Detection of stable QTLs for grain protein content in rice (*Oryza sativa* L.) employing high throughput phenotyping and genotyping platforms

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Lack of appropriate donors, non-utilization of high throughput phenotyping and genotyping platforms with high genotype × environment interaction restrained identification of robust QTLs for grain protein content (GPC) in rice. In the present investigation a BC$_3$F$_4$ mapping population was developed using grain protein donor, ARC10075 and high-yielding cultivar Naveen and 190 lines were genotyped using 40 K Affimetrix custom SNP array with the objective to identify stable QTLs for protein content. Three of the identified QTLs, one for GPC ($qGPC1.1$) and the other two for single grain protein content ($qSGPC2.1$, $qSGPC7.1$) were stable over the environments explaining 13%, 14% and 7.8% of the phenotypic variances, respectively. Stability and repeatability of these additive QTLs were supported by the synergistic additive effects of multi-environmental-QTLs. One epistatic-QTL, independent of the main effect QTL was detected over the environment for SGPC. A few functional genes governing seed storage protein were hypothesised inside these identified QTLs. The $qGPC1.1$ was validated by NIR spectroscopy-based high throughput phenotyping in BC$_3$F$_5$ population. Higher glutelin content was estimated in high-protein lines with the introgression of $qGPC1.1$ in telomeric region of short arm of chromosome 1. This was supported by the postulation of probable candidate gene inside this QTL region encoding glutelin family proteins.

Malnutrition is responsible for about 24,000 deaths per day worldwide. Rice is staple food for more than half of the world population. It has a significant contribution in daily calorie-intake as millions of poor families depend mainly of rice for their nutrition. Rice supplies abundant carbohydrate as its kernel constitutes mainly of starch (>80%) but protein (7–8%) is the source of concern. However, the protein quality measured by protein digestibility index and amino acid composition is the best among cereals, which makes it preferable for the food and feed industries. Efforts were made during past three decades by rice breeders to improve the protein content in rice grain, but significant and stable improvement could not be achieved due to the involvement of many small effect genes/quantitative trait loci (QTLs) substantially affected by environment. The QTLs for grain protein content (GPC) in rice have been identified in almost all chromosomes, though majority of them are present on chromosomes 1, 2, 6, 7, 10 and 11. But multi-environmental stable and robust QTL for this trait was rare. This was due to the lack of high throughput genotyping platform leading to low density linkage map, low population size, lack of high throughput phenotyping procedure and lack of validation in different cropping season and environments. Moreover, this trait is not only governed by additive gene effect but also significantly influenced by the
complex gene interaction including dominance, epistatic and genotype × environment interaction (G E I) component effects as realized by many researchers. But, in spite of quite high probability of getting epistasis and GEI-QTLs, no notable epistatic or multi-environment trial QTL (MET-QTL) was detected in rice for this trait. With the recent advancements in rice genomics research, more robust and reproducible markers such as single nucleotide polymorphic (SNPs) markers have been utilized to make SNP chips of various magnitude, i.e. on medium density Illumina’s rice platform, high density 50 K Illumina Infinium array platform (RiceSNP50) and Affymetrix custom array such as 44 K and 50 K SNP chips platform in rice. In addition, Near Infrared (NIR) spectroscopy has been used by researchers to screen large number of germplasm for protein content in several cereals and in high throughput phenotyping of breeding lines.

In bi-parental mapping, population for detection of robust QTL for a particular trait required significant differences of two parents for that trait. For detecting QTLs for GPC, rarely very high protein genotype and low protein counterpart had been used which restricted trait variability and availability of robust QTL. Several rice germplasm with high GPC have been identified over the environments. They however were low yielder and had many undesirable features. Backcross breeding could be an effective approach for minimizing the undesirable effects coming from un-adapted donor parents. Backcross population is not only useful for detecting robust QTLs but also to generate introgression lines for use as pre-breeding lines or as high yielding elite cultivars. The advanced backcross QTL (AB-QTL) analysis has been successfully employed in detecting and transferring QTLs from un-adapted germplasm into advanced breeding lines in many plant species. In rice, AB-QTL analysis has helped to detect many QTLs for several grain quality traits. But the use of two diverse parents (regard to origin, nature, type and adaptability) often poses many problems such as lack of proper chromosomal pairing, pollen sterility in backcross lines leading to segregation distortion (SD) etc. Zhan and Xu suggested that being the potential evolutionary force, the SD loci should be effectively utilized in mapping genes using appropriate packages. Among the statistical packages utilized for mapping QTLs, a SAS-based programme Proc QTL, QTL IciMapping V4 and DistortedMap handle SD markers safely and effectively to identify regions influencing trait expression. Inside the putative or multi-environment QTLs region, functional genes which ultimately governed the phenotype were found using bioinformatics tool in previous studies on rice. In the present study high genetic variability governed by high protein donor followed by high throughput SNP-array based genotyping were exercised with the aim of detection of robust QTLs for grain protein content with plausible influence of epistasis and genotype × environment interaction. This investigation also explored the scope of high throughput phenotyping using NIR spectroscopy to validate stable QTLs in advanced near isogenic line (NIL) population. Finally it focused on the delineation of QTLs loci to find functional genes inside QTLs and tried to associate them with higher protein and protein fraction content in the selected stable high protein introgressed (NILs) over the environments.

Results and Discussion

Phenotypic analysis. ANOVA for plant height (cm) (PH), maturity duration (MD), number of panicles/plant (PN), panicle length (cm) (PL), grains/panicle (GRAIN), 100 grain weight (g) (GWT), plant yield (g) (PY), grain protein content (%) (GPC), single grain protein content (mg/g) (SGPC) in both kharif 2013 (Env.1) and rabi season 2014 (Env.2) individually and over the seasons (Env.1 + Env.2) revealed the significant variation in population for all the traits (Supplementary Table 1). High heritability (h² = 0.75–0.78) of GPC in individual environment was observed. But this was moderate to low (h² = 0.45) across environments calculated from pooled data. In contrary, SGPC revealed relatively higher heritability (h² = 0.55) over the environments (Supplementary Table 1). Moreover, higher phenotypic variance of SGPC also indicated its suitability for QTL analysis. These facts indicated that SGPC was three environmentally more stable than the GPC trait and therefore, transfer of this trait could be more feasible. Except PY and GWT all other traits followed normal distribution and both absolute values of skewness and kurtosis were less than 1.0, indicating suitability of data for QTL analysis (Supplementary Table 4). Comparative lower (p < 0.01) negatively associated with PY in two seasons and over the environments. Path coefficient analysis (Supplementary Table 4) also revealed most significant direct effect of PN and GRAIN on PY, while no significant effect of GPC and SGPC was observed on PY.

Analysis of variance revealed significant differences (p < 0.001) of genotypes, environment and genotype × environment interaction for grain protein content (GPC) with nearly similar trend for single grain protein content (SGPC) in genotype (G) and environment (E) (p < 0.001) as well as G x E (p < 0.01). The significantly higher (p < 0.001) mean GPC of mapping population was observed in rabi season 2014 (Env.2) as compared to both the kharif seasons (Env.1 and Env.3). Comparative lower (p < 0.001) average SGPC was also found in Env.1 than in the Env. 2. Better water and nutrient management and higher light intensity in rabi season might have contributed to better grain filling and protein content in rice. ARC10075 had higher GPC and SGPC values than the control. Hence, ARC10075 and environment Env.2 were considered as reference combinations for identifying the best genotype in any specific environment. Lines, PLN-32, PLN-64, PLN-58 and PLN-56 in Env.2 were found superior in GPC while PLN-64 was also found superior in SGPC in Env. 2. Interaction plots and ANOVA suggested that the genotype × environment interaction effects were significant (p < 0.01) for both GPC and SGPC. The trend lines (Fig. 2) also showed that for both GPC and SGPC, all three environments were not parallel. Therefore, the presence of genotype × environment interaction effect was obvious.
Distribution of sNps in chip, genotyping and linkage analysis. Among the four types of genes used for the 40,894 SNP chip designing, majority (96.6%) were single copy (SC) genes. The rest were from agronomically important cloned rice genes (AGCR) (2.27%) and multi-copy rice (MCR) (1.14%) genes. Further, 21100 (51.6%) single copy genes were unique to rice (SCR) and 18397 (45%) conserved single-copy genes were common to wheat and rice (CSCWR) (Fig. 3a). This SNP chip had 38% SNPs from exons, 42% from introns and 20% from 5′ and 3′ UTR regions (Fig. 3b). The SNPs from exon regions could be further classified into non-synonymous (20% of total SNPs) and synonymous (18% of total SNPs) types. The non-synonymous SNPs are important for detection of probable functional genes for the trait concerned. The presence of large number of these SNPs, makes this chip more effective for associating genotypes with the desired phenotype, i.e. high protein content. Overall, the SNPs were distributed among all 12 rice chromosomes with an average of one SNP per 9.54 kb (Fig. 3c). The number of SNPs varied from 983 (chromosome 10) to 8428 (chromosome 1) with an average of 3407.83 per chromosome.

All the genotypes passed the development quality check (DQC) with a high cut-off value of > 0.82, and the maximum DQC value was 0.99. Except for one sample, all others possessed high genotyping call rates of > 95%, with an average of 99.9%. Out of the 40894 markers in SNP chips, 5492 SNPs accounting 13.43% were found to be homo-polymorphic between ARC10075 and Naveen. These markers were highly informative as many of them were located inside genes. The remaining 82.96% (33925) and 1.66% (680) were non-polymorphic and hetero-polymorphic, respectively. Only 506 SNPs accounting 1.24% were not detected in this assay (GAP) (Supplementary Table 5). GTC Software efficiently separated homozygous and heterozygous cluster (Supplementary Fig. 1). The proportion of genome of Naveen in backcross derived lines varied from 46.88 to 95.62% with an average of 81.8% based on homo-polymorphic SNP markers. The proportion of genome of the donor ARC10075 varied from 2.97 to 40.37% (average 13.06%). The rest genome with an average of 5.12% was heterozygote among backcross derived lines. Out of the homo-polymorphic markers between the parents, 87%
showed segregation distortion (SD) through \( \chi^2 \) test \((p > 0.01)\) and were distributed on all the 12 chromosomes. Segregation distorted markers occurred due to unwanted selection pressure imparted by pollen sterility, incompatibility, epistatic and environmental interaction, etc. We employed one accession (ARC10075) as donor for high GPC which was collected from North-eastern part of India, which is considered as the secondary centre of origin for rice. Assam rice collection (ARC) represents diversity of this region. Some of the germplasm belonging to the part of the country adjoining Myanmar, China, and Indonesia have many traits intermediate to those of indica and japonica. Therefore, chances of having sterility were high as observed frequently in inter sub-specific crosses, especially in backcross progenies which led to segregation distortion (SD). In general, distorted markers did not have much effect on the position and effect estimations of QTL; moreover, their effects can be ignored in large-size mapping populations \(^{40,41}\). In the previous studies \(^{4,41}\) large number of markers (40–55%) showing SD were successfully utilized to map grain quality traits. In traditional linkage mapping, there is all likelihood of losing all these informative markers in QTL analysis. In the present study, we handled SD markers along with non-SD markers using ‘SDL mapping’ in the QTL IciMapping V4 software which helped in restoration of order and position of the distorted markers to safely use in QTL detection. By employing these options of mapping, we could use all available markers, whether Mendelian or otherwise and could save valuable resources. A high density linkage map with 12 linkage groups on 12 rice chromosomes was generated. The average genetic to physical distance of 1 cM = 0.2 Mb. The total map distance was 2480 cM with an average 0.46 cM marker-interval. Through DistortedMap v.1 software, it was found that all markers were qualified for SD mapping analysis. Although little higher average (0.67 cM) marker distance was noticed which could be the effect of epistatic SD locus.

**Single environment QTLs.** A total of 14 main effect additive single environment QTLs for GPC and SGPC were detected by inclusive composite interval mapping (ICIM). Three of them were found in more than one environment (Table 1). Compared to GPC, more number of additive QTLs were detected for SGPC in single and multi-environment. Previously also researcher\(^ {46}\) did not find any consistent environmentally stable QTLs for GPC, but detected stable QTLs for protein index (PI) which was almost identical with SGPC, used in the present study. In kharif season 2013 (Env.1), one QTL (qGPC1.1) at 11 cM position was identified for GPC on chromosome 1 with a logarithm of odds ratio (LOD) value of 3.83 which explained 13.86% phenotypic variance. In this environment, four other QTLs for SGPC were identified. One of the pleotropic QTL (qSGPC1.1) shared the same position with qGPC1.1 explaining 10.37% phenotypic variance with a LOD value of 2.9. The other three QTLs for SGPC (qSGPC2.1, qSGPC7.1, qSGPC11.1) had LOD values of 3.32, 3.31 and 2.88 with 6.7%, 7.68% and 6.42% phenotypic variance explained (PVE), respectively. In rabi season 2014 (Env.2), still higher number of QTLs for
both GPC and SGPC were detected. Of the two QTLs for GPC, one (qGPC1.1) was common with the previous environment (Env.1) explaining 13.35% phenotypic variance with LOD value of 4.02. The new putative QTL (qGPC 2.1) was detected at 170 cM position on chromosome 2 which had 17.35% PVE with LOD value of 3.19. Eleven QTLs for SGPC were found in Env.2 on chromosomes 1, 2, 3, 7, 8, and 12. Two of them were common with previous environment (Env.1). They were qSGPC2.1 and qSGPC7.1 with LOD values of 3.53 and 3.33, respectively which explained 14.64% and 7.81% phenotypic variance (Fig. 4).

Apart from GPC and SGPC, QTLs were detected for other traits such as panicle length (PL), panicle number/plant (PN), grains/panicle (GRAIN) which were normally distributed. In Env.1, 15 putative QTLs (PL-1, PN-13, GRAIN-1) distributed among chromosomes, 1, 2, 4, 5, 8, 9, 10, 11 and 12 explaining 6% to 32.5% phenotypic variance (PVE) (Supplementary Table 6) and in Env.2, 5 putative QTLs (PL-3, GRAIN-2) distributed in chromosomes, 1, 2, 6 and 7 with 6.1% to 24.67% PVE (Supplementary Table 7) were detected. But none of them was detected over the environments. Simple interval mapping which is based on maximum likelihood may not be as efficient as ICIM, but it can provide information on small effect QTLs independent from variance of other QTLs. IM was used to identify QTLs for GPC and SGPC in the present investigation. Multi-environmental consistent QTLs such as qGPC1.1 and qSGPC2.1 which were identified by ICIM, were also found in interval mapping. Position of another consistent QTL, qSGPC7.1 was little shifted in this analysis. Apart from them single environment putative QTLs, qSGPC1.1, qSGPC11.1, qSGPC1.2, qSGPC1.3, qSGPC3.1, qSGPC7.2, qSGPC8.1, qSGPC8.2,
qSGPC12.1 which were detected by ICIM, were also found through IM. In addition 14 other putative QTLs on chromosome 1, 3, 4, 8, and 11 were found by this analysis (Supplementary Table 8).

Epistatic QTLs and MET-QTLs for GPC and SGPC. The epistatic interaction can not be ignored because such attempt may lead to underestimation of total genetic effects of a trait. The proper detection of the direction of epistatic interaction, i.e. synergistic or non-synergistic effect on other QTLs can guide the breeder to introgress multi-QTLs for one or many traits. Epistatic interaction for grain protein content was reported in other cereals43,46, such attempt may lead to underestimation of total genetic effects of a trait. The proper detection of the direction of epistatic interaction, i.e. synergistic or non-synergistic effect on other QTLs can guide the breeder to introgress multi-QTLs for one or many traits. Epistatic interaction for grain protein content was reported in other cereals43,46.

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have GEI effect, they can be safely used in the molecular breeding programme. On the other hand, it also indicated that all these MET-QTLs had significant positive effect on positive allele which improved the phenotypic expression leading to higher GPC and SGPC in rice in favourable environments. Finally the epistasis × environment interaction effect (aae) was an important component of QTL × environment (QE) interaction effects. In MET analysis 6 pair (Supplementary Table 9) and 48 pair (Supplementary Table 12) of QTLs associated with GPC (Fig. 5b) and SGPC, respectively, were found with epistatic effects (aa) and epistasis × environment (aae) effects. But none of them was found associated with the main effect QTLs. One main epistatic QTL was adjacent to the

Figure 4. QTLs detected for SGPC in rabi season (Env.2) showing common QTLs qSGPC2.1 and qSGPC7.1 found in also in kharif season (Env.1).

Figure 5. (a) Cyclic diagram of epistatic QTLs (ep-QTL) for single grain protein content (SGPC) in rabi season 2014 (Env.2) and (b) epistatic × environmental (aae) interaction QTLs for grain protein content (GPC). The dotted lines indicate the interacting SNP marker pairs situated on the same or different chromosomes with corresponding LOD scores owing to their epistatic effects. Marker position (cM) is mentioned inside the oval located on chromosome. One common ep-QTL in kharif season 2013 (Env.1) is located between 6 cM position on chromosome 11 and 255 cM position on chromosome 1 (a).
Spectroscopy-based high throughput non-destructive protein estimation method in kharif types in segregating generations for grain protein. We attempted QTL mapping for GPC using near infrared (NIR) sampling and tedious analysis procedure is a major bottleneck for mapping QTLs and selecting superior genotypes for GPC and also established the NIR spectroscopy as the valid high throughput phenotyping tool for detection of stable QTL ($qGPC1$). The GPC varied from 6.56% to 12.89% with mean value of 9.33%. We detected the same from Naveen/ARC10075. NIL population showed the normal distribution with positive skewness for apparent GPC similar phenotypic variance (12.18%) as observed in $Env$. Validation of main effect QTL for GPC using high throughput phenotyping. The need of destructive sampling and tedious analysis procedure is a major bottleneck for mapping QTLs and selecting superior genotypes in segregating generations for grain protein. We attempted QTL mapping for GPC using near infrared (NIR) spectroscopy-based high throughput non-destructive protein estimation method in $kharif$ 2014 ($Env$).3. Calibrated NIR spectroscopy for GPC was used to determine the apparent grain protein content in NILs ($BC_{F_2}$) derived from Naveen/ARC10075. NIL population showed the normal distribution with positive skewness for apparent GPC (Supplementary Fig. 2). The GPC varied from 6.56% to 12.89% with mean value of 9.33%. We detected the same QTL ($qGPC1$) as was found through phenotyping by conventional micro-Kjeldahl method. It also explained almost similar phenotypic variance (12.18%) as observed in $Env$.1 and $Env$.2 (13.85%). This observation validated QTL for GPC and also established the NIR spectroscopy as the valid high throughput phenotyping tool for detection of stable QTL for GPC in rice.

Delineation of QTL loci and identification of probable candidate genes. Normal distribution for GPC and SGPC indicated the involvement of many QTLs for grain protein in rice. However, some regulatory genes were reported to be involved in controlling GPC in seeds in many cereal crops such as barley, wheat and rice. GWAS and candidate gene based association study identified a gene $HVNAM$ controlling GPC in barley. In what also, a high grain protein gene, $Gpc-B1$ was introgressed to improve protein content without affecting grain yield. In the present experiment, the functional genes presents inside and adjacent to the QTLs were identified (Table 3). One main effect additive QTL $qGPC1.1$ was found stable over the environments. The peak SNP marker inside this QTL (locus- 1:611041-111041) in $Env$.1 was CSCWR_Os01g02590__61041 which is located inside a conserved single-copy gene common to wheat and rice. This QTL interval region corresponded to a span of 186 $O. sativa$ $Japonica$ genes starting from Os01g0111600 to Os01g0119500. This QTL interval was narrow (locus- 1:811041-911041) in $Env$.2 corresponding only 34 coding and non-coding genes staring from Os01g0115100 to Os01g0116000 with same peak locus as was detected in $Env$.1. Multi environmental (MET) QTLs, $Eq-GPC1.1$ and $Eq-SGPC1.1$ were pleotropic (locus- 1:811041-911041) inside the main effect robust QTL, $qGPC1.1$. Among genes located inside these QTLs one gene $Os01g0111900$ (locus- 1:625986-627009) encoded glutelin family protein. This gene was located just 0.93 cM apart from the QTL peak. Inside one putative single environment QTL, $qGPC1.3$ (locus- 1:3737716–3837716) encoding seed storage protein (pro-glutelin content in seed, floury endosperm) was located just 0.23 cM apart from the QTL peak (Q-TARO annotation). Adjacent to one putative QTL, $qGPC1.2$ (locus- 1:39016361–39116361), around 0.83 Mb upstream region, $OsAAP6$ gene was present which was amino acid transporter enhancing GPC. On chromosome 2, one putative QTL, $qGPC2.1$ (locus- 2:9476316–10476316) and in the MET-QTL $Eq-GPC2.1$ with relatively narrow interval (locus- 2:9926316–10026316) contained two genes $Os02g0268100$ and $Os02g0268300$ which were 1.5 cM apart from the QTL peak. They also encoded glutelin protein. A gene cluster encoding glutelin fragment proteins and prolamin box binding factors is also found near to it. On chromosome 3, inside one putative QTL $qSGPC3.1$ (locus- 3:33532659–37322659) and MET-QTL $Eq-GPC3$ (locus- 3:35228948–35328948) one gene $Os03g0826500$ encoded anthranilate synthase alpha 1 related to higher grain protein content (Q-TARO annotation). This was located 3.01 cM and 1.1 cM apart from the peak of main QTL and MET-QTL, respectively. On chromosome 11, one putative QTL, $qSGPC11.1$ and two MET-QTLS, $Eq-qSGPC11.1$ and $Eq-GPC11.1$ (locus- 11:3737716–3837716) had the peak marker SCR200_Os11g07480_87716. One gene $OsAsp1$ influencing seed protein synthesis was located 0.62 Mb downstream of this QTL peak.

Analogy with previous findings on QTLs for grain protein in rice. A few QTLs identified in present study were located near or inside the QTLs and genes for GPC reported earlier. The main effect additive QTL, $qSGPC1.3$ was located near to $qPr1$ at 12 Mb region on chromosome 1. On the same chromosome, another...
QTL, qSGPC1.2 was identified at 39.07 Mb position which was very near to a reported QTL qPC1.5 and cloned gene (OsAAP6) inside this QTL at 38.13 Mb region. One epistatic QTL, ep-qSGPC-1 which was identified over the season was also located adjacent to qPC-1 at 24.39 Mb region with 10.5% phenotypic variance explained (PVE)13. On chromosome 2, one MET-QTL, Eq-SGPC2.2 was located adjacent to a main effect QTL qPC2 for grain protein content4,5,6. On the same chromosome, one QTL qSGPC2.1 identified in more than one environments and in MET analysis was detected near another major QTL qPro-2 explaining 41% PVE for grain protein at 4.3 Mb position10. On chromosome 3 only one main effect QTL qSGPC3.1 detected in the present study was located near to a reported QTL, qPC-3.3,4. On chromosome 5, 1.35 Mb downstream of one stable QTL qSGPC7.1 and MET-QTL Eq-SGPC7.1 (locus- 7:22197522–22297522) three gene cluster (Os07g0570100, Os07g0570300, Os07g0570500) encoding peptidase protein was located13. A few other QTLs (qCP7, qPr7, PC7) reported earlier3,4,8 were also located near the present QTL qSGPC 7.1. Another putative QTL, qSGPC8.1 on chromosome 8 was located just adjacent to a QTL for grain protein content qPro-8 at 1.2 Mb position11. We found another putative QTL qSGPC11.1 and MET-QTL on the same position which was located very near to a QTL, qGPC-11 detected through association mapping at 4.3 Mb position with peak SNP located on gene OsAsp1.33.

### High protein elite NILs: their significance in mapping and validation of robust QTL

Seven high yielding introgression lines (BC$_1$F$_2$) were selected for high GPC and phenotypic resemblance with Naveen. They had comparable maturity duration (121–127 days) and plant height (108–115 cm) with Naveen (124 days, 113 cm). All lines had significantly higher GPC and SGPC in both rabi and kharif 2015 (Env-4 and Env-5). The average protein yields of these lines were also higher than those of their high yielding parent (Supplementary Table 13). These selected lines along with another three high protein lines without phenotypic resemblance with Naveen were analysed for genomic composition. They had 81–87% genome from Naveen and 10.7–16.7% from ARC10075. GGT selection lines along with another three high protein lines without phenotypic resemblance with Naveen were analysed for genomic composition. They had 81–87% genome from Naveen and 10.7–16.7% from ARC10075. GGT analysis revealed that except for two all the selected high protein lines had the genomic region with the stable QTL qGPC1.1 in telomeric region (~0.8 Mb) of short arm of chromosome 1 (Fig. 7). Gene (Os01g0111900) present in this region (locus- 1:625986–627709) synthesized glutelin protein. Significantly, except two, all high protein lines had higher (p > 0.01) glutelin content than the recurrent parent, Naveen. Glutelin contains essential amino acids like glutamic acid, aspartic acid, and proline which are essential for the growth and development of plants.
lysine and is the major constituent of protein body II, which is more digestible than protein body I, which contains mostly prolamin54. Therefore, higher accumulation of glutelin ensures better protein quality in most of these lines. It was reported earlier that improvement of grain protein content reduced the protein quality and resulted in hardening of the cooked rice grains due to increase in prolamin fraction. The ratio of prolamin to glutelin fractions ranged mostly prolamins54. Therefore, higher accumulation of glutelin ensures better protein quality in most of these lines.

Table 3. Predicted functional genes present inside and adjacent to the main, epistatic and MET-QTLs and their distances from the QTLs peak.

| Trait/QTL | Environment | chromosome | Peak marker | Position (Mb) | Number of genes in QTL interval | Starting - ending gene at QTL interval | Nearest functional gene (RAP DB ID/Q-TARO ID) | Gene function | Gene Position (Mb) | Distance (gene-QTL) (Mb) |
|-----------|-------------|------------|-------------|--------------|-------------------------------|---------------------------------------|---------------------------------|---------------|-------------------|------------------------|
| qGPC1.1 | Env.1, Env.2, Env.3, MET | 1 | CSCWR, Os01g02950_61041 | 0.861041 | 186 | Os01g0111600-Os01g0119500 | Os01g0111900 | Glutelin family protein. (Os01i0111900-01) | 0.6264975 | 0.23454 |
| qSGPC1.1 | Env.1 | 2 | SCR200, Os02g17350_76316 | 9.976316 | 251 | Os02g02685700-Os02g02812000 | Os02g0268100 | Similar to Glutelin (Fragment). (Os02i0268100-01) | 9.5814845 | 0.394 |
| qSGPC7.1 | Env.1, Env.2, MET | 7 | SCR100, Os07g57440_17971 | 22.24752 | 28 | Os07g0556500-Os07g0558500 | Os07g0570100, Os07g0570300, Os07g0570500 | Gene cluster of three peptidase proteins | 23.614 | 1.35 |
| qSGPC11.1 | Env.1, MET | 11 | SCR200, Os11g07480_87716 | 3.788 | 35 | Os11g0175300-Os11g0177200 | Os11g0184800 (OsAsp1) | OsAsp1 | 4.34 | 0.62 |
| ep-qSGPC11.1 Epistatic in Env.1 and Env.2 | 11 | SCR100, Os11g08270_33249 | 4.333 | — | — | Os11g0184800 (OsAsp1) | OsAsp1 | 4.34 | 0.01 |
| qGPC2.1 | Env.2 | 2 | SCR200, Os02g17350_76316 | 9.976316 | 251 | Os02g02685700-Os02g02812000 | Os02g0268100 | Similar to Glutelin (Fragment). (Os02i0268100-01) | 9.5814845 | 0.394 |
| Eq-GPC2.1 | MET | 2 | SCR200, Os02g17350_76316 | 9.976316 | 251 | Os02g0268100 | Os02g0272900-Os02g0274100 | Similar to Glutelin (Fragment). (Os02i0268100-01) | 9.5814845 | 0.394 |
| qSGPC1.2 | Env.2 | 1 | SCR100, Os01g066690_66361 | 39.06636 | 28 | Os01g0897700-Os01g0899100 | Os01g0878700 (OsAAP6) | Amino acid transporter, transmembrane domain containing protein | 39.89 | 0.83 |
| qSGPC1.3 | Env.2 | 1 | CSCWR, Os01g14920_57875 | 8.357875 | 36 | Os01g0251400-Os01g0253800 | Os01g0254000 (Sar1c) | Seed storage protein. Pro-gluandin content in seed. Floursy endosperm. | 8.41 | 0.06 |
| qSGPC3.1 | Env.2 | 3 | CSCWR, Os03g04360_22659 | 36.32266 | 383 | Os03g0840200-Os03g0862200 | Os03g0826500 | Anthranilate synthase alpha 1 | 35.57 | 0.75 |
| Eq-GPC3.1 MET | 3 | SCR200, Os01g02348_78948 | 35.28 | 32 | Os03g0838400-Os03g0826500 | Os03g0826500 | Anthranilate synthase alpha 1 | 35.57 | 0.29 |

Expression profile of the functional gene located within QTL loci. Most of the probable functional genes (Table 3) inside the QTLs showed up-regulation in seed as a whole, aleurone layer, panicle tissues and root (Supplementary Fig. 3) based on RNA-seq data in Rice expression database (RED). Using RiceXPro database (RXP_0012) an expression heat map was generated to compare the gene expression profile of the 11 probable functional genes located inside QTLs in embryonic and endosperm-specific tissues at 7-, 10-, 14-, 21-, 28- and 42- days after flowering (DAF), respectively. The heat map (Fig. 8) clearly demonstrated the up-regulation of majority of the genes, including genes for enhancing storage proteins viz. glutelin and prolamin, preferentially at endosperm in all the time-points considered under study. Preferential up-regulation of functional genes for high GPC during endosperm development suggests higher accumulation of total protein in the selected introgression lines listed in Table 4.

Conclusion
We found that had we discarded SD markers, the stable QTLS like qGPC1.1 or qSGPC2.1 and qSGPC7.1 and their MET-QTLS would have remained undetected. Such QTLS detected through high throughput genotyping were not reported earlier. One of the reason could be the employment of unique germplasm in the present study which consistently showed high GPC (12%) as compared to low yielding counterpart (8%) leading to high genetic variation. The most stable QTL detected in our investigation qGPC1.1 was validated in an additional environment (Env.3) employing high throughput phenotyping technique. Another putative QTL for SGPC (qSGPC1.1) was found pleiotropic to the former. Inside this QTL region, one gene (Os01g0111900) was found which encoded
glutelin family protein. Physical (0.6–1 Mb) and linkage map (7–10 cM) position was highly corresponding. RiceXPro database revealed upregulation of this gene in endosperm during seed development. This corresponded with higher glutelin content in introgressed lines. Positive correlation was reported earlier between total protein and glutelin content in rice grain6. In our experiment, we also found similar trend in NILs with high GPC.

Fine mapping of this region using mapping population derived from high-protein-NIL × Naveen is in progress to detect tightly linked marker for marker assisted selection. Another stable QTL, qSGPC7 was detected near to a cluster of genes encoding three peptidase proteins53. More than 15 genes are responsible for glutelin synthesis which accounts more than 80% of the total seed storage protein. We reported a few other probable functional genes which were located inside or adjacent to the identified QTLs for GPC and SGPC in the present study.

Materials and Methods

Plant materials and development of mapping population. Through evaluation of 248 germplasm we identified a few (Supplementary Table 15) which consistently showed high protein content in both brown and polished rice. But they were mostly low yielders (<3000 kg/ha grain yield). The ARC 10075 was one of them with an average 12–13% GPC in brown rice25. This germplasm was crossed with a high yielding (4500 kg/ha grain yield) popular variety, Naveen with an average 8% GPC. F1 plants were backcrossed with the recurrent parent, Naveen.
to get 25 BC₁F₁ lines. Finally, 200 lines of BC₃F₄ and BC₃F₅ were developed by three consecutive backcrossing followed by single seed descent method.

Field experiments and phenotypic evaluation. One hundred ninety lines from the backcross population (BC₃F₄) were planted in three rows, 15 plants in each row with 20 cm row to row and 15 cm plant to plant spacing in augmented randomized block design along with replicated checks (Naveen and ARC 10075) following standard package of agronomic practices in kharif season 2013 (Env.1), rabi season 2014 (Env.2) and kharif season 2014 (Env.3) at the experimental farm of ICAR-National Rice Research Institute, Cuttack, Odisha, India. Ten randomly selected plants were used to study agronomic traits, including plant height (PH), maturity duration (MD), number of panicles/plant (PN), panicle length (cm) (PL), number of grains/panicle (GRAIN), 100 grain weight (GWT) and plant yield (PY). Seven selected introgression lines (BC₃F₅) with high GPC across the three environments and phenotypic resemblance with the recurrent high yielding parent, Naveen were again raised in replicated plots (25sq m) in kharif (Env.4) and rabi season 2015 (Env.5). Nitrogen, phosphorus, and potassium were supplied @ 80 kg, 60 kg, and 40 kg per hectare, respectively in Env.4 and @ 120 kg, 60 kg and 60 kg per hectare, respectively in Env.5. Phosphorus (as single super phosphate) was applied as a basal dose, and half of the total nitrogen (as urea) and potassium (as muriate of potash) were applied in basal and rest half were applied in two equal doses at 30 days after transplanting and at 50% flowering. The grain yield from the 25 sq m plot was converted to kg/ha.

Estimation of grain protein content. The GPC of all entries in Env.1 and Env.2 and also from the replicated large plots of selected lines in Env.4 and Env.5 were determined by the standard micro-Kjeldahl method by taking ten grains of brown rice (grains devoid of husk, but with the brown bran layer intact). The grain protein content was calculated by multiplying percent nitrogen content by 5.95. Single grain protein content (SGPC) was estimated on weight basis (mg/g) from the average protein content of 10 grains. Samples of known values for GPC of these lines and other germplasm were used in calibration and validation of NIR spectrophotometry for GPC in brown rice in our laboratory. The apparent grain protein content of mapping population in kharif season 2014 (Env.3) was estimated using 15 g dehusked grain in calibrated NIR spectroscopy.

Fractionation of grain protein and estimation of grain quality traits. Extraction of rice proteins was performed by standard protocol. Rice flour (6–7 g) was defatted with n-hexane. Standard steps were followed to separate protein fractions in the order of albumins, globulins, prolamin and glutelins. The extracted proteins were freeze-dried and stored at −70 °C. The protein content of each fraction was measured according to Lowry et al. The amylose content was measured as par standard procedure. Briefly, 100 mg sample was wetted with 1 ml ethanol followed by addition of 9 ml 1 N NaOH with shaking and placing the tube in a boiling water bath for 10 min. After adding 1 ml 1 N acetic acid and 2 ml Iodine reagent, the volume was made to 100 ml with water and the absorbance was measured at 620 nm. Gelatinization temperature was indirectly estimated in terms of the extent of alkali spreading value (ASV) measured using a seven-point scale ranging from score-1 (least spread) to score-7 (highest spread). The analysis of cooked rice elongation, CRE % = (ACL − BCL)/BCL × 100 (ACL: after cooking length, BCL: before cooking length) and other cooking parameters were done as described by Wang et al.
Statistical analysis. The phenotypic data were subjected to analysis of variance, genotypic and phenotypic coefficient of variances, genetic advance and heritability by using statistical package WINDOSTAT version 8.6, Indostat Service, Hyderabad. Heritability (h²) in broad sense is calculated from \( \sigma_g^2 / \sigma_p^2 \), where \( \sigma_g \) is genotypic variance and \( \sigma_p \) is phenotypic variance. Phenotypic data were statistically analysed and the normal distribution of phenotypic data was verified by K-S test at level of \( \alpha = 0.01 \) by using software, SPSS version 15.0 (SPSS, Chicago, IL, USA). For genotype × environment interaction studies, ANOVA was performed considering independent variables viz. genotype, environment, blocks within environment and genotype × environment as fixed effects and GPC and SGPC as the response variables using PROC GLM following the standard procedures.\(^{61,62}\) Graphs were plotted using PROC SGPLOT procedure of SAS 9.3 software. The t-test was employed for detection of significant differences if any for mean GPC and SGPC in multi-environments.

SNP array design and validation. Seventy one mer 50,000 SNP sequences were downloaded from OryzaSNP@MSU databases (http://rice.plantbiology.msu.edu). These SNPs were found uniformly distributed throughout the 12 chromosomes having good representation from coding and UTR regions. They were taken mostly from single copy (SC) genes and multi-copy rice (MCR) genes and the rest from agronomically important cloned rice genes (AGCR)\(^{31}\). The SNP sequences were shared with Affymetrix Bioinformatics team at Santa Clara, California, US for in-silico selection of markers for chip design. In-silico validation of the assay involved preliminary screening of the designed array file for each selected SNP. Both forward and reverse probes of each SNP were assigned with p-convert values, derived from a random forest model to predict the probability of SNP conversion on the array. The model considers factors including the probe sequence, binding energy and expected degree of non-specific hybridization to multiple genomic regions. SNP probes with high p-convert values are expected to convert on the SNP array with a high probability. Potential probes were designed for each SNP in both the forward and reverse direction, each of which was designated as ‘recommended,’ ‘neutral,’ or ‘not recommended’ based on p-convert values through which the SNP data sets were easily filtered. Thus, SNP probes were designed by screening 50,000 SNP loci of which an extremely high proportion of 40, 894 loci (90.8%) showed high-quality scores with p-convert values of >0.40, and the vast majority of them having p-convert values of >0.6, which were successfully synthesized on the array chip. The SNPs that were highly repetitive in the genome and contained ambiguities were removed. The resulting SNPs, selected for uniform spacing across the genome not having any other SNP, indel, translocation within 10 bp were selected for high resolution mapping of genetic loci in complex traits.

Genomic DNA preparation, SNP genotyping, allele calling and data analysis. Genomic DNA was extracted from young leaf tissues of 10 seedlings of parental lines and each of the 190 lines using CTAB method\(^{49}\). The quantity and quality of genomic DNA of each sample was determined using nano-drop spectrophotometer and 1% agarose gel. The samples with OD\(_{260}/OD_{280} \geq 1.8 \) and OD\(_{260}/OD_{230} \geq 1.5 \) and more than 10 Kb intact genomic DNA were used for SNP genotyping. An aliquot of 20\( \mu \)l (a total of 200 ng) of diluted gDNA from each sample was used for target probe preparation and genotyping using high-resolution Affymetrix custom array of 40894 SNP chip. The assays were performed on Gene Titan platform; the high throughput automated working station. Microarray tiled with probes specific to a genomic position of interest. Amplified total genomic DNA was fragmented it and hybridized to the array. Hybrization solution probes (9 mer) was paired a specific “happen” to a specific base. DNA ligation was used to covalently bind only the correct base followed by washing, staining, fixing and scanning. Hybridization to the Bead Chip and imaging of the arrays were performed by the Imperial Life Science (P) Ltd., Gurgaon, Haryana, India. The Affymetrix Gene Titan assay was based on 2 colors for genotyping; one probe for heterozygous locus detection while 2 probes for homozygous locus, by an allele-specific single base extension/ligation step. The data files generated after scans were CEL files. The analysis was performed on Affymetrix Genotype Console (GTC) Software version: 4.1. The samples below of DQC <0.82 and SNP call rate <95% were removed from the analysis and genotyping call was directly exported from the software. For clustering of SNP, we also used GTC software to call as separate homozygous and heterozygous cluster.

Linkage map construction and QTL analysis. All polymorphic markers including segregation distortion loci were mapped using SLD option (segregation distortion locus mapping) taking inclusive composite interval mapping (ICIM) and interval mapping (IM) implemented in QTL IciMapping V4 (http://www.isbreeding.net). SLD mapping using this software helped in restoration of order and position of the distorted markers on linkage map. This was additionally verified by Distorted Map v.1 software\(^{62}\). For identification of main effect of additive and digenic epistatic QTL in each environment and for each trait, the ‘IM-ADD,’ ‘ICIM-ADD’ and ‘ICIM-EPI’ functions, respectively, of the software were utilized\(^{64,65}\). Logarithm of odds (LOD) score peaks \( \geq 2.5 \) were used to declare the presence of a putative QTL in a given genomic region. A threshold LOD of 5.0 with probability values for entering variables (PIN) of 0.01 was used to declare significant epistatic-QTLs. The ‘Multi-Environment Trials’ (MET) function of the software was also utilized to determine the consensus positions for the major QTL and identification of additive × environment interaction effect QTLs (AE-QTL). MET- QTLs were considered if they accounted for >5% of the variance.

Bioinformatics tool to identify functional genes located inside or close to the identified QTLs. Genes directly related to the synthesis of storage proteins of rice grain, viz. glutelin, globulin, prolamin and albumin, were downloaded along with their physical position from Rice Annotation project Database\(^{65}\) and Oryzabase\(^{66}\). Functionally validated genes related to increase in grain protein content were also downloaded along with their physical positions from the gene information table available in QTL Annotation Rice Online Database\(^{67}\). The gene located inside the QTL interval region or within 1.0 Mb or nearly 4 cM (considering average
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Author Contributions
K.C. developed mapping population; K.C., B.C.M., L.K.B. and A.S. conducted field experiments and statistical analysis; L.B. and N.U. selected and filtered SNPs for designing array; K.C., L.B., N.U., S.R., M.C. and N.R.P. analyzed genotyping data and conducted genetic analysis; A.D., T.B.B., S.S., S.G.S., S.S.S. and N.M. analysed biochemical parameters, K.C., L.B., K.O.C., M.C., S.R., N.U. and S.G.S. interpreted the results and wrote the manuscript; K.C., L.B. and S.G.S. conceived, designed and supervised the study.

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