.202 | 0.001 | TraesCS6A02G058500   | 30876811      | 30873143          | L-type lectin-domain containing receptor kinase IX.1 LURP-one-like protein (DUF567) | APO gene in rice improved grain yield per plant (Terao et al. 2010)       |
| AX-95219967 | 6A  | 31036496      | N60, N120 (GY)     | 1.11E-06    | 0.201 | 0.001 | TraesCS6A02G058700   | 31036808      | 31034388          | LURP-one-like protein (DUF567)                                           | APO gene in rice improved grain yield per plant (Terao et al. 2010)       |
| AX-95070275 | 6A  | 31048271      | N60, N120 (GY)     | 7.49E-07    | 0.204 | 0.001 | TraesCS6A02G063700   | 34033068      | 34029127          | F-box protein At5g03970-like                                             | APO gene in rice improved grain yield per plant (Terao et al. 2010)       |
| AX-94970334 | 6A  | 31474354      | N60, N120 (GY)     | 1.30E-06    | 0.199 | 0.001 | TraesCS6A02G064600   | 34285946      | 34283159          | predicted protein serine/arginine-rich splicing factor RS41 isoform X2      | abiotic stress tolerance (Palusa et al. 2007)                              |
protein accumulation and replication of chloroplasts 6, chloroplastic transcription termination factor MTERF4, chloroplastic-like S-(+)-linalool synthase, chloroplastic-like protein STRUBBELIG-RECEPTOR FAMILY 5-like papain-like cysteine proteinase control male-sterility, organ development, cell proliferation in Arabidopsis (Chevalier et al. 2005) up-regulation of multiple pathogenesis-related proteins and biosynthesis of secondary metabolites (Niño et al. 2020), proteolysis and physiological processes (Liu et al. 2018) APO gene in rice improved grain yield per plant (Terao et al. 2010) APO gene in rice improved grain yield per plant (Terao et al. 2010) chloroplast division (Zhang et al. 2015), regulation of mitochondrial DNA replication as well as gene transcription and translation (Tang et al. 2019) chloroplast or mitochondria development (Quesada 2016) monoterpenone (C10) biosynthesis, resistance to the bacterial blight pathogen Xanthomonas oryzae pv. oryzae (Taniguchi et al. 2013) role in stress response (Brands and Ho, 2002), auxin transport to root tips (Janiak et al. 2017) transport of secretory proteins from the Golgi complex (Mo et al. 2007)
| Accession | Chromosome | NGK1 | NGK2 | BP1 | BP2 | E1 | Pearson | q | Gene Description | Functional Description |
|-----------|------------|------|------|-----|-----|-----|----------|----|------------------|---------------------------------------------------------------------|
| AX-94607905 | 2A | 571213772 | N0, (DSW); N120 (DRW, DSW) | 9.13E-07 | 0.199 | 0.001 | TraesCS2A02G337700 | 571150423 | 571149635 | non-specific lipid-transfer protein-like protein At2g13820 | predicted protein, lipid binding and transport, xylem differentiation (Motose et al. 2004, Kobayashi et al. 2011) |
| AX-94923560 | 2A | 729858414 | N60 (FRW, DRW, FSW, DSW); N120 (DRW, DSW) | 1.95E-06 | 0.128 | 0.001 | TraesCS2A02G501900 | 729858636 | 729852393 | acyl-coenzyme A thioesterase A | predicted protein, acyl-CoA hydrolase activity (Cheng et al. 2006); improves grain filling rate in rice (Zhao et al. 2019); lipid metabolism |
| AX-94705680 | 2B | 598802253 | N120 (DRW); N60 (FSW) | 1.29E-06 | 0.198 | 0.001 | TraesCS2B02G418100 | 598959096 | 598956372 | predicted protein | subtilisin-like protease SBT1.7 |
| AX-95203088 | 2B | 613322090 | N60 (FSW, DSW); N120 (DRW, DSW) | 1.72E-06 | 0.194 | 0.001 | TraesCS2B02G426600 | 613234917 | 613205484 | predicted protein | unnamed protein product |
| AX-94601746 | 2B | 743105753 | N60 (FSW), N120 (DRW) | 4.85E-07 | 0.203 | 0.001 | TraesCS2B02G546300 | 743331453 | 743328441 | predicted protein | Protein CutA 1, chloroplastic |
| AX-9487553 | 2D | 580238575 | N0 (FSW, DSW); N60 FRW, DRW, DSW); N120 (DRW, FSW) | 6.41E-07 | 0.137 | 0.001 | TraesCS2D02G479500 | 580238874 | 580235096 | predicted protein | Cadmium content and leaf margin trait (https://www.uniprot.org/uniprot/Q109R6), copper ion binding (Burkhead et al. 2003); signal transduction (Arnesano et al. 2003); nitrogen regulatory response in bacterial and eukaryotic chloroplast (Ninfa and Atkinson, 2000, Arcondéguy et al. 2001) |
| AX-94735141 | 2D | 581570385 | N60 (DRW, DSW, FSW); N120 (FRW, DRW, FSW, DSW) | 2.32E-06 | 0.127 | 0.001 | TraesCS2D02G480200 | 581526532 | 581523421 | predicted protein | anthocyanidin reductase activity and flavonoid biosynthetic process (Winkel-Shirley, 2001) |
| AX-94474729 | 2D | 584545802 | N60 (FRW, DRW) | 1.82E-06 | 0.13 | 0.001 | TraesCS2D02G481800 | 584545905 | 584542725 | predicted protein | Anthocyanin reductase |
| AX-94835810 | 2D | 584799948 | N0 (DRW, FSW, DSW); N60 (DRW, DSW); N120 (DRW, FSW, DSW) | 2.69E-06 | 0.175 | 0.002 | TraesCS2D02G482500 | 584804705 | 584799573 | predicted protein | tRNA modification, drought, salt and cold stress response, root and plant development in rice |
| AX-95223893 | 2D | 584861391 | N0 (FRW, DRW, FSW, DSW); N60 (DRW, DSW) | 3.55E-07 | 0.186 | 0.002 | TraesCS2D02G482800 | 584864531 | 584861189 | tRNA (guanine(10)-N2)-methyltransferase homolog | predicted protein, tRNA modification, drought, salt and cold stress response, root and plant development in rice |
| Accession | Chromosome | Start | End | Gene | Description |
|-----------|------------|-------|-----|------|-------------|
| AX-95003296 | 2D | 586331032 | 586332990 | unnamed protein product | regulation of cellular expansion and differentiation in Arabidopsis, ATP and carbohydrate binding, defence and signalling (Uniprot) |
| AX-94477325 | 2D | 586572446 | 586573324 | uncharacterized protein LOC109744903 | regulates rice seed length (Kitagawa et al. 2010); male meiosis, anther dehiscence, and fertility in rice (Zhou et al. 2011) |
| AX-95197137 | 2D | 586839201 | 586841990 | receptor-like serine/threonine-protein kinase SD1-8 | confers durable and broad-spectrum resistance to wheat powdery mildew (Cao et al. 2011); regulates stress-responsive gene expression in rice (Diédhiou et al. 2008), Negative regulator of immune responses in Arabidopsis (Lin et al. 2015) |
| AX-94525577 | 2D | 587107149 | 587108996 | predicted protein | confers durable and broad-spectrum resistance to wheat powdery mildew (Cao et al. 2011); regulates stress-responsive gene expression in rice (Diédhiou et al. 2008), Negative regulator of immune responses in Arabidopsis (Lin et al. 2015) |
| AX-94702180 | 2D | 587292781 | 587294568 | putative kinesin motor domain-containing protein | regulates Pi uptake by modulating PHT1;1 expression in Arabidopsis (Wang et al. 2014); age-triggered leaf senescence (Chen et al. 2017); Benzothiadiazole-inducible blast resistance (Shimono et al. 2007); resistance against *F. graminearum* in wheat (Bahrini et al. 2011); broad-spectrum resistance to wheat powdery mildew (Cao et al. 2011) |
| AX-94487982 | 2D | 588675894 | 588677693 | WRKY45-like transcription factor | regulates Pi uptake by modulating PHT1;1 expression in Arabidopsis (Wang et al. 2014); age-triggered leaf senescence (Chen et al. 2017); Benzothiadiazole-inducible blast resistance (Shimono et al. 2007); resistance against *F. graminearum* in wheat (Bahrini et al. 2011); broad-spectrum resistance to wheat powdery mildew (Cao et al. 2011) |
| AX-95190381 | 2D | 59102700 | 591038790 | serine-threonine protein kinase | regulates Pi uptake by modulating PHT1;1 expression in Arabidopsis (Wang et al. 2014); age-triggered leaf senescence (Chen et al. 2017); Benzothiadiazole-inducible blast resistance (Shimono et al. 2007); resistance against *F. graminearum* in wheat (Bahrini et al. 2011); broad-spectrum resistance to wheat powdery mildew (Cao et al. 2011) |
| Accession | Designation | Gene Symbol | Description |
|-----------|-------------|-------------|-------------|
| AX-95018936 | 2D 595159320  N60 (DRW); N120 (DRW, DSW) | TraesCS2D02G500500 | JmjC domain-containing protein regulation of RNA silencing, DNA methylation (Qian et al. 2019), Brassinosteroid (BR) signalling pathway, affecting flowering, and biorhythm and bud regeneration (Yokoo et al. 2014) |
| AX-94457170 | 2D 596252217 N0 (DSW); N120 (FSW) | TraesCS2D02G502700 | Adenine nucleotide alpha hydrolases-like superfamily protein hydrolase activity and root hair cell differentiation (https://www.uniprot.org/uniprot/Q84JS5); response to salt stress (Jung et al. 2015); involved in male sterility (Mok and Mok 2001) |
| AX-94962360 | 2D 596914793 N0 (DRW, FSW, DSW); N60 (FRW, DRW, DSW); N120 (FSW) | TraesCS2D02G503000 | Plasma membrane H+ ATPase Plant adaptation to environmental stresses (Janicka-Russak and Kabala, 2015), P deficiency and Al toxicity (Yu et al. 2015, Wang et al. 2014), transport of various nutrients (nitrate, phosphate and potassium) through roots, elongation of hypocotyls in Arabidopsis (Takahashi et al. 2012); NH4+ metabolism in rice roots (Weng et al. 2020); auxin-mediated cell elongation during wheat embryo development (Rober-Kleber 2003) |
| AX-94829391 | 2D 601212171 N0 (DSW); N60 (DRW) | TraesCS2D02G507800 | Nuclease S1 nucleic acid degradation during plant programmed cell death (Lesniewicz et al. 2013) |
| AX-94786006 | 2D 610277424 N60 (FRW, DRW, FSW); N120 (FRW) | TraesCS2D02G521400 | 3-oxoacyl-[acyl-carrier-protein] synthase III, chloroplastic fatty acid biosynthesis and metabolism, lipid biosynthesis and metabolism (https://www.uniprot.org/uniprot) |
AX-94695716  
2D  
702726797  
N60 (FSW, DSW); N120 (FRW, DRW, DSW)  
2.82E-06  
0.193  
0.002  
protein, role in rice root development (Ding et al. 2015)

AX-95136655  
3B  
234490336  
N0, N60, N120 (DTF)  
1.70E-06  
0.161  
0.016  
TraesCS3B02G201300  
233224384  
233224014  
protein, DEHYDRATION-INDUCED 19-like protein, drought tolerance in rice (Wang et al. 2014) and Arabidopsis through up-regulation of pathogenesis-related PR1, PR2, and PR5 gene expressions (Liu et al. 2013); response to salt and water stress (https://www.uniprot.org/uniprot/Q84J70), accelerate flowering (Hwang et al. 2019)

AX-95113687  
6A  
595578832  
N120 (RSA); N60 (Tips)  
3.04E-06  
0.256  
0.006  
TraesCS6A02G371000  
595564219  
595563589  
predicted protein, response to salt and water stress (Singh et al. 2017); growth and immune response in Arabidopsis (Yun et al. 2013)

AX-94513497  
6A  
595627899  
N0 (RSA, RV); N60 (RSA, RV, Tips); N120 (RSA, RV, Tips)  
1.89E-07  
0.311  
0.001  
TraesCS6A02G371300  
595628211  
595624936  
predicted protein, Protein transport, response to salt stress (https://www.uniprot.org/uniprot/Q9LFP1), tolerance to water stress (Singh et al. 2017); growth and immune response in Arabidopsis (Yun et al. 2013)

AX-94911804  
6A  
595776559  
N0 (RSA, RV); N60 (RSA, RV, Tips); N120 (RSA, RV, Tips, Forks)  
5.24E-07  
0.307  
0.002  
TraesCS6A02G371800  
595778154  
595774917  
vesicle-associated membrane protein 713

AX-94565231  
6D  
683646420  
N0 (RSA); N60 (RSA, TRL, RV, Tips); N120 (RSA, RV, Forks, Tips)  
8.41E-07  
0.305  
0.003  

AX-94676800  
7A  
376797697  
N60 (FSW); N120 (DRW)  
2.79E-06  
0.195  
0.001  
TraesCS7A02G293700  
376870968  
376869867  
calcium/calmodulin-regulated receptor-like kinase 2, response to cold, plant tolerance to salt and ABA stress (Yang et al. 2010)

p-value: significance level of marker-trait association; R²: percent phenotypic variance explained by the SNP; FDR: false discovery rate, p-value, R² and FDR is represented as the mean value across seasons
Table 4 The top 10 nitrogen insensitive (NIS) and 10 nitrogen sensitive (NS) breeding lines with contrasting grain yield (GY; kg ha\(^{-1}\)) derived from pooled mean over two seasons and three treatments.

| Category | Designation | Pop  | Mean GY | 2018_N0 | 2019_N0 | 2018_N60 | 2019_N60 | 2018_N120 | 2019_N120 |
|----------|-------------|------|---------|---------|---------|---------|---------|---------|---------|
| NIS      | HT661       | Pop1 | 3289    | 2865    | 2869    | 4093    | 3843    | 3190    | 2873    |
| NIS      | HT712       | Pop1 | 3237    | 3145    | 3568    | 3435    | 3458    | 2902    | 2915    |
| NIS      | HT722       | Pop1 | 3195    | 3102    | 3349    | 3158    | 3338    | 2892    | 3328    |
| NIS      | HT726       | Pop1 | 3172    | 2982    | 2502    | 3978    | 3522    | 2845    | 3203    |
| NIS      | HT723       | Pop1 | 2970    | 2802    | 2791    | 3207    | 2930    | 3222    | 2867    |
| NIS      | HT1913-2    | Pop4 | 2933    | 2332    | 2526    | 2909    | 2616    | 3693    | 3523    |
| NIS      | HT1870      | Pop4 | 2911    | 2122    | 2433    | 2632    | 2745    | 3665    | 3871    |
| NIS      | HT727       | Pop1 | 2911    | 2377    | 1833    | 3370    | 3012    | 3610    | 3266    |
| NIS      | HT1908-2    | Pop4 | 2900    | 2641    | 2148    | 2851    | 2368    | 3553    | 3837    |
| NIS      | HT764-1     | Pop2 | 2897    | 2440    | 2897    | 3528    | 2872    | 2747    | 2896    |
| NS       | HT1723      | Pop3 | 1885    | 1158    | 760     | 1806    | 1813    | 2675    | 3096    |
| NS       | HT696       | Pop1 | 1750    | 825     | 1261    | 1660    | 1950    | 2455    | 2350    |
| NS       | HT704       | Pop1 | 1661    | 827     | 1245    | 1948    | 1274    | 2297    | 2372    |
| NS       | HT644       | Pop1 | 1658    | 989     | 1384    | 1838    | 1474    | 1822    | 2443    |
| NS       | HT845       | Pop2 | 1625    | 1111    | 1232    | 1700    | 1793    | 2038    | 1878    |
| NS       | HT1882-3    | Pop4 | 1594    | 918     | 1186    | 1256    | 1646    | 1982    | 2576    |
| NS       | HT647       | Pop1 | 1590    | 846     | 985     | 1082    | 1535    | 2440    | 2652    |
| NS       | HT765       | Pop2 | 1519    | 768     | 1162    | 1375    | 1606    | 1720    | 2484    |
| NS       | HT665       | Pop1 | 1413    | 542     | 1010    | 1187    | 1077    | 2370    | 2292    |
| NS       | HT847       | Pop2 | 1359    | 683     | 717     | 1258    | 1333    | 1980    | 2182    |
**Fig 1.** Schematic representation of the breeding strategy used to develop the nested synthetic wheat introgression libraries.
Fig 2. Plots of Pearson’s r-values showing the correlation among traits measured (A) at N0, (B) N60 and (C) N120 level. The blue colour indicates positive correlation and red colour indicated the negative correlation among different traits, the variation in colour intensity is representing the strength of the correlation among the traits. *significance at <5% level, **significance at <1% level, ***significance at <0.1% level.
Fig. 3 (A) Population structure within the nested synthetic wheat introgression libraries. The population structure plots with each vertical bar representing a breeding line coloured according to the particular group to which the breeding line has been assigned. The breeding lines assigned to more than one of group represents the degree of their admixed set of the alleles. (B) The Kinship matrix displayed as the heat map, where the red indicates the highest correlation between the pairs of breeding lines and yellow indicates the lowest correlation. (C) The Scree plot indicating the most of the variability explained by first three PCs for association study. (D) The three-dimensional view of the principal components explaining the genotypic variation among breeding lines constituting the introgression libraries. (E) The appropriate number of the sub-populations determined from the largest delta K=3
Fig. 4 Manhattan plot and qq plot for the yield and yield related traits across seasons at three differential level of nitrogen (N0, N60, N120) (A) grain yield (GY) (B) Days to 50% flowering (DTF) (C) shoot biomass (SB) (D) spikelets per spike (SPS) and (E) number of productive tillers
Fig 5 Schematic representation of the SNP distribution along the 21 chromosomes of wheat. The chromosome map showing genomic regions where MTAs for different NUE related trait, root traits, yield and yield related traits. The numbers below each chromosome indicate chromosome numbers. The bp representing the physical position of the SNPs on the chromosome in base pair.

**Fig 5** Schematic representation of the SNP distribution along the 21 chromosomes of wheat. The chromosome map showing genomic regions where MTAs for different NUE related trait, root traits, yield and yield related traits. The numbers below each chromosome indicate chromosome numbers. The bp representing the physical position of the SNPs on the chromosome in base pair.
Fig. 6 GGE biplot showing the performance of 352 nested synthetic wheat introgression lines across seasons and treatments (N0, N60, N120).

The environment view refers to the three-differential level of nitrogen application: N0, N60 and N120. The genotype view refers to the 352 nested synthetic wheat introgression lines. The numeric number refers to the coding for the introgression lines, which is given in detail in Supplementary table S7.
Fig. 7: The grain yield performance of top 20 breeding lines derived from the nested introgression libraries possessing high and stable grain yield (GY; kg ha\(^{-1}\)) across two seasons and three treatments. The numeric values above the bar graph indicate the mean grain yield (GY; kg ha\(^{-1}\)) performance of breeding lines across seasons.
Fig. 8 The allelic constitution of the selected promising breeding lines, wild accessions of *Ae. tauschii*, cultivated and synthetic wheats for the (A) root related traits and (B) grain yield.
**Figure 1**

Schematic representation of the breeding strategy used to develop the nested synthetic wheat introgression libraries.
Figure 2

Plots of Pearson's r-values showing the correlation among traits measured (A) at N0, (B) N60 and (C) N120 level. The blue colour indicates positive correlation and red colour indicated the negative correlation among different traits, the variation in colour intensity is representing the strength of the correlation among the traits. *significance at <5% level, **significance at <1% level, ***significance at <0.1% level.
Figure 3

(A) Population structure within the nested synthetic wheat introgression libraries. The population structure plots with each vertical bar representing a breeding line coloured according to the particular group to which the breeding line has been assigned. The breeding lines assigned to more than one of group represents the degree of their admixed set of the alleles. (B) The Kinship matrix displayed as the heat map, where the red indicates the highest correlation between the pairs of breeding lines and yellow indicates the lowest correlation. (C) The Scree plot indicating the most of the variability explained by first three PCs for association study. (D) The three-dimensional view of the principal components explaining the genotypic variation among breeding lines constituting the introgression libraries (E) The appropriate number of the sub-populations determined from the largest delta K=3
Manhattan plot and qq plot for the yield and yield related traits across seasons at three differential level of nitrogen (N0, N60, N120) (A) grain yield (GY) (B) Days to 50% flowering (DTF) (C) shoot biomass (SB) (D) spikelets per spike (SPS) and (E) number of productive tillers.

Figure 4
Figure 5

Schematic representation of the SNP distribution along the 21 chromosomes of wheat. The chromosome map showing genomic regions where MTAs for different NUE related trait, root traits, yield and yield related traits. The numbers below each chromosome indicate chromosome numbers. The bp representing the physical position of the SNPs on the chromosome in base pair.
Figure 6

GGE biplot showing the performance of 352 nested synthetic wheat introgression lines across seasons and treatments (N0, N60, N120). The environment view refers to the three-differential level of nitrogen application: N0, N60 and N120. The genotype view refers to the 352 nested synthetic wheat introgression lines. The numeric number refers to the coding for the introgression lines, which is given in detail in Supplementary table S7.

Figure 7
The grain yield performance of top 20 breeding lines derived from the nested introgression libraries possessing high and stable grain yield (GY; kg ha-1) across two seasons and three treatments. The numeric values above the bar graph indicate the mean grain yield (GY; kg ha-1) performance of breeding lines across seasons.

Figure 8

The allelic constitution of the selected promising breeding lines, wild accessions of Ae. tauschii, cultivated and synthetic wheats for the (A) root related traits and (B) grain yield.

Supplementary Files
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- SupplementaryTablesandFigures01062021.xlsx